Organic enrichment in droplet residual particles relative to out of cloud over the northwest Atlantic: Analysis of airborne ACTIVATE data

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23 Abstract.

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24 Cloud processing is known to generate aerosol species such as sulfate and secondary organic aerosol, yet there is a scarcity of airborne data to examine this issue. The NASA Aerosol 25 Cloud meTeorology Interactions oVer the western ATlantic Experiment (ACTIVATE) was 26 designed to build an unprecedented dataset relevant to aerosol-cloud interactions with two 27 28 coordinated aircraft over the northwest Atlantic, with aerosol mass spectrometer data used from 29 four deployments between 2020-2021 to contrast aerosol composition below, in (using a counterflow virtual impactor), and above boundary layer clouds. Consistent features in all time 30 periods of the deployments (January-March, May-June, August-September) include the mass 31 fraction of organics and relative amount of oxygenated organics (m/z 44) relative to total organics 32 (f44) increasing in droplet residuals relative to below and above cloud. Detailed analysis comparing 33 data below and in cloud suggests a possible role for in-cloud aqueous processing in explaining 34 35 such results; an intriguing aspect though requiring more attention is that only approximately a 36 guarter of the cloud cases (29 of 110) showed higher organic mass fractions either below or above 37 cloud. Of those 29 cases, the majority (25) showed higher organic mass fraction below cloud base 38 where the cloud processing signature is presumably more evident as compared to above cloud. These results are consistent with the few past studies analyzing droplet residuals pointing to higher 39 organic enrichment than in adjacent cloud-free areas. The data findings are important as other 40 41 datasets (e.g., reanalysis) suggest that sulfate is both more abundant than organics (in contrast to 42 this work) and more closely related to drop number concentrations in the winter when aerosolcloud interactions are strongest; here we show that organics are more abundant than sulfate in the 43 droplet residuals and that aerosol interaction with clouds potentially decreases particle 44 hygroscopicity due to the <u>increase</u> in organic:sulfate ratio for droplet residuals relative to 45 surrounding cloud-free air. These results are important in light of the growing importance of 46 organics over the northwest Atlantic in recent decades relative to sulfate owing to the success of 47 48 regulatory activity over the eastern United States to cut sulfur dioxide emissions.

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52 1. Introduction

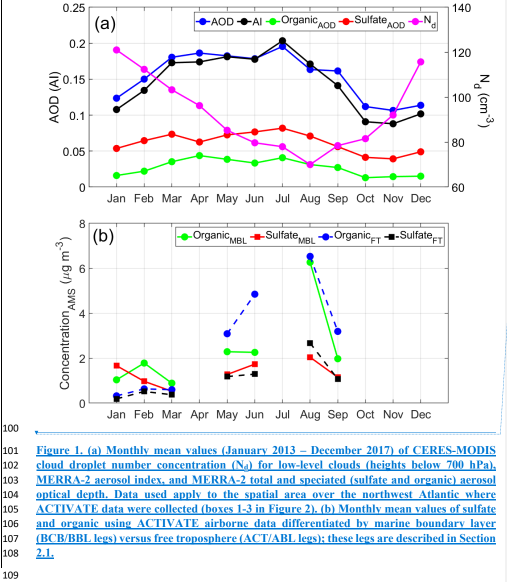
The nature of aerosol-cloud interactions over the northwest Atlantic Ocean is uncertain 53 even though the region has been the target of decades of atmospheric research (Sorooshian et al., 54 55 2020). These interactions include a subset of aerosol particles called cloud condensation nuclei 56 (CCN) that activate into cloud droplets, which subsequently undergo aqueous processing to transform into a particle after evaporation varying in size and composition relative to the original 57 CCN. An aspect of these steps that is poorly characterized is the composition of the droplet 58 59 residuals in cloud relative to particles below and above clouds, which requires airborne measurements. The NASA Aerosol Cloud meTeorology Interactions oVer the western ATlantic 60 Experiment (ACTIVATE) was designed to collect in situ and remote sensing data in and around 61 62 clouds during different seasons in a region with a wide range of weather conditions (Painemal et 63 al., 2021) and air mass sources (Corral et al., 2021), qualifying as a suitable dataset to examine 64 this very issue.

65 The annual cycle of aerosol and cloud drop number concentrations (Nd) varies northwest Atlantic, with aerosol parameters (e.g., aerosol optical depth, aerosol index) peal 66 summer months in contrast to N_d being highest the winter (Figure 1). This discrepand 67 68 reconciled by Dadashazar et al. (2021a) who showed that conditions linked to cold air ou events (e.g., enhanced turbulence, higher marine boundary layer (MBL) height, higher low 69 liquid cloud fraction) promote stronger aerosol-cloud interactions in the winter to help a 70 particles into drops with higher efficiency than other times of the year. Gradient boosted reg 71 72 tree analysis revealed that the most influential aerosol parameter in predicting N_d was either s 73 mass concentration of sulfate (winter) or organics (summer). However, those results were 74 on reanalysis data without any indication of causal effects between aerosol composition and 75 microphysics. Airborne in situ data are needed to unravel the composition details in and 76 clouds. Of particular interest related to aerosol chemical characterization around clouds is g 77 evidence in the literature that in-cloud aqueous processing can generate not only sulfate (B al., 2000; Ervens, 2015) but also secondary organic aerosol (SOA) (Blando and Turpin, 78 79 Warneck, 2003; Sorooshian et al., 2006a; Ervens et al., 2011; Heald et al., 2011), w 80 hypothesized to manifest itself in enhanced organic mass fractions in droplet residuals rela 81 below and above cloud. Past work over the northwest Atlantic has pointed to the importa 82 secondary formation via gas-to-particle conversion processes in influencing the organic budget of aerosol particles (de Gouw et al., 2005; Schroder et al., 2018; Shah et al., 83 Furthermore, chemical analysis of droplet residuals can potentially lend insight into proper 84 85 the CCN activating into droplets from the current or a future cloud passage, with pas showing an important role for organics (Russell et al., 2000; Drewnick et al., 2007; Mertes 86 87 2007; Hawkins et al., 2008; Asa-Awuku et al., 2015).

88 The goal of this study is to compare aerosol mass spectrometer data over the nor 89 Atlantic below, in, and above clouds for different times of the year (January-March, May August-September). Case studies of flights during cold air outbreaks probe deeper to 90 understand the nature of aerosol and droplet residual particle composition during these even 91 stronger aerosol-cloud interactions as compared to other times of the year (Dadashazar 92 93 2021a; Painemal et al., 2021). The results have implications for aerosol-cloud interacti 94 droplet residual composition is shown here to deviate from that of aerosol out of cloud. 95 important to lend insight into properties of the CCN activating into drops and/or pointing to 96 role for cloud processing over the northwest Atlantic to alter aerosol properties.

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110 2. Methods

111 2.1 Field Campaign Description

We use airborne in situ data collected aboard the HU-25 Falcon from deployments 1 (14 113 February - 12 March 2020), 2 (13 August - 30 September 2020), 3 (27 January - 2 April 2021), 114 and 4 (13 May - 30 June 2021) of the ACTIVATE mission. Data necessary for this study were 115 only available for two flights in deployment 3 (29 January and 3 February) owing to an aircraft 116 117 maintenance issue reducing the size of the available payload. ACTIVATE employs a dual aircraft 118 approach with the Falcon acquiring in situ data for trace gases, aerosol particles, and clouds in the MBL while a King Air flies overhead at ~9 km conducting remote sensing measurements and 119 launching dropsondes (Sorooshian et al., 2019). Typical flights are ~3-4 hours based out of NASA 120 Langley Research Center in Hampton, Virginia. The Falcon flies in what are termed "ensembles", 121 which comprise legs in the following nominal order: below cloud base (BCB), above cloud base 122 (ACB), BCB, ACB, minimum altitude leg at ~150 m (Min. Alt.), above cloud top (ACT), below 123 cloud top (BCT), and then descent back to BCB to start a new ensemble. Cloud-free ensembles 124 include the following legs: Min. Alt., below boundary layer top (BBL), above boundary layer top 125 126 (ABL), and then descent back down to Min. Alt. to start a new ensemble. The Falcon flies at ~120 127 m s⁻¹, with the duration (length) of each leg and cloud ensemble being ~3.3 min (~24 km) and 35 min (~250 km), respectively. Cloud-free ensembles were approximately 15 min (~100 km). The 128 129 repeated nature of these ensembles has built a large statistical database relevant to aerosol-cloud-130 meteorology interactions. <u>Clear</u> ensembles were generally closer to the coast.

