Characterization of Aerosol Hygroscopicity Over the Northeast Pacific Ocean: Impacts on Prediction of CCN and Stratocumulus Cloud Droplet Number Concentrations

B. C. Schulze1, S. M. Charan2, C. M. Kenseth2, W. Kong2, K. H. Bates3, W. Williams4, A. R. Metcalf4, H. H. Jonsson5, R. Woods5, A. Sorooshian6,7, R. C. Flagan8,2, and J. H. Seinfeld8,2

1Department of Environmental Science and Engineering, California Institute of Technology, Pasadena, CA, USA, 2Division of Chemistry and Chemical Engineering, California Institute of Technology, Pasadena, CA, USA, 3Center for the Environment, Harvard University, Cambridge, MA, USA, 4Department of Environmental Engineering and Earth Sciences, Clemson University, Anderson, SC, USA, 5Naval Postgraduate School, Monterey, CA, USA, 6Department of Chemical and Environmental Engineering, University of Arizona, Tucson, AZ, USA, 7Department of Hydrology and Atmospheric Sciences, University of Arizona, Tucson, AZ, USA, 8Division of Engineering and Applied Science, California Institute of Technology, Pasadena, CA, USA

Abstract During the Marine Aerosol Cloud and Wildfire Study (MACAWS) in June and July of 2018, aerosol composition and cloud condensation nuclei (CCN) properties were measured over the N.E. Pacific to characterize the influence of aerosol hygroscopicity on predictions of ambient CCN and stratocumulus cloud droplet number concentrations (CDNC). Three vertical regions were characterized, corresponding to the marine boundary layer (MBL), an above-cloud organic aerosol layer (AC-OAL), and the free troposphere (FT) above the AC-OAL. The aerosol hygroscopicity parameter (κ) was calculated from CCN measurements (κ_{CCN}) and bulk aerosol mass spectrometer (AMS) measurements (κ_{AMS}). Within the MBL, measured hygroscopicities varied between values typical of both continental environments (~0.2) and remote marine locations (~0.7). For most flights, CCN closure was achieved within 20% in the MBL. For five of the seven flights, assuming a constant aerosol size distribution produced similar or better CCN closure than assuming a constant “marine” hygroscopicity (κ = 0.72). An aerosol-cloud parcel model was used to characterize the sensitivity of predicted stratocumulus CDNC to aerosol hygroscopicity, size distribution properties, and updraft velocity. Average CDNC sensitivity to accumulation mode aerosol hygroscopicity is 39% as large as the sensitivity to the geometric median diameter in this environment. Simulations suggest CDNC sensitivity to hygroscopicity is largest in marine stratocumulus with low updraft velocities (<0.2 m s\(^{-1}\)), where accumulation mode particles are most relevant to CDNC, and in marine stratocumulus or cumulus with large updraft velocities (>0.6 m s\(^{-1}\)), where hygroscopic properties of the Aitken mode dominate hygroscopicity sensitivity.

1. Introduction

Marine stratocumulus (MSc) clouds, commonly observed off the Western coasts of North America, South America, Africa, and Australia, cover nearly one fifth of the Earth’s surface and exert a large impact on its radiative balance (Wood, 2012). These cloud decks are particularly relevant to global climate due to their high albedo contrast with the underlying ocean and relatively low altitude, resulting in stronger shortwave reflectance than longwave absorption (Brenguier et al., 2000; Randall et al., 1984; Wood, 2012). Previous estimates suggest that a ~12% increase in the albedo of these clouds would produce a negative radiative forcing equivalent in magnitude to that of doubling atmospheric CO\(_2\) concentrations (Latham et al., 2008; Stevens & Brenguier, 2009). Remote sensing, parcel modeling, and large eddy simulation (LES) studies have all established that MSc exhibit substantial albedo susceptibility to variations in cloud droplet number concentrations (CDNC) (Berner et al., 2015; Chen et al., 2011; Oreopoulos & Platnick, 2008; Platnick & Twomey, 1994; Sanchez et al., 2016). Understanding the sensitivity of MSc CDNC to aerosols acting as
cloud condensation nuclei (CCN) is therefore a critical aspect of reducing uncertainty in climate change predictions (Seinfeld et al., 2016).