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132 2.2 Airborne Instrument Details

The central dataset relevant to aerosol composition in this study comes from the Aerodyne 133 High-Resolution Time-of-Flight Aerosol Mass Spectrometer (AMS) (DeCarlo et al., 2008). The 134 135 instrument measures submicrometer non-refractory aerosol composition in 1 Hz Fast-MS mode with data averaged to 25-second time resolution. We make use of specific mass spectral markers 136 137 including m/z 43 (mostly $C_2H_3O^+$) and 44 (CO_2^+), which represent oxygenated organic fragments, 138 with the ratios of the markers relative to total organic mass referred to as f_{43} and f_{44} , respectively. 139 AMS measurements were conducted downstream of an isokinetic double diffuser inlet (Brechtel 140 Manufacturing Inc.) in cloud-free conditions and downstream of a counterflow virtual impactor 141 (CVI) inlet (Brechtel Manufacturing Inc.) in clouds (Shingler et al., 2012). For classification of data as cloud and cloud-free, we use a liquid water content (LWC) threshold of 0.05 g m⁻³ based 142 143 on data from the Fast Cloud Droplet Probe (FCDP; $D_p \sim 3 - 50 \ \mu m$) (SPEC Inc.; Kirschler et al., 144 2022). This LWC threshold has been used in recent work using ACTIVATE data (Dadashazar et 145 al., 2021a). Data for both rain water content and ice water content were used from a twodimensional stereo probe vertical direction (2DS-V; $D_p \sim 29 - 1465 \mu m$). We also use a proxy for 146 hygroscopicity in the form of f(RH), which is the ratio of total light scattering between relative 147 148 humidities of 80% and 20% as measured by tandem nephelometers (TSI Inc, St. Paul, MN, USA; Model 3563) (Ziemba et al., 2013). 149

Note that while cloud water samples were also chemically characterized, those data are outside the scope of this work as the partial speciation of organics in the cloud water samples makes it hard to compare to AMS total organics. Furthermore, particle-into-liquid sampler (PILS) data are not used owing to lengthier time resolution (~5 min) and chemical smearing during sample collection (Sorooshian et al., 2006b) preventing a clear assignment of data to individual legs in ensembles.

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Deleted: data for out-of-cloud and in-cloud data, and (ii) because the

Deleted: organic fraction could not be quantified owing to only being able to speciate selected organic acids.

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166 2.3 Complementary Datasets

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167 2.3.1 HYSPLIT and CWT Maps

We obtained 5-day back-trajectory data from NOAA's Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Stein et al., 2015; Rolph, 2017) ending at the Falcon position during any of the 29,164 cloud-free AMS data points, Note that this includes data during cloud ensembles but only when cloud liquid water content was < 0.05 g m⁻³, and thus data during BCB and ACT legs are included, We relied on the National Centers for Environmental Prediction/National Center for Atmospheric Research (NCEP/NCAR) reanalysis data using the "Model vertical velocity" method and obtained data points every 6 hours along trajectories.

As this study is mainly focused on organics and sulfate, concentration-weighted trajectory (CWT) maps were generated using HYSPLIT back-trajectories in conjunction with speciated AMS data (Figures S1-S2) to show the predominant sources for each of these two aerosol components (e.g., Hsu et al., 2003). As demonstrated by past works for other regions (e.g., Dadashazar et al., 2019), the method assigns a weighted concentration to grid cells based on mean concentrations passing through each grid cell from all the considered trajectories. CWT profile maps are produced using the GIS-based software called TrajStat (Wang et al., 2009).

182 **2.3.2 MERRA-2**

We use both total and speciated (sulfate and organic) aerosol optical depth (AOD) at 550 nm from the Modern-Era Retrospective analysis for Research and Applications-Version 2 (MERRA-2) (Gelaro et al., 2017) between January 2013 and December 2017 near Aqua's overpass time (13:30 local time). We also show results for aerosol index (AI), which is the product of AOD and the Ångström parameter. As the latter accounts for aerosol size, AI is better related to columnar CCN as compared to AOD (Nakajima et al., 2001). Data are used for the spatial area over the northwest Atlantic where ACTIVATE data were collected (boxes 1-3 in Figure 2).

190 2.3.3 CERES-MODIS

Cloud droplet number concentrations (Nd) are presented for the ACTIVATE region 191 192 following the specific calculations and filtering methods of Dadashazar et al. (2021a) using Clouds and the Earth's Radiant Energy System (CERES) edition 4 products (Minnis et al., 2011; Minnis 193 et al., 2021). CERES retrieval algorithms are applied to MODerate resolution Imaging 194 195 Spectroradiometer (MODIS)-Aqua radiances as obtained during daytime overpasses around 13:30 196 local time. Level 3 cloud data were used between January 2013 and December 2017 at $1^{\circ} \times 1^{\circ}$ resolution for low-level clouds (> 700 hPa) based on CERES-MODIS edition 4 Single Scanning 197 198 Footprint (SSF) products (Loeb et al., 2016). N_d was calculated with an adiabatic cloud model (Grosvenor et al., 2018): 199

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$$N_d = \frac{\sqrt{5}}{2 \pi k} \left(\frac{f_{ad} C_w \tau}{Q_{ext} \rho_w r_e^5} \right)^{1/2}$$

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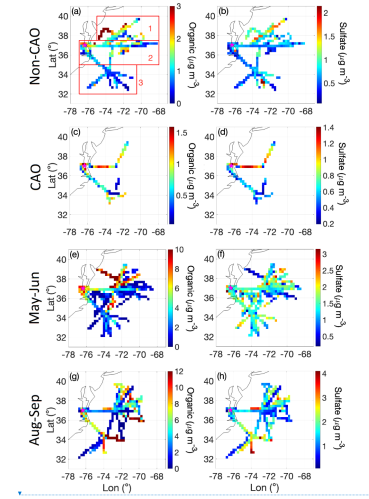
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207 208 where k represents the droplet spectrum width (assumed to be 0.8 over the ocean), re is cloud drop effective radius, τ is cloud optical depth, Q_{ext} is the unitless extinction efficiency factor (assumed 209 to be 2 for liquid droplets), and ρ_w is the density of water (1 g cm⁻³). N_d data are used when low-210 211 level liquid cloud fraction exceeded 40%. Data are used for the same spatial area as MERRA-2 data (i.e., boxes 1-3 in Figure 2). Deleted: 1 212 213 214 2.4 Classification of Cold Air Outbreak Flights Deleted: flights We determine whether flights occurred during cold air outbreaks (CAOs) leveraging 215 216 methods in recent ACTIVATE studies (Seethala et al., 2021; Corral et al., 2022). Briefly, Visible 217 Infrared Imaging Radiometer Suite (VIIRS) imagery (NASA Worldview) is used to visually identify cloud streets that are characteristic of CAOs. Flight notes and weather forecast slides were 218 219 used as additional confirmation, followed by data from dropsondes released from the King Air 220 following the method described in Papritz et al. (2015). 221 222 3. Results Deleted: 3. Formatted: List Paragraph, Numbered + Level: 1 + 223 A motivation of this study is the opposite annual pattern of N_d and aerosol parameters Numbering Style: 1, 2, 3, ... + Start at: 1 + Alignment: Left + Aligned at: 0" + Indent at: 0.25" 224 shown in Figure 1a. Notable is that sulfate AOD exceeds that of organic AOD for all months based 225 on MERRA-2 data, which has been shown before in the region (Braun et al., 2021). The 226 ACTIVATE airborne data show that while the total concentrations of both aerosol components are 227 higher in the summer months (similar to related aerosol parameters in Figure 1a), a difference 228 compared to MERRA-2 speciated AODs is that organic levels exceed those of sulfate (except 229 January in the MBL), regardless of whether the data were collected in the MBL (i.e., BBL and 230 BCB legs) or free troposphere (i.e., ACT and ABL legs) (Figure 1b). Hegg et al. (1997) concluded 231 for the month of June based on a chemical apportionment study using aerosol column optical depth 232 data off the mid-Atlantic coast of the United States that the three most abundant components (in 233 decreasing order) were water, carbonaceous compounds, and then sulfate. This is an important 234 result with implications for aerosol characteristics such as hygroscopicity. For instance, higher 235 organic:sulfate mass ratios in the MBL correspond to suppressed hygroscopic growth factors at 236 high relative humidities ($\geq 85\%$) (Hersey et al., 2009). For comparison, airborne measurements 237 in winter and summer periods over the eastern North Atlantic (Wang et al., 2022) showed sulfate concentrations exceeding those of organics up to the same altitudes in this study (~ 1.6 km). 238 239 240 3.1 Multi-season **Overview** of AMS Composition **Deleted:** overview **Deleted:** composition 241 Relative to all AMS species, sulfate and organics are the dominant aerosol components by mass with combined mass fractions being near 75% usually regardless of season or location 242 243 relative to clouds (Tables S1-S2; spatial maps in Figure 2); this is consistent with their predictive Deleted: Table 1 244 capability for N_d over the northwest Atlantic (Dadashazar et al., 2021a). Nitrate and ammonium Deleted: 1 245 were the next most abundant components, with chloride being much lower. The highest organic 7