The CDNC and albedo of MSc are substantially influenced by the abundance of below-cloud CCN. A recent satellite analysis suggested that variability in below-cloud CCN concentration may be responsible for ~45% of the variability in the radiative effect of marine boundary layer clouds (Rosenfeld et al., 2019). This influence results from the fact that increased CCN abundance enhances cloud reflectivity at constant liquid water path (Twomey, 1977) and has the potential to reduce MSc precipitation rates, increasing cloud lifetime (Ackerman et al., 1993; Albrecht, 1989; Goren & Rosenfeld, 2012; Rosenfeld, 2006). As a result, a major component of the uncertainty in the estimated indirect aerosol forcing has been attributed to the prediction of below-cloud CCN concentrations (Rosenfeld et al., 2014; Sotiropoulou et al., 2007). While the aerosol size distribution is generally thought to be the most important determinant of CCN activity (e.g., Dusek et al., 2006; Ervens et al., 2007; McFiggans et al., 2006; Reutter et al., 2009), particle composition has also been shown to exert a substantial influence (Jimenez et al., 2009; Liu & Wang, 2010; Mei et al., 2013; Quinn et al., 2008; Sanchez et al., 2016).

The propensity of a given aerosol particle to act as a CCN can be described using Köhler theory (Köhler, 1936; Seinfeld et al., 2016), provided sufficient information is known regarding particle size and solute properties (e.g., molecular weight, solubility, density, and activity). A novel framework, κ Köhler theory, condenses these solute characteristics into a single parameter κ (the aerosol hygroscopicity) that can be easily incorporated into large-scale models (Petters & Kreidenweis, 2007). Substantial effort has, therefore, been devoted to quantifying κ values in a multitude of environments (Ervens et al., 2010; Gunthe et al., 2009; Pringle et al., 2010; Rose et al., 2010; Thalman et al., 2017). While κ values characteristic of inorganic aerosol components are relatively well-established, atmospheric organic aerosol is composed of numerous, highly diverse organic compounds, complicating representation of organic hygroscopicity using a single parameter (Kanakidou et al., 2005). Experimental studies have characterized κ values of secondary organic aerosol (SOA) (e.g., Asa-Awuku et al., 2010; Duplissy et al., 2008, 2011; Frosch et al., 2013; Lambe et al., 2011; Massoli et al., 2010; Zhao et al., 2015), and field studies have characterized the typical range of organic κ values (κorg) observed in the atmosphere (Chang et al., 2010; Gunthe et al., 2009; Levin et al., 2014; Mei et al., 2013; Thalman et al., 2017; Wang et al., 2008). Generally, ambient κorg values are found to be 0.1–0.2 for aged aerosol and primary marine organics and ~0 for freshly emitted combustion aerosol (e.g., soot) (Kreidenweis & Asa-Awuku, 2014). A linear relationship has been noted between observed κorg values and organic aerosol oxygen-to-carbon (O:C) ratios in both the laboratory and the field (Chang et al., 2010; Lambe et al., 2011; Mei et al., 2013; Wang et al., 2019).

Ambient particle hygroscopicity data have been combined with aerosol size distribution measurements in CCN closure studies to assess the extent to which Köhler theory can be used to predict ambient CCN concentrations (e.g., Almeida et al., 2014; Asa-Awuku et al., 2011; Cubison et al., 2008; Medina et al., 2007; McFiggans et al., 2006; Moore et al., 2012; Ren et al., 2018; VanReken et al., 2003). Analyzing the accuracy of predicted CCN concentrations can provide insight into the influence of specific aerosol characteristics on CCN activity (Bougiatioti et al., 2011; Cubison et al., 2008; Medina et al., 2007; VanReken et al., 2003; Wang et al., 2010). For instance, size-resolved compositional (i.e., hygroscopicity) data are often required to accurately reproduce observed CCN concentrations in locations dominated by organic aerosol (Bhattu & Tripathi, 2015; Medina et al., 2007; Ren et al., 2018), while aerosol mixing state has been shown to strongly impact total CCN concentrations in urban environments (Cubison et al., 2008; Ervens et al., 2010; Quinn et al., 2008). By analyzing data from five ambient measurement campaigns, Ervens et al. (2010) found that for aerosol measured farther than a few tens of kilometers from the emission source, CCN activity could be predicted within a factor of two independent of either aerosol mixing state (i.e., internal or external) or organic solubility (i.e., insoluble or slightly soluble). Wang et al. (2010) further demonstrated that CCN concentrations can often be reproduced within 20% assuming internal mixing of aerosol components if the overall κ of the aerosol population is >0.1. The direct impact of variability in aerosol hygroscopicity on CCN concentrations is often assessed by assuming an invariant chemical composition, represented as a fixed κ, in CCN closure analyses. Field campaigns in continental environments ranging from polluted megacities to the pristine tropical rainforest have shown that CCN concentrations could be reproduced within 20% and 50%, respectively, assuming a constant κ = 0.3 (Gunthe et al., 2009; Rose et al., 2010), a value representative of average continental conditions (Andreae & Rosenfeld, 2008; Pringle et al., 2010). However, in coastal regions, MBL aerosol can result from a mixture of
distinct marine and continental emissions (e.g., Coggon et al., 2014; Mardi et al., 2018; Modini et al., 2015; Sorooshian et al., 2009), which complicates aerosol representation using regional or global models. CCN closure analysis can provide insight into the uncertainties in CCN concentrations that may result from inaccurate model representation of aerosol composition in these environments.

Due to the importance of the persistent stratocumulus cloud decks over the N.E. Pacific to global climate, aerosol characteristics in this region have received considerable attention. However, the diverse range of particle sources, including shipping exhaust (Coggon et al., 2012; Murphy et al., 2009; Prabhakar et al., 2014; Wonaschütz et al., 2013), primary and secondary natural marine emissions (Modini et al., 2015; Prabhakar et al., 2014; Sorooshian et al., 2009), anthropogenic and biogenic continental emissions (Coggon et al., 2014; Hegg et al., 2010; Moore et al., 2012), wildfire plumes (Brioude et al., 2009; Mardi et al., 2018), and aged aerosol from the Asian continent (Roberts et al., 2006, 2010), combined with strong temporal and spatial variability due to variable meteorological conditions, has hindered determination of general characteristics of the marine atmosphere in this location. This complexity is reflected in the diversity of hygroscopicity measurements previously reported in the marine boundary layer (MBL) and free troposphere (FT). For instance, average $\kappa$ values reported from MBL measurements have varied from ~0.2–0.3 (Moore et al., 2012; Roberts et al., 2010) to ~0.5–0.7 (Royalty et al., 2017; Yakobi-Hancock et al., 2014). Measurements in the FT, while sparse, have been even more variable ($\kappa$ ~0.05–1.0) (Roberts et al., 2006, 2010). While these measurements could largely be reconciled assuming various mixtures of continental (0.27 ± 0.2) and marine (0.72 ± 0.2) aerosol, determining the major emissions sources and meteorological patterns dictating these changes is important for improving model representation of the region (Pringle et al., 2010). CCN-based measurements of aerosol hygroscopicity and the resulting information about small particle composition can be especially useful in this regard, as knowledge of small particle composition can provide substantial insight into particle sources.

While hygroscopicity and mixing state characterization are important components of understanding the CCN activity of ambient aerosol, the dynamic processes controlling supersaturation, droplet nucleation, and droplet growth within clouds lead to nonlinear relationships between aerosol properties and CDNC. As a result, aerosol-cloud parcel modeling is instrumental to fully understand the role of aerosol hygroscopicity and mixing state on CDNC. Reutter et al. (2009) used such a model to distinguish three regimes of aerosol activation, defined as the aerosol-limited, updraft-limited, and transitional regimes, based on the ratio of updraft velocity to aerosol number concentration at the cloud base. The dependence of CDNC on aerosol hygroscopicity, while limited relative to other parameters such as particle number concentration and updraft velocity, was found to vary substantially between regimes. Additional modeling revealed that CDNC sensitivity to aerosol hygroscopicity is highly dependent on the below-cloud aerosol size distribution, with sensitivity increasing substantially with smaller median radii (Ward et al., 2010). Sanchez et al. (2016) concluded that modeled stratocumulus albedo is insensitive to the assumed hygroscopicity of the organic aerosol fraction; however, the sensitivity of CDNC to bulk hygroscopicity has yet to be fully evaluated in this environment.