253 254 255 256 257 258 259 260 261 262 262 263	concentrations were in August-September assisted in part by transported wildfire emissions from western North America (Mardi et al., 2021). Mean vertical profiles of organics in each season (Figure 3) show that in all months, but especially May-June and August-September, there is an enhancement at altitudes exceeding 200 m in the northernmost parts of the study region. Organic aerosol CWT maps reveal significant influence from continental sources based on the highest concentrations along trajectories coming from the U.S. East Coast (Figure 51). In terms of the nature of the organic aerosol fraction, vertical profiles of f_{44} were fairly similar between seasons and areas of the study region (Figure 2), ranging in mean value for the various leg types in Table 51 between 0.11 and 0.27. For reference, the f_{44} of atomized oxalic acid, a tracer for cloud processing in the absence of biomass burning and coarse aerosol (Hilario et al., 2021 and references therein), is 0.36 (Lambe et al., 2011).	Deleted: S5 Deleted: 2 Deleted: S5 Deleted: 1
264	In contrast to organics, sulfate exhibits more spatially homogenous concentrations over the	
265	northwest Atlantic (Figure 2) owing largely to ocean-emitted dimethylsulfide that undergoes gas	Deleted: 1
266	and in-cloud oxidation such as what was shown for the eastern North Atlantic (Ovadnevaite et al.,	
267	2014). This is supported by how sulfate's seasonal CWT maps (Figure <u>\$2</u>) differ from those of	Deleted: 3
268	organics with comparable concentrations widespread over the northwest Atlantic relative to the	
269	continent. The August-September CWT map for sulfate reveals more high concentration areas	
270	(note the different color bar scale for Aug-Sep in Figure <u>S2</u>) over the continent with concentrations	(Deleted: 3
271	exceeding those over most of the ocean; this is presumably due to more secondary formation	
272	stemming from local sulfur dioxide emissions over the eastern U.S. (Yang et al., 2018) aided in	
273	part by higher temperatures and humidity (Corral et al., 2021) that co-vary with other conditions	
274	favorable for sulfate production such as stagnation and certain air flow patterns (Tai et al., 2010).	
275	Figure 3 demonstrates that neither sulfate or organics exhibit a clear reduction with altitude	Deleted: S5
276	pointing towards a potential source aloft that might include long-range transport and/or secondary	
277	production.	
278	Although based on only two consecutive days of flight data, results from Leaitch et al.	
279	(2010) are relevant in that they sampled below, in, and above boundary layer clouds over the	
280	northwest Atlantic. On the first day with more marine influence, sulfate was more abundant than	
281	organics in fine particles below cloud. In contrast, the second day had more continental influence	
282	with organic levels exceeding those of sulfate below cloud, which was often the case during	
283	ACTIVATE (Table <u>S1</u>). They concluded with a parcel model that the impact of anthropogenic	Deleted: 1
284	carbonaceous components on the cloud albedo effect can exceed that of anthropogenic sulfate,	
285	which motivates attention to the droplet residual composition discussed next.	Deleted: , which is
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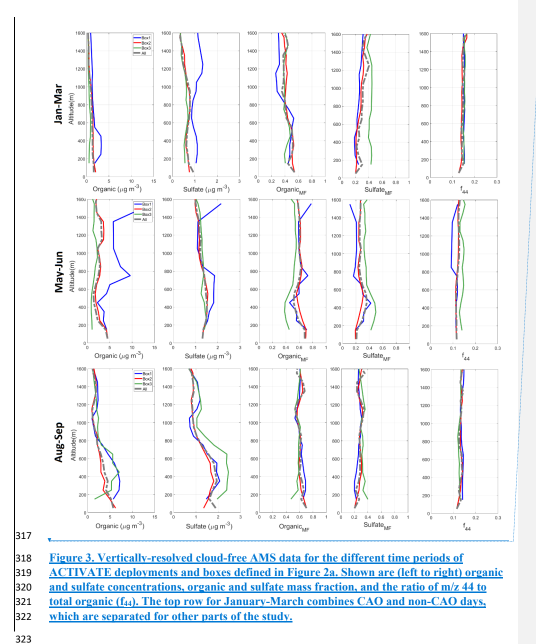
Deleted: Table 1. Average concentrations of submicrometer aerosol species measured by an airborne AMS for different seasons associated with ACTIVATE deployments 1-4. Non-CAO and CAO categories include samples collected between January and March. CVI = droplet residual particle measurements in cloud; BCB = below cloud base, ACT = above cloud top, BBL = below boundary layer top, ABL = above boundary layer top. Corresponding standard deviations and number of points are provided in Table S1.→¶

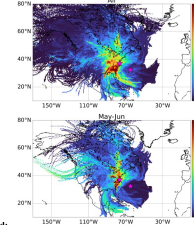
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	CVI	BCB
Organic (µg m ⁻³)	-	1.07/0.67/1.49/3.2
Sulfate ($\mu g m^{-3}$)	-	0.93/0.79/1.71/1.
Nitrate (µg m ⁻³)	-	0.40/0.21/0.07/0.
Ammonium (µg m ⁻³)	-	0.45/0.32/0.36/0.
Chloride (µg m ⁻³)	-	0.03/0.02/0.03/0.0
Organic _{MF}	0.55/0.60/0.68/0.61	0.40/0.34/0.35/0.4
Sulfate _{MF}	0.24/0.19/0.14/0.14	0.39/0.45/0.53/0.
Nitrate _{MF}	0.05/0.05/0.05/0.05	0.08/0.07/0.02/0.0
Ammonium _{MF}	0.09/0.08/0.07/0.09	0.13/0.13/0.10/0.0
Chloride _{MF}	0.06/0.08/0.06/0.10	0.01/0.01/0.01/0.0
f ₄₄	0.33/0.34/0.24/0.37	0.15/0.13/0.11/0.

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Figure 2. Spatial map of cloud-free AMS data for organics and sulfate collected during deployments 1-4 of ACTIVATE spanning from February 2020 to June 2021. Non-CAO and CAO represent non_ccold air outbreak and cold air outbreak days, <u>respectively</u>, between January and March. Spatial boxes labeled 1-3 in (a) correspond to domains used for calculations in other parts of this study. Grid cells are 0.25° × 0.25° and represent an average of data across all vertical levels flown between 0.02 and 8.1 km. Color bar scales differ by panel to highlight variability better within a panel.

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Deleted: 150°W 110°W 70°W 30°W Figure 2. Concentration weighted trajectory maps for organic aerosol concentrations as measured by an AMS on the Falcon during different ACTIVATE deployments (All data, Jan-Mar 2020 and 2021, May-Jun 2021, August-September 2020). These are based on 29,164 cloud-free AMS data points. The pink stars represent NASA Langley Research Center (Hampton, Virginia) and Bermuda for reference. Color bar scales differ to show variability better within a given panel.¶

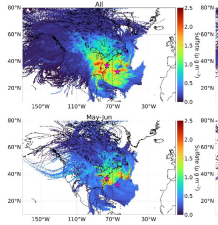


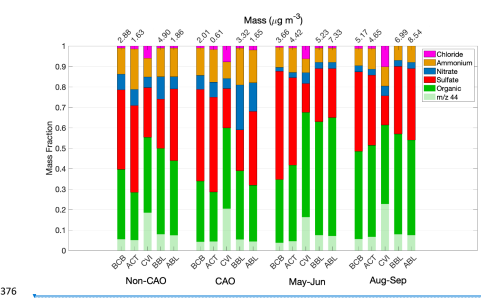
Figure 3. Concentration weighted trajectory maps for sulfate aerosol concentrations as measured by an AMS on the Falcon during different ACTIVATE deployments (All data, Jan-Mar 2020 and 2021, May-Jun 2021, August-September 2020). These are based on 29,164 cloud-free AMS data points. The pink stars represent NASA Langley Research Center (Hampton, Virgin(...[1])

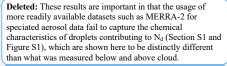
3.2 Droplet Residual Composition 363

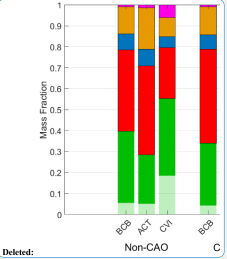
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A striking result in all seasons is that organic mass fraction was higher downstream of the 364 CVI in droplet residual particles in contrast to adjacent BCB and ACT legs in cloudy ensembles 365 (Figure 4 and Table S1). To compensate, sulfate mass fractions decreased in droplet residuals. 366 Furthermore, f44 increased in droplet residuals as compared to BCB and ACT data in each season, 367 368 indicative of more contribution of oxygenated organic species like carboxylic acids. There was no 369 significant difference in the mass fraction profiles between seasons for a fixed leg type (Figure 4).

370 The organic mass fraction and f44 changes in droplet residuals can be explained at least in 371 part by some combination of preferential activation of CCN with these special properties and/or 372 aqueous processing in droplets to generate oxygenated organics. Although not the focus here, the high chloride mass fractions in droplet residuals (Figure 4) can be explained by how sea salt would 373 374 preferentially activate into drops owing to its large size and that the AMS has some ability (albeit 375 not efficient) to detect sea salt chloride (Zorn et al., 2008; Ovadnevaite et al., 2012).









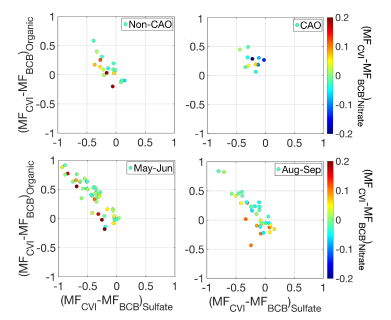
377 Figure 4. Seasonal comparison of AMS mass fractions, including the relative contribution of 378 m/z 44 to total organic (f₄₄). Numbers above each bar represent the mean total AMS mass concentration for that category; note that absolute masses are not reported downstream of

379 a CVI owing to high uncertainties. Note that the Non-CAO and CAO categories represent 380

381 all flight data in January-March (deployments 1 and 3) that were separated using the criteria

in Section 2.4. 382

We next examine scatterplots of organic mass fraction (i.e., organic mass divided by total 391 AMS mass) differences between each cloud leg with CVI-AMS data and its closest BCB leg in 392 the same cloud ensemble versus analogous sulfate mass fraction differences for the same pair of 393 legs (Figure 5). Aqueous processing to preferentially increase one of the two species relative to 394 395 the other would presumably translate into a positive value on the more preferred species' axis; in 396 other words, if there was more organic aerosol formation in clouds via aqueous processing 397 relative to sulfate, it would register as a positive (negative) value on the y (x) axis. Regardless of 398 season, the results reveal a consistent feature of increasing (decreasing) organic (sulfate) mass fraction downstream of the CVI relative to BCB samples, suggestive of aqueous processing 399 shifting the composition to be more organic-rich. For the very few points laying to the bottom 400 left of the origin, nitrate is often more enhanced in those droplet residual samples relative to 401 402 BCB data. Although not shown, the results in Figure 5 are similar to if ACT data were used in 403 place of BCB data.



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Figure 5. Scatterplot of the difference in organic mass fraction in cloud legs with CVI data and below cloud base (BCB) legs for an individual cloud ensemble relative to the analagous difference for sulfate mass fraction between the same pair of legs. Markers are colored by the analagous difference in nitrate mass fraction. Panels represent different seasons with winter deployments (January-March) separated into CAO and non-CAO days.

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413 difference between CVI and BCB/ACT data in each season (Figure 6). Ambient organic aerosol

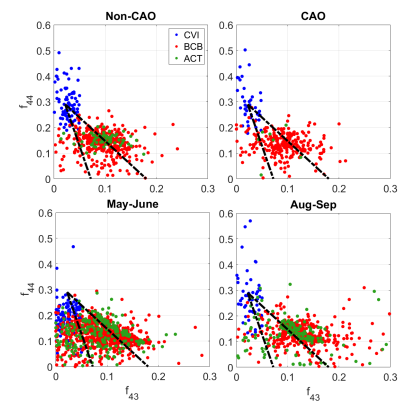
414 typically converge at the top left of the triangle representative of more atmospheric aging leading

415 to low volatility oxygenated organic aerosol species. The CVI data are systematically higher and

416 to the left of the triangle plot in each season. In contrast, the BCB and ACT data are lower and to

417 the right of the triangle plots without much distinction, suggestive of a similarly lower level of

418 oxygenation relative to droplet residuals.



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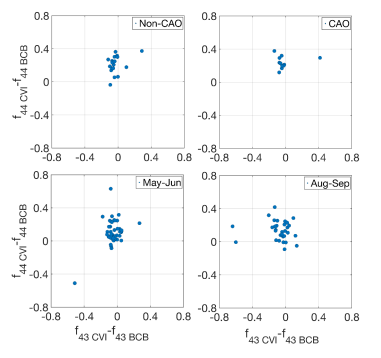
Figure 6. Comparison of f₄₄ and f₄₃ for individual BCB and ACT legs out of cloud, in addition
to CVI data in cloud legs. <u>Panels represent</u> different <u>seasons with winter</u> deployments
(January-March) separated into CAO and non-CAO days. Superimposed on the plots are
triangles corresponding to how former work (Ng et al., 2010) compared these ratios. Points
with organic mass concentration less than 0.5 µg m⁻³ were omitted from this analysis.

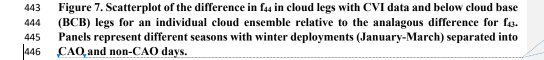
The CVI droplet residuals are more oxidized because of some combination of aqueous processing effects to yield more oxidized organic species, or because CCN with higher f₄₄ activated Deleted: Data are separated between time periods coinciding with ... Deleted: ACTIVATE

433 into droplets. To probe more into which of the two aforementioned processes could be more 434 responsible for the cluster of CVI points at the top left of the triangle plots, we next examine 435 (analogous to Figure 5) scatterplots of f_{44,CVI} - f_{44,BCB} versus f_{43,CVI} - f_{43,BCB}, where data are compared between the pair of cloud and BCB legs closest to one another in individual cloud 436 437 ensembles (Figure 7). If there was no difference in organic composition between a pair of legs, a marker representing that pair would be at the origin. Aqueous processing is presumed to result in 438 439 a positive (negative) value on the y (x) axis. Each season consistently exhibits points positioned to the top left of the origin suggestive of aqueous processing leading to the enhanced oxygenation 440 441 of the organic fraction in droplet residuals relative to BCB legs.

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Deleted: Note that this analysis omitted consideration of ACT legs as the predominant source of droplets is from activation of sub-cloud aerosol particles.





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447 A discussion on possible contributing factors (other than aqueous processing) to the
448 different chemical signature in CVI samples relative to adjacent cloud-free areas is warranted,
449 First, we note that 23% of BCB/CVI pairs of data points (25 out of 110) exhibited higher organic
450 mass fraction in the BCB leg relative to droplet residuals (Figure <u>8</u>). This number increases to 26%
451 when considering if either the BCB or ACT organic mass fraction was higher than the

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Y	Deleted: including processes occurring in the CVI inlet
	Deleted: S6), demonstrating that the null case exists without an organic enhancement downstream of the CVI. The

corresponding CVI data in cloud for an ensemble. Clearly the cases where a higher organic mass
 fraction was observed out of cloud seems to be most prevalent below cloud suggesting that location
 is where a cloud processing signature can be more reliably observed. These 26% of the cases
 studied demonstrate that the null case exists without an organic enhancement downstream of the

468 <u>CVI, reducing concerns over instrument and sampling artifacts.</u>

469 In terms of the contamination due to the inlet's material of construction, the CVI inlet was 470 designed with both stainless steel and aluminum yielding negligible organic contamination 471 (Shingler et al., 2012). A way to test this is with conducting CVI sampling in cloud-free conditions. 472 Figure S3 shows a representative time series of AMS data during a flight (research flight 10 on 28 473 February 2020) with numerous cloud passes and periods when there was still sampling 474 downstream of the CVI inlet outside of cloud. During those three key periods shown out of cloud 475 with CVI sampling, sulfate and organic levels exhibit concentrations close to zero and with 476 concentrations considerably lower than CVI data in cloud. Compared to sulfate, there is more 477 variability in organic levels downstream of the CVI regardless of whether sampling was in or out 478 of cloud or even whether sampling was done using the isokinetic inlet out of cloud. The data reveal 479 that at small time scales there is variability in the organic:sulfate ratio behind the CVI in cloud, specifically when comparing the clouds at 16:18-16:29 versus 16:39-16:43 with the former being 480 481 more organic rich. This representative time series provides confidence in the inlet itself not being 482 the source of the significant changes observed downstream the CVI throughout the first four 483 ACTIVATE deployments. This case flight is examined more in Section 3.3.

The heated counterflow in the CVI reduces positive artifacts from volatile gaseous species partitioning into sampled droplets such as with volatile organic compounds (VOCs) to form organics or with nitric acid to form nitrate (Prabhakar et al., 2014); in contrast, the heated counterflow would presumably evaporate some fraction of the existing nitrate and organics in the CCN that activated into droplets unlike sulfate which is not volatile. Thus, the heated inlet would tend to favor sulfate in the cloud droplet residuals and could not explain the enhanced organic residual observations here.

Inlets including the CVI can be prone to droplet shatter such as with large drizzle drops (> 100 μ m) (Twohy et al., 2013), although drizzle was not always frequent and the particulate artifacts generated would still be representative of droplet residuals. It seems implausible that such drop shatter would lead to an organic enrichment especially as the chemical results we report were consistent across the entire study region. AMS results were compared to both rain water content and ice water content without evidence of a distinct relationship between precipitation levels and whether or not there was a higher organic mass fraction behind the CVI relative to out of cloud.

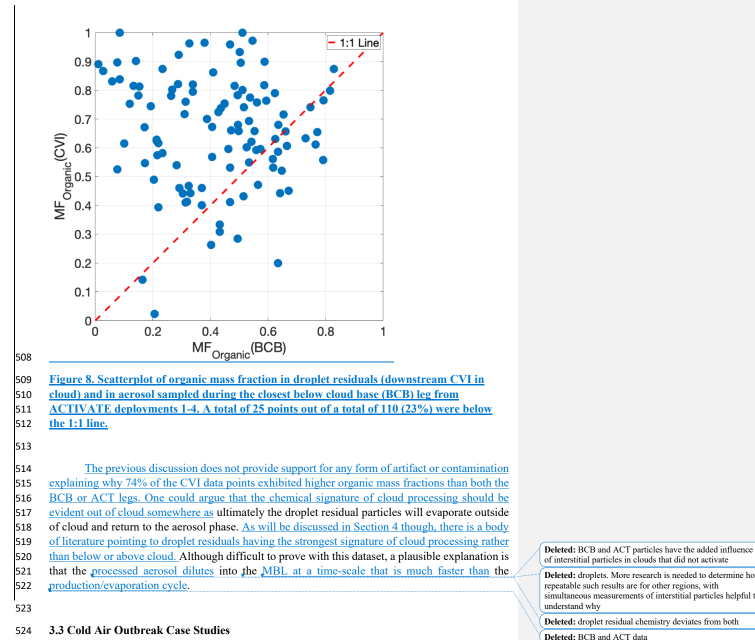
498 <u>It is also noteworthy that there can be considerable variability in AMS composition along</u>
 499 level legs (BCB, in cloud, ACT) pointing to how a signature of cloud processing out of cloud can
 500 be reduced when averaging data. Figure S3 demonstrates variability along individual legs that is
 501 not consistent with the organic:sulfate ratio always being enhanced downstream of the CVI.