The present study uses measurements of aerosol composition and CCN activity collected during the Marine Aerosol Cloud and Wildfire Study (MACAWS), combined with an aerosol-cloud parcel model, to gain insight into near-coastal aerosol hygroscopicity and its influence on prediction of CCN and MSc CDNC. Hygroscopicity measurements are combined with airmass backward trajectories and meteorological parameters to attribute observed particle characteristics to distinct sources when possible. CCN closure analyses are performed to investigate the impact of compositional and mixing state assumptions on CCN predictions. Finally, aerosol-cloud parcel model simulations constrained with MSc microphysical measurements are used to directly investigate the sensitivity of stratocumulus CDNC to aerosol hygroscopicity, mixing state, and size distribution properties.

### 2. Methodology

#### 2.1. MACAWS Field Mission

The 2018 Marine Aerosol Cloud and Wildfire Study (MACAWS) consisted of 16 research flights operated out of the Center for Interdisciplinary Remotely-Piloted Aircraft Studies (CIRPAS) in Marina, California, during June and July. Measurements were performed on-board the CIRPAS Navy Twin Otter aircraft.
The scientific objectives of individual flights included characterization of marine aerosols and clouds, sampling of shipping vessel exhaust plumes, and investigation of nearby wildfire emissions. The present study focuses on seven research flights primarily aimed at characterization of the relationship between marine aerosol and the overlying stratocumulus cloud deck. Paths of these seven flights are depicted in Figure 1. Flight strategies typically involved a series of level legs at varying altitudes within the MBL and overlying FT. Slant or spiral soundings were generally performed before and after a series of level legs.

2.2. Twin Otter Instrumentation

The navigational and meteorological instrumentation utilized by the Twin Otter aircraft is described in detail by Sorooshian et al. (2018). Ambient aerosol was sampled using a forward-facing sub-isokinetic inlet (Hegg et al., 2005). Aerosol and cloud droplet number concentrations were characterized using a variety of instruments, including multiple condensation particle counters (CPC, TSI 3010, $D_p > 10$ nm; ultrafine CPC, TSI UFCPC, $D_p > 3$ nm), a passive cavity aerosol spectrometer probe (PCASP, $D_p \sim 0.11–3.4$ μm), and forward scattering spectrometer probe (FSSP, Particle Measuring Systems [PMS], $D_p \sim 1.6–45$ μm). Cloud liquid water content was measured using a PVM-100A probe (Gerber et al., 1994), and a threshold value of 0.02 g m$^{-3}$ was used to distinguish in-cloud sampling (Dadashazar et al., 2018; MacDonald et al., 2018).

Cloud condensation nuclei (CCN) number concentrations were measured at four supersaturations (SS) (0.1%, 0.3%, 0.43%, and 0.57%) using a Droplet Measurement Technologies (DMT) dual-column streamwise thermal-gradient cloud condensation nuclei counter (CCNC) (Lance et al., 2006; Roberts & Nenes, 2005). The CCNC operates by applying a linear temperature gradient to a cylindrical sampling tube with continuously wetted walls. As the thermal diffusivity of water vapor exceeds the diffusivity of air, supersaturated conditions are produced along the sampling column centerline. For this study, activated droplets grown to sizes larger than 0.75-μm diameter were counted and sized by an optical particle counter. The sheath and sample flows of each column were maintained at 0.45 and 0.05 L min$^{-1}$, respectively. Instrument pressure was maintained at 750 mb using a flow orifice and active pressure control system at the instrument inlet. Each column of the CCNC was calibrated using ammonium sulfate particles following standard methods as described in Rose et al. (2008). Calibrations were performed before and after the campaign, and observed
deviations in applied SS for a given temperature gradient imply uncertainties of ~6%, similar to the 5% value typical of field campaigns, as reported by Rose et al. (2008).

Aerosol size distributions and number concentrations for \( D_p \) between ~15 and 800 nm were measured with a custom-built scanning mobility particle sizer (SMPS) consisting of a differential mobility analyzer (DMA, TSI 3081) coupled to a condensation particle counter (TSI 3010). The DMA is operated in a closed-system configuration with a recirculating sheath and excess flow of 2.67 L min\(^{-1}\) and an aerosol flow of 0.515 L min\(^{-1}\). The column voltage was scanned from 15 to 9,850 V over a ~2-min interval.