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repeatable such results are for other regions, with simultaneous measurements of interstitial particles helpful to

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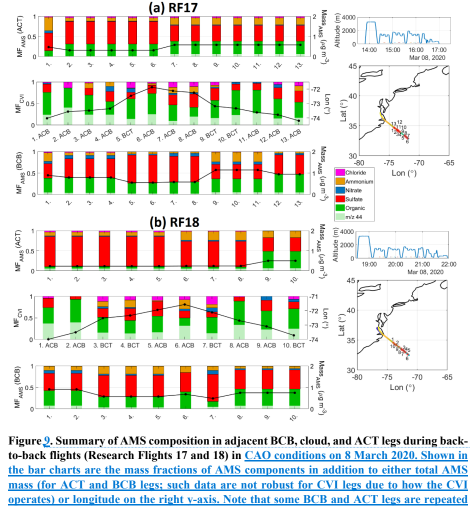
Owing to interest in the winter season having the strongest aerosol-cloud 533 534 interactions (Dadashazar et al., 2021a; Painemal et al., 2021), here we examine six_case study 535 research flights (RFs) during CAOs to understand the compositional characteristics below, inside, 536 and above clouds. We focus more on the representative day of 8 March 2020, (Figure 9), which 537 included two consecutive flights (RFs 17 and 18) based out of Hampton, Virginia profiling aerosol 538 and cloud properties in CAO conditions. These two flights were investigated in past work showing 539 enhanced new particle formation in ACT legs (Corral et al., 2022) and that entrainment of free 540 tropospheric air dilutes MBL CCN concentrations (Tornow et al., 2022). The other four flights 541 (Figure 10: RFs 5-6 on 22 February 2020; Figure 11: RFs 10-11 on 28 February 2020) exhibited the same general results as those shown for 8 March with higher organic mass fractions and f44 in 542 543 the cloud legs.

544 Figure 9 shows the AMS composition profile on the out-and-back flights on 8 March, 545 which involved flying out to a point and repeating the same path back to the airfield. Stacked on 546 top of each other in Figure 9 are the corresponding legs within individual cloud ensembles including (from top to bottom) ACT, either BCT or ACB legs with CVI data, and BCB. RF17 in 547 548 the morning comprised 13 different cloud legs with corresponding BCB and ACT legs. The BCB 549 and ACT mass fraction profiles were similar with sulfate being most abundant (mass fractions: 550 0.34-0.65) followed closely by organics (mass fractions: 0.15-0.42). The f₄₄ fraction of the 551 organics in BCB and ACT legs was quite low (0.00-0.16). The cloud data show a very different 552 profile with organics dominating the mass profile (mass fractions: 0.41-0.86) followed usually by sulfate (mass fractions: 0.00-0.30). Furthermore, there was a significant jump in f₄₄ in the CVI data 553 554 (0.21-0.48). RF18 later in the day re-traced the same flight path and included 10 sets of matching cloud-BCB/ACT legs showing again a similar jump in both organic mass fraction and f44 in droplet 555 residuals. In the second flight there was more variability in the BCB and ACT pairs, with higher 556 557 sulfate mass fractions (0.34-0.75) in the ACT legs throughout most of the flight excluding the last 558 two sets of legs. The total AMS mass concentrations were slightly higher in the BCB legs (0.49-559 0.91 µg m⁻³) relative to ACT legs (0.24-0.50 µg m⁻³).

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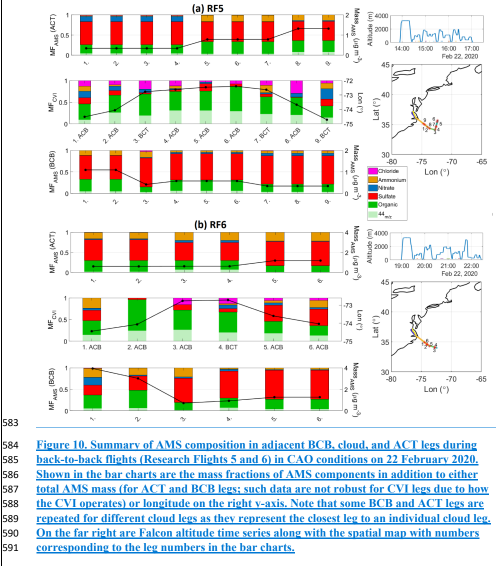
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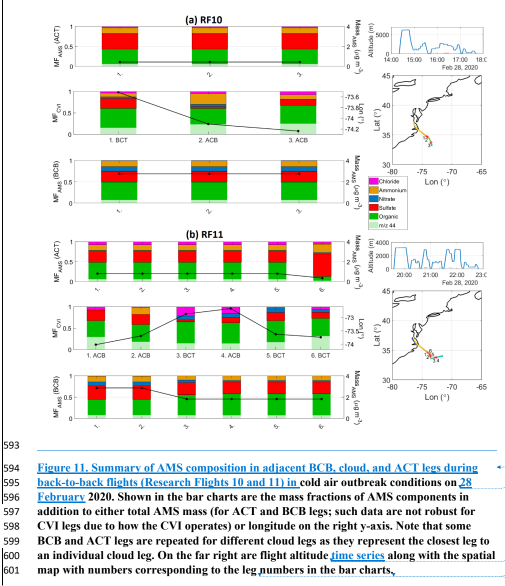
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for different cloud legs as they represent the closest leg to an individual cloud leg. On the far right are Falcon altitude during the flight along with the spatial map with numbers corresponding to the leg set numbers in the bar charts.

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603 4. Discussion

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608 Our results represent unique atmospheric data that are scarce in the literature owing to the difficulty of obtaining aerosol chemical data below, in, and above cloud in close spatiotemporal 609 610 proximity across many flights in different times of the year. Figure 1 provides implications of the results in terms of differences with MERRA-2 speciated AOD. Although we cannot 611 unambiguously prove it with the dataset, the results suggest that the most likely explanation for 612 613 organic and f₄₄ enrichment in droplet residuals has to do with aqueous processing rather than preferential activation of CCN with enhanced values of the organic:sulfate ratio and f₄₄. That the 614 615 droplet residuals shift to a more organic-rich signature with more oxygenated organics has implications for the aerosol particle properties remaining after droplet evaporation as they shift in 616 617 composition and size. Interestingly this study shows though that such a signature out of cloud was absent for 74% of the cloud cases as organic mass fraction was higher in cloud versus either below 618 or above cloud. These findings are significant in terms of motivating additional research, especially 619 620 as other studies discussed below also have shown higher levels of organic mass fraction of ratios of oxygenated organics relative to total organic mass in CVI samples as compared to out of cloud. 621

Coggon et al. (2012) showed increased AMS organic:sulfate ratios with altitude in the 622 623 MBL over the northeast Pacific Ocean coincident with increased liquid water content, which was 624 attributed to aqueous processing effects to generate more organics relative to sulfate; this was also suggested by past work in that region with a PILS (Sorooshian et al., 2007). Coggon et al. (2012) 625 626 showed that organics and sulfate were typically the most abundant AMS species both below cloud 627 and in droplet residuals with comparable mass fractions and no consistent trend of either one dominating the droplet residual composition; however, they showed that in 70% of their cloud 628 629 cases that the CVI data exhibited higher organic mass fraction relative to out of cloud. Past 630 measurements off the California coast and over Texas revealed enhanced f₄₄ values in droplet residuals relative to below and above cloud data and also relative to interstitial aerosol particles in 631 cloud (Sorooshian et al., 2010). That study showed similarly enhanced values of other ratios in 632 droplet residuals indicative of more oxygenated organics (e.g., PILS oxalate:AMS m/z 44, PILS 633 oxalate: AMS organic). Over the Texas area, PILS measurements of oxalate relative to AMS sulfate 634 and organic revealed significant enhancements (factors up to 4 and 13, respectively) downstream 635 636 a CVI relative to cloud-free conditions at similar altitudes (Wonaschuetz et al., 2012); furthermore, 637 they showed that organic mass fractions increased together with oxalate:organic and oxalate:sulfate ratios as a function of residual cloud fraction, which was a metric representing 638 639 "cloud processing history" of an air parcel in shallow cumulus cloud fields. CVI-AMS data from 640 a surface site studying warm tropospheric clouds on Mt. Åreskutan in central Sweden in July 2003 showed that organics and nitrate activated with higher ease than sulfate (Drewnick et al., 2007); 641 even though our results suggest the droplet residual changes in composition are likely driven by 642 aqueous processing, it is relevant that organics have been shown in at least another region to 643 644 activate more easily than sulfate.

645 While a measurement of hygroscopicity of the droplet residuals was not available, we 646 instead examine aerosol hygroscopicity from BCB legs as that is the area out of cloud most commonly exhibiting higher organic mass fractions relative to in cloud. Even if the signature out 647 of cloud is not as clear as one would expect presumably owing to dilution effects, still the 648 influence of cloud processing on organics inevitably should exist to some extent making the 649 650 subsequent discussion valuable. Having more organics relative to sulfate may reduce 651 hygroscopicity at high RHs (e.g., Hersey et al., 2009), but a compensating factor could be that 652 the organics are more oxygenated, which would increase the hygroscopicity of the organic 653 fraction itself.

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Moved down [1]: Having more organics relative to sulfate may reduce hygroscopicity at high RHs (e.g., Hersey et al., 2009), but a compensating factor could be that the organics are more oxygenated, which would increase the hygroscopicity of the organic fraction itself.

Moved down [2]: shows an inverse relationship between f(RH) and organic mass fraction across all the BCB legs in ACTIVATE deployments 1-4, which is similar to what has been observed over the continental U.S. (Shingler et al., 2016); using the linear best fit line shows that the representative f(RH) value for pure organic aerosol (i.e., organic mass fraction of 1.0) was 1.22 in contrast with 0.92 over the continental United States (Shingler et al., 2016). The f(RH) value for pure inorganic aerosol (i.e., organic mass fraction of 0.0) was 1.39.

Moved down [3]: along with previous discussion suggests that aerosol interaction with clouds decreases particle hygroscopicity at an RH of 80% although future work will look deeper into aerosol hygroscopic properties over the

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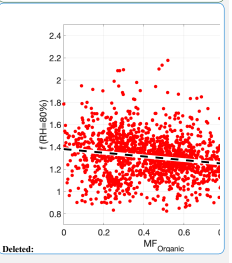
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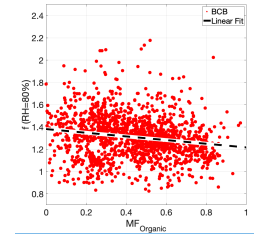
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Figure <u>12</u>. Relationship between f(RH) and organic mass fraction for BCB legs during ACTIVATE deployments 1-4. Markers are based on f(RH) data synched to the time resolution of the AMS data. The f(RH) values from the linear fit at a MF_{organic} value of 0.0 1.0 are 1.39 and 1.22, respectively.

701 Figure 12 shows an inverse relationship between f(RH) and organic mass fraction across 702 all the BCB legs in ACTIVATE deployments 1-4, which is similar to what has been observed over 703 the continental U.S. (Shingler et al., 2016); using the linear best fit line shows that the 704 representative f(RH) value for pure organic aerosol (i.e., organic mass fraction of 1.0) was 1.22 in 705 contrast with 0.92 over the continental United States (Shingler et al., 2016). The f(RH) value for 706 pure inorganic aerosol (i.e., organic mass fraction of 0.0) was 1.39. Results of Figure 12 along 707 with previous discussion suggests that aerosol interaction with clouds decreases particle 708 hygroscopicity at an RH of 80% although future work will look deeper into aerosol hygroscopic 709 properties over the ACTIVATE region. This is especially relevant as regulatory activities have reduced sulfate levels over the eastern U.S. in recent decades promoting higher relative amounts 710 of organics (Bates et al., 2005; Hand et al., 2012) with downwind impacts on the northwest Atlantic 711 712 due to offshore flow (Keene et al., 2014; Aldhaif et al., 2021; Dadashazar et al., 2021b). 713

714 5. Conclusion

715 A large airborne dataset collected over the northwest Atlantic as part of the NASA 716 ACTIVATE mission reveals a distinctly different chemical signature in cloud droplet residuals 717 (lower sulfate mass fraction, higher organic mass fraction, and higher f₄₄) relative to particles 718 below and above cloud for approximately 75% of the cloud cases examined. Several case study 719 flights during cold air outbreak conditions are profiled showing the aforementioned compositional 720 changes in droplet residuals. Detailed analysis suggests this shift in composition is driven more by 721 in-cloud aqueous processing rather than preferential activation of CCN with such chemical 722 characteristics. Of the 29 cases (out of 110) with higher organic mass fraction either above or Deleted: 9

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726 below clouds versus droplet residuals, 25 (4) exhibited higher organic mass fraction below (above) 727 cloud suggestive of the cloud processing signature being more prevalent below cloud. These 728 results are analogous to past work in other regions using different instrumentation showing maximum values of various metrics relevant to organics (e.g., f44, organic mass fraction) 729 730 downstream of a CVI in cloud relative to either below or above cloud (Sorooshian et al., 2010; 731 Coggon et al., 2012; Wonaschuetz et al., 2012). More work is needed to continue validating whether aqueous processing is the primary reason for the composition changes and to determine 732 733 if these results apply to other regions.

The results of this study <u>motivate</u> increased attention to both in-cloud formation of oxygenated organics and the composition of particles activating into droplets over the northwest Atlantic. <u>This work has implications for</u> aerosol-cloud interactions in this region as datasets often relied on in the absence of airborne data such as reanalysis data suggest a different story where sulfate is more enhanced than organics year-round (in contrast to the airborne data) (e.g., Braun et al., 2021). <u>The high relative abundance of organics needs more attention</u>, especially in light of the increasing relative amount of species in aerosol particles other than sulfate due to regulatory

741 activities over the U.S. (Hand et al., 2012).

Moved up [4]: Several case study flights during cold air outbreak conditions are profiled showing the aforementioned compositional changes in droplet residuals.

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751 Data Availability.

752	ACTIVATE Airborne	Data:
753	https://doi.org/10.5067/ASDC/ACTIVATE_Aerosol_AircraftInSitu_Falcon_	Data_1
754	(NASA/LARC/SD/ASDC,	2020a),
755	https://doi.org/10.5067/ASDC/ACTIVATE Cloud AircraftInSitu Falcon D	ata 1
756	(NASA/LARC/SD/ASDC, 2020b),	and
757	https://doi.org/10.5067/ASDC/ACTIVATE_MetNav_AircraftInSitu_Falcon_	Data_1
	• • •	

758 (NASA/LARC/SD/ASDC, 2020c).

Author contributions. HD conducted the analysis. AS and HD prepared the manuscript. All authors contributed by providing input and/or participating in airborne data collection.

761 *Competing interests.* The authors declare that they have no conflict of interest.

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Organic enrichment in droplet residual particles relative to out of cloud over the northwest Atlantic: Analysis of airborne ACTIVATE data Hossein Dadashazar¹, Andrea F. Corral¹, Ewan Crosbie^{2,3}, Sanja Dmitrovic⁴, Simon Kirschler^{5,6}, Kayla McCauley⁷, Richard Moore², Claire Robinson^{2,3}, Joseph Schlosser¹, Michael Shook², K. Lee Thornhill², Christiane Voigt^{5,6}, Edward Winstead^{2,3}, Luke Ziemba², Armin Sorooshian^{1,4,7} ¹Department of Chemical and Environmental Engineering, University of Arizona, Tucson, AZ, USÂ ²NASA Langley Research Center, Hampton, VA, USA ³Science Systems and Applications, Inc., Hampton, VA, USA ⁴James C. Wyant College of Optical Sciences, University of Arizona, Tucson, AZ, USA ⁵Institute of Atmospheric Physics, German Aerospace Center ⁶Institute of Atmospheric Physics, University Mainz, Germany ⁷Department of Hydrology and Atmospheric Sciences, University of Arizona, Tucson, AZ, USA

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23 Section S1. Discussion of Figure S1

A motivation of this study is the opposite annual pattern of N_d and aerosol parameters 24 shown in Figure S1a. Notable is that sulfate AOD exceeds that of organic AOD for all months 25 26 based on MERRA-2 data, which has been shown before in the region (Braun et al., 2021). The ACTIVATE airborne data show that while the total concentrations of both aerosol components are 27 28 higher in the summer months (similar to related aerosol parameters in Figure S1a), a difference 29 compared to MERRA-2 speciated AODs is that organic levels exceed those of sulfate (except 30 January in MBL), regardless of whether the data were in the marine boundary layer (i.e., BBL and 31 BCB legs) or free troposphere (i.e., ACT and ABL legs) (Figure S1b). Hegg et al. (1997) concluded for the month of June based on a chemical apportionment study using aerosol column optical depth 32 33 data off the mid Atlantic coast Of the United States that the three most abundant components (in 34 decreasing order) were water, carbonaceous compounds, and then sulfate. This is an important 35 result with implications for aerosol characteristics such as hygroscopicity. For instance, higher organic:sulfate mass ratios in the marine boundary layer correspond to suppressed hygroscopic 36 37 growth factors at high relative humidities (> 85%) (Hersey et al., 2009). For comparison, airborne measurements in winter and summer periods over the eastern North Atlantic showed sulfate 38 concentrations exceeding those of organics up to the same altitudes (~1.6 km) in this study (Wang 39 et al., 2022). 40

42 Table S1. Average concentrations of submicrometer aerosol species measured by

43 an airborne AMS for different seasons associated with ACTIVATE deployments 1-4. Also

44 shown are mass fractions of each AMS species relative to total AMS mass, in addition to the

45 ratio of m/z 44 to total organic mass (i.e., f44). Non-CAO and CAO categories include samples

46 <u>collected between January and March. CVI = droplet residual particle measurements in</u>
 47 cloud; BCB = below cloud base, ACT = above cloud top, BBL = below boundary layer top,

47 <u>cloud; BCB = below cloud base, ACT = above cloud top, BBL = below boundary layer top,</u>
 48 <u>ABL = above boundary layer top. Corresponding standard deviations and number of points</u>

49 are provided in Table S2.