Aerosol chemical composition was measured using a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS, Aerodyne Research Inc., hereafter referred to AMS) (DeCarlo et al., 2006). Incoming air enters the AMS through a 100-μm critical orifice, after which an aerodynamic lens produces a particle beam that is accelerated under high vacuum. The particle beam is flash-vaporized on a resistively heated surface (600°C), and the resulting gases are ionized by electron impact (70 eV). Individual ion identity is determined using a high-resolution time-of-flight mass spectrometer. Due to the limited amount of aerosol mass present over the MBL, data were collected in high-sensitivity V-mode. The ionization efficiency (IE) of the AMS was calibrated using dry, 350-nm ammonium nitrate particles before each flight. Data were averaged over 1-min intervals, and all data were analyzed using standard AMS software (SQUIRREL v1.57 and PIKA v1.16l) within Igor Pro 6.37. The collection efficiency (CE) was determined using the composition-dependent calculator within the SQUIRREL and PIKA software packages (Middlebrook et al., 2012). Elemental H:C and O:C ratios were calculated using the “Improved-Ambient” elemental analysis method for AMS mass spectra (Canagaratna et al., 2015).

Positive matrix factorization (PMF) analysis (Paatero & Tapper, 1994) was performed on the improved AMS mass spectra in order to distinguish major classes and transformation processes of measured OA. Three factors were extracted, two of which factors correspond to OA subtypes characteristic of the MBL and above-cloud aerosol layer (AC-OAL), respectively, and resemble low-volatility oxygenated organic aerosol (LV-OOA). The third factor, which was rarely observed, is likely a result of primary anthropogenic emissions and resembles hydrocarbon-like organic aerosol (HOA). Further discussion of PMF data preparation and factor interpretation is included in the supporting information.

### 2.3. Determination of Aerosol Hygroscopicity

Aerosol hygroscopicity was calculated using two distinct methods based on measurements with the CCNC and AMS, respectively. Assuming a particle population is internally mixed, the critical activation diameter \( D_{p,c} \) (the diameter at which all larger particles will activate into cloud droplets) produced by a given SS can be determined by integrating the particle size distribution until the total CN concentration is equivalent to the measured CCN concentration:

\[
N_{CCN} = \int_{D_{p,c}}^{\infty} n_{CN} dD_p
\]

Knowledge of the critical diameter can then be used to calculate a single parameter representation of aerosol hygroscopicity from Köhler theory (Petters & Kreidenweis, 2007):

\[
s = \frac{D_{wet}^3 - D_{p,c}^3}{D_{wet}^3 - D_{p,c}^3 (1 - \kappa_{CCN})} \exp \left( \frac{4\sigma M_w}{RT \rho_w D_{wet}} \right)
\]

where \( s \) is the equilibrium supersaturation, \( D_{p,c} \) is the critical activation diameter, \( D_{wet} \) is the droplet diameter, \( R \) is the universal gas constant, \( T \) is the absolute temperature, \( \rho_w \) is the molar density of water, \( M_w \) is the molecular weight of water, and \( \sigma \) is the surface tension of the droplet at the point of activation. Following Rose et al. (2010), \( \kappa \) was determined by applying the observed activation diameter and varying both \( D_{wet} \) and \( \kappa \) until \( s \) is equivalent to the applied supersaturation of the CCNC and the maximum of a Köhler curve of CCN activation. The droplet surface tension is assumed equal to that of water for comparison with other studies (Collins et al., 2013; Petters & Kreidenweis, 2007; Roberts et al., 2010; Yakobi-Hancock et al., 2014). Hygroscopicity values calculated using this method are referred to as “CCN-derived.” Since the likelihood of particle activation at a given SS tends to be a stronger function...
of size than composition (Dusek et al., 2006), $\kappa_{\text{CCN}}$ values correspond to particles with diameters near the calculated critical diameter.

A Monte Carlo approach was used to estimate the uncertainty in CCN-derived kappa values (Wang et al., 2019). A detailed description is provided in the supporting information. For a given measurement of the aerosol size distribution and CCN number concentration, the distribution of possible $\kappa_{\text{CCN}}$ values calculated by varying these input parameters (i.e., CCN number concentration and size distribution) within their respective uncertainties is lognormally distributed. As a result, uncertainties attributed to $\kappa_{\text{CCN}}$ are not symmetric about the geometric mean values. In general, we estimate 1σ uncertainties of $+55%/-40\%$ for $\kappa_{\text{CCN}}$ calculated at SS = 0.3%, $+75\%/−45\%$ at SS = 0.43%, and $+100\%/−50\%$ to values calculated at SS = 0.57%. Due to the low CCN number concentrations observed at SS = 0.1% ($<100$ cm$^{-3}$) and possibility of counting unactivated particles (expected to only be a few per cm$^{-3}$), $\kappa_{\text{CCN}}$ at SS = 0.1% are not reported, as small absolute deviations in particle number concentration measured by the CCNC and DMA due to differential inlet losses could strongly influence the resulting $\kappa_{\text{CCN}}$ estimates.

Hygroscopicity estimates can also be made using component volume fractions measured by the HR-ToF-AMS using the following equation (Petters & Kreidenweis, 2007):

$$\kappa_{\text{AMS}} = \sum_{i} \varepsilon_i \kappa_i$$

where $\varepsilon_i$ and $\kappa_i$ represent the volume fraction and hygroscopicity of the $i$th NR-PM$_1$ component, respectively. While this calculation cannot capture the contribution of refractory components (sea salt, mineral dust, etc.), further analysis suggests their contribution is minor, as discussed in the supporting information. Organic aerosol density was assumed to be 1.4 g cm$^{-3}$ for volume fraction calculations given the remote nature of the environments sampled and the oxidized character of the measured organic aerosol (e.g., O:C ratios of MBL and AC-OAL PMF factors were 0.91 and 0.76, respectively) (Hallquist et al., 2009; Roberts et al., 2010). The hygroscopicity of individual inorganic components is calculated using

$$\kappa_i = \left( \frac{M_w}{\rho_w} \right) \left( \frac{\rho_i}{M_i} \right) v_i$$

where $M_w$ and $\rho_w$ are the molar mass and density of water, respectively, and $M_i$, $\rho_i$, and $v_i$ are the molar mass, density, and van’t Hoff factor of the inorganic component. Inorganic aerosol was dominated by sulfate and ammonium. The relative abundances of ammonium sulfate, ammonium bisulfate, and sulfuric acid were calculated using the molar ratio of ammonium to sulfate (Asa-Awuku et al., 2011; Nenes et al., 1998). Ammonium sulfate and bisulfate were assigned van’t Hoff factors of 2.5, while sulfuric acid was assigned $\kappa = 0.9$ to align with previous measurements (Petters & Kreidenweis, 2007). Modifying the van’t Hoff factors of ammonium sulfate and ammonium bisulfate and assumed $\kappa$ of sulfuric acid within reasonable limits had a negligible influence on the presented results. Chloride measured by the AMS was assumed to represent sodium chloride and was assigned a hygroscopicity of 1.28 (Petters & Kreidenweis, 2007). AMS-measured nitrate aerosol was assumed to be ammonium nitrate with a hygroscopicity of 0.67 (Petters & Kreidenweis, 2007). The hygroscopicity of the organic component ($\kappa_{\text{org}}$) was assumed to be either 0 (non-hygroscopic), 0.1 (slightly-hygroscopic), or a function OA composition using a parameterization based on bulk O:C ratios developed in the literature (Lambe et al., 2011). Comparisons of $\kappa_{\text{CCN}}$ and $\kappa_{\text{AMS}}$ values, analysis of PMF factor composition, and evaluation of CCN-closure calculations are used to evaluate these different $\kappa_{\text{org}}$ estimates.

An uncertainty analysis similar to that described for $\kappa_{\text{CCN}}$ values was performed for $\kappa_{\text{AMS}}$ values and is described in detail in the supporting information. For median conditions in the MBL and FT, the relative uncertainty in $\kappa_{\text{AMS}}$ is estimated to be $−10–20\%$, due primarily to uncertainty in the estimated hygroscopicity of the organic component ($\kappa_{\text{org}}$). In the AC-OAL, the dominant contribution of organic aerosol increases the relative uncertainty to $−50\%$; however, due to the low absolute $\kappa_{\text{AMS}}$ values observed in the AC-OAL, the absolute uncertainty is only $−0.1$ or less.
2.4. Aerosol-Cloud Parcel Model

The aerosol-cloud parcel model used in this study employs a user-specified updraft velocity to induce adiabatic cooling of an air parcel, leading to water vapor supersaturation. The predicted parcel supersaturation at each time step is determined by the relative rates of production through adiabatic cooling and loss through condensation of water vapor onto activated cloud droplets (Pruppacher & Klett