		(Non-CAO/CAO/May-Jun/Aug-Sep)			
	CVI	BCB	ACT	BBL	ABL
Organic (µg m ⁻³)	-	1.07/0.67/1.49/3.27	0.61/0.19/2.62/3.04	2.59/1.16/3.49/4.46	0.94/0.57/5.28/5
Sulfate (µg m ⁻³)	-	0.93/0.79/1.71/1.35	0.53/0.26/1.23/1.11	0.80/0.57/1.17/1.77	0.51/0.45/1.26/2.1
Nitrate (µg m ⁻³)	-	0.40/0.21/0.07/0.16	0.19/0.05/0.14/0.11	0.79/0.93/0.17/0.21	0.14/0.32/0.26/0.1
Ammonium (µg m ⁻³)	-	0.45/0.32/0.36/0.36	0.28/0.10/0.41/0.37	0.67/0.65/0.38/0.53	0.26/0.30/0.51/0.6
Chloride (µg m ⁻³)	-	0.03/0.02/0.03/0.03	0.02/0.01/0.02/0.02	0.05/0.01/0.02/0.02	0.01/0.01/0.02/0.0
Organic _{MF}	0.55/0.60/0.68/0.61	0.40/0.34/0.35/0.48	0.28/0.29/0.42/0.51	0.50/0.39/0.63/0.57	0.44/0.32/0.65/0.5
Sulfate _{MF}	0.24/0.19/0.14/0.14	0.39/0.45/0.53/0.39	0.42/0.46/0.43/0.34	0.24/0.20/0.26/0.33	0.35/0.36/0.24/0.3
Nitrate _{MF}	0.05/0.05/0.05/0.05	0.08/0.07/0.02/0.03	0.08/0.07/0.03/0.03	0.11/0.22/0.03/0.03	0.06/0.14/0.03/0.0
Ammonium _{MF}	0.09/0.08/0.07/0.09	0.13/0.13/0.10/0.08	0.20/0.16/0.12/0.10	0.14/0.18/0.08/0.07	0.14/0.16/0.07/0.0
Chloride _{MF}	0.06/0.08/0.06/0.10	0.01/0.01/0.01/0.01	0.01/0.02/0.01/0.01	0.01/0.01/0.01/0.00	0.01/0.03/0/00/0.0
f_{44}	0.33/0.34/0.24/0.37	0.15/0.13/0.11/0.14	0.26/0.16/0.12/0.15	0.16/0.14/0.12/0.14	0.17/0.14/0.11/0.1
A			·	·	

52 <u>Table S2.</u> Standard deviations and the number of points for parameters measured by the

53 AMS instrument. Note that numbers of points refer to entire legs for which calculations

54 were conducted using raw data rather than the raw data points per leg. Non-CAO and

55 CAO categories include samples collected between January and March. CVI = droplet

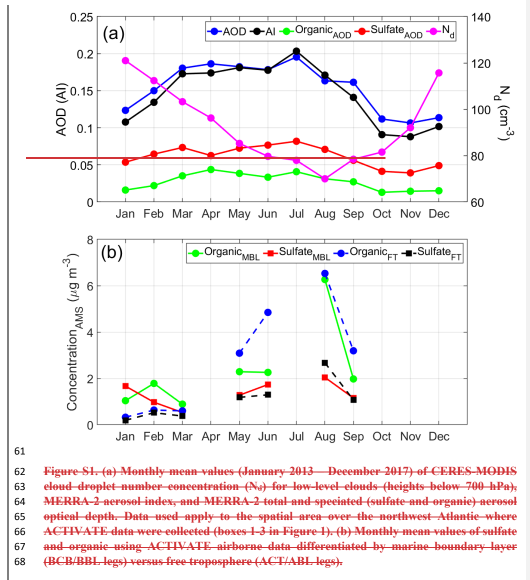
56 residual particle measurements in cloud; BCB = below cloud base, ACT = above cloud top,

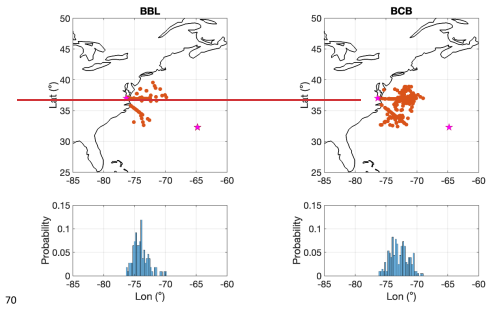
57 BBL = below boundary layer top, ABL = above boundary layer top. Corresponding

58 average values are provided in Table <u>1 of the manuscriptS1</u>.

		(Non-CAO/CAO/May-Jun/Aug-Sep)						
		CVI	BCB	ACT	BBL	ABL		
	Organic (µg m ⁻³)	-	0.63/0.43/1.84/3.93	1.06/0.14/5.08/4.50	2.13/0.69/2.58/3.25	1.01/0.57/3.92/5.86		
	Sulfate (µg m ⁻³)	-	0.52/0.50/0.68/0.83	0.45/0.16/1.04/1.30	0.38/0.35/0.53/0.82	0.27/0.42/0.71/2.10		
	Nitrate (µg m ⁻³)	-	0.82/0.44/0.07/0.16	0.37/0.07/0.27/0.15	1.05/0.82/0.19/0.18	0.25/0.48/0.25/0.18		
	Ammonium (µg m ⁻³)	-	0.52/0.33/0.25/0.37	0.32/0.08/0.50/0.45	0.58/0.50/0.22/0.38	0.21/0.36/0.35/0.67		
	Chloride (µg m ⁻³)	-	0.03/0.02/0.02/0.03	0.05/0.01/0.02/0.03	0.06/0.01/0.02/0.02	0.01/0.01/0.01/0.02		
	Organic _{MF}	0.15/0.16/0.18/0.20	0.14/0.11/0.21/0.23	0.21/0.15/0.21/0.21	0.16/0.09/0.15/0.16	0.17/0.15/0.18/0.23		
	Sulfate _{MF}	0.12/0.14/0.13/0.12	0.15/0.15/0.21/0.22	0.18/0.16/0.18/0.19	0.15/0.07/0.16/0.18	0.15/0.16/0.17/0.22		
	Nitrate _{MF}	0.05/0.06/0.06/0.05	0.10/0.08/0.01/0.02	0.09/0.07/0.03/0.03	0.08/0.11/0.02/0.02	0.04/0.09/0.03/0.02		
	Ammonium _{MF}	0.10/0.09/0.10/0.14	0.06/0.06/0.05/0.07	0.10/0.08/0.08/0.09	0.05/0.04/0.04/0.04	0.08/0.06/0.04/0.05		
	Chloride _{MF}	0.08/0.08/0.08/0.14	0.01/0.01/0.01/0.01	0.02/0.01/0.01/0.02	0.01/0.01/0.01/0.01	0.01/0.05/0.00/0.00		
	f_{44}	0.14/0.21/0.51/0.59	0.12/0.04/0.07/0.11	0.19/0.11/0.09/0.11	0.02/0.04/0.03/0.04	0.09/0.03/0.03/0.08		
59	no. points	180/96/386/228	32/21/70/41	31/22/67/41	24/8/45/27	24/12/97/53		

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71 Figure S2. Midpoint locations of both below bounday layer top (BBL) legs in cloud-free

72 ensembles and below cloud base (BCB) legs in cloudy ensembles during ACTIVATE's

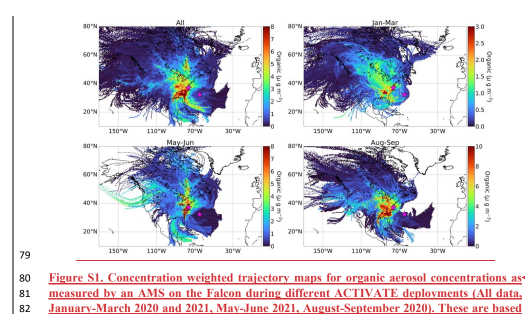
73 deployments 1-4 in 2020 and 2021. The bottom panels show probability histograms of the

74 location of the two leg types relative to longitude. The analagous results for above

75 boundary layer top (ABL) and above cloud top (ACT) legs resemble these since the ABL

76 and ACT legs occur fairly soon after BBL and BCB legs, respectively, within an ensemble.

	(Non-CAO/CAO/May-Jun/Aug-Sep)					Field Code Changed
	CVI	BCB	ACT	BBL	ABL	_
Organic (µg m ⁻³)	-	0.63/0.43/1.84/3.93	1.06/0.14/5.08/4.50	2.13/0.69/2.58/3.25	1.01/0.57/3.92/5.86	5
Sulfate (µg m ⁻³)	-	0.52/0.50/0.68/0.83	0.45/0.16/1.04/1.30	0.38/0.35/0.53/0.82	0.27/0.42/0.71/2.10	
Nitrate (µg m ⁻³)	-	0.82/0.44/0.07/0.16	0.37/0.07/0.27/0.15	1.05/0.82/0.19/0.18	0.25/0.48/0.25/0.48	3
Ammonium (µg m ⁻³)	-	0.52/0.33/0.25/0.37	0.32/0.08/0.50/0.45	0.58/0.50/0.22/0.38	0.21/0.36/0.35/0 67	7
Chloride (µg m ⁻³)	-	0.03/0.02/0.02/0.03	0.05/0.01/0.02/0.03	0.06/0.01/0.02/0.02	0.01/0.01/0.01/0.02	2
Organic _{MF}	0.15/0.16/0.18/0.20	0.14/0.11/0.21/0.23	0.21/0.15/0.21/0.21	0.16/0.09/0.15/0.16	0.17/0.15/0.18/0.23	3
Sulfate _{MF}	0.12/0.14/0.13/0.12	0.15/0.15/0.21/0.22	0.18/0.16/0.18/0.19	0.15/0.07/0.16/0.18	0.15/0.16/0.17/0.22	2
Nitrate _{MF}	0.05/0.06/0.06/0.05	0.10/0.08/0.01/0.02	0.09/0.07/0.03/0.03	0.08/0.11/0.02/0.02	0.04/0.09/0.03 0.02	2
Ammonium _{MF}	0.10/0.09/0.10/0.14	0.06/0.06/0.05/0.07	0.10/0.08/0.08/0.09	0.05/0.04/0.04/0.04	0.08/0.06/0.04/0.05	5
Chloride _{MF}	0.08/0.08/0.08/0.14	0.01/0.01/0.01/0.01	0.02/0.01/0.01/0.02	0.01/0.01/0.01/0.01	0.01/0.05/0.00/0.00	
f_{44}	0.14/0.21/0.51/0.59	0.12/0.04/0.07/0.11	0.19/0.11/0.09/0.11	0.02/0.04/0.03/0.04	0.09/0.03/0.03/0.08	3
no. points	180/96/386/228	32/21/70/41	31/22/67/41	24/8/45/27	24/12/97/53	



on 29,164 cloud-free AMS data points. The pink stars represent NASA Langley Research

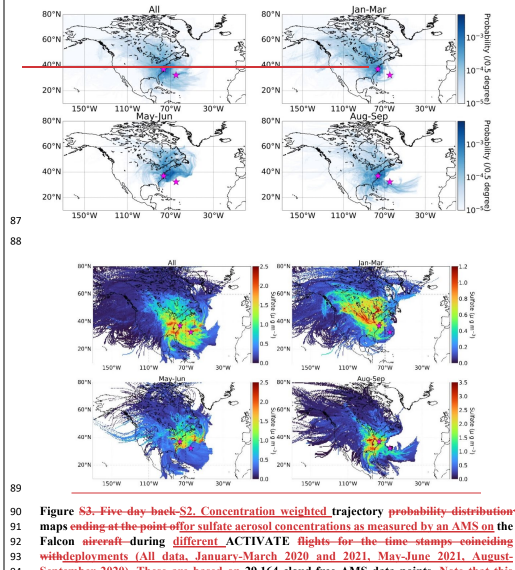
Center (Hampton, Virginia) and Bermuda for reference. Color bar scales differ to show

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variability better within a given panel.

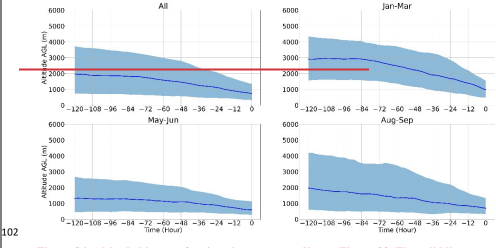


September 2020). These are based on 29,164 cloud-free AMS data points. Note that this includes data during cloud ensembles but only when cloud liquid water content was < 0.05 g m⁻³, and thus data during BCB and ACT legs are included. "All" shows the cumulative results of the other three panels. The January March panel combines CAO and non-CAO

98 days, which are separated for other parts of the study. The pink stars represent NASA

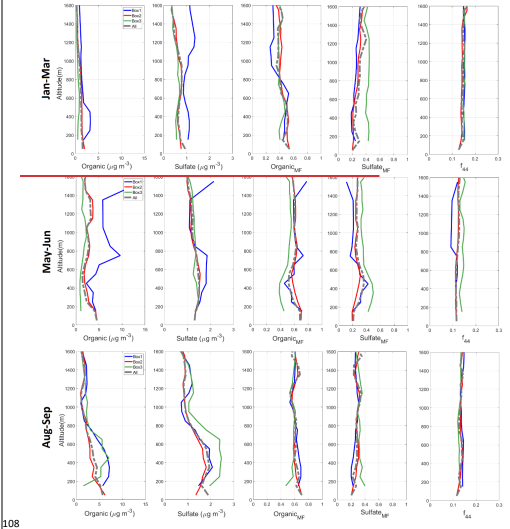
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- Langley Research Center (Hampton, Virginia) and Bermuda for reference. <u>Color bar scales</u> <u>differ to show variability better within a given panel.</u>
- 100

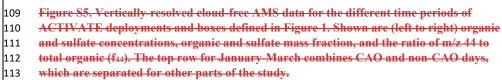


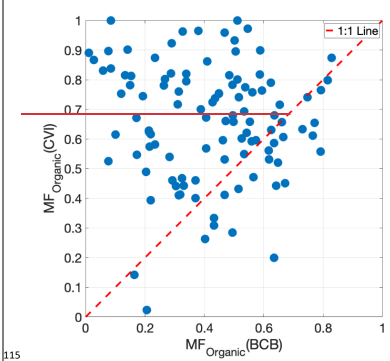
103 Figure S4. Altitude history of trajctories corresponding to Figure S3. The solid line

- represents the median and the shading corresponds to the 25th/75th percentiles. "All" shows
 the cumulative results of the other three panels. The January-March panel combines CAO
 and non-CAO days, which are separated for other parts of the study.
- 107







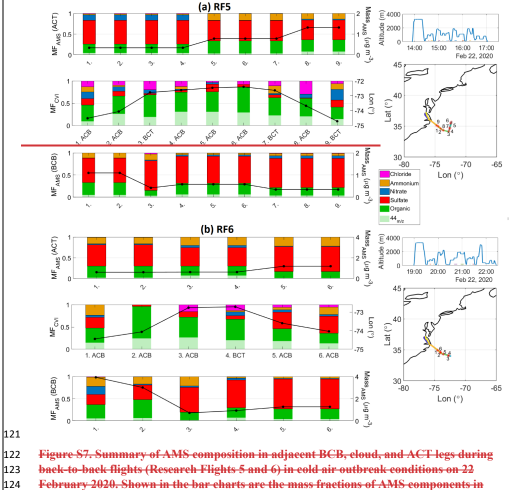






117 cloud) and in acrosol sampled during the closest below cloud base (BCB) leg from

- 118 119 ACTIVATE deployments 1-4. A total of 25 points out of a total of 110 (23%) were below
- the 1:1 line.



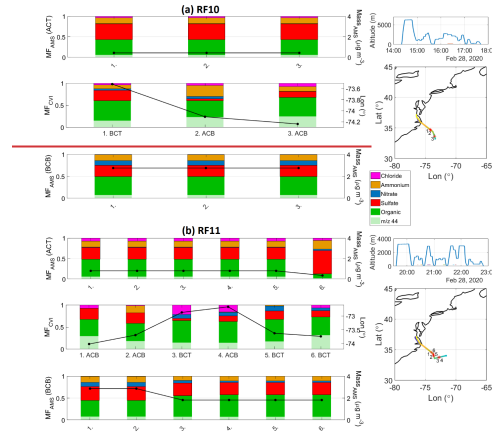
125 addition to either total AMS mass (for ACT and BCB legs; such data are not robust for

126 CVI legs due to how the CVI operates) or longitude on the right y-axis. Note that some

127 BCB and ACT legs are repeated for different cloud legs as they represent the closest leg to

128 an individual cloud leg. On the far right are flight altitude time series along with the spatial

129 map with numbers corresponding to the leg numbers in the bar charts.



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Figure S8. Summary of AMS composition in adjacent BCB, cloud, and ACT legs during 132 133 back-to-back flights (Research Flights 10 and 11) in cold air outbreak conditions on 28 February 2020. Shown in the bar charts are the mass fractions of AMS components in 134 addition to either total AMS mass (for ACT and BCB legs; such data are not robust for 135 136 CVI legs due to how the CVI operates) or longitude on the right y-axis. Note that some 137 BCB and ACT legs are repeated for different cloud legs as they represent the closest leg to 138 an individual cloud leg. On the far right are flight altitude time series along with the spatial 139 map with numbers corresponding to the leg numbers in the bar charts.

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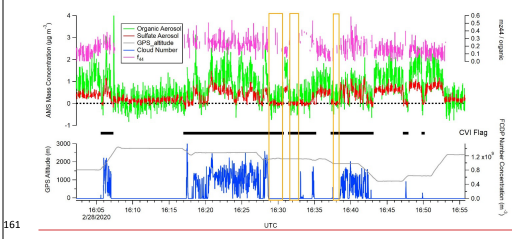
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¹⁶²

- 163 Figure S3. Time series from research flight 10 (28 February 2020) of (top) AMS mass
- 164 concentrations of sulfate and organic in addition to f44, and (bottom) Falcon altitude and
- 165 FCDP droplet number concentration. The CVI Flag horizontal bar indicators represent
- 166 when aerosol sampling was conducted downstream of the CVI, with the vertical orange

167	bars indicating specifically those CVI times where sampling was out of cloud (i.e., data	
168	serving as blanks).	Formatted: Font color: Auto
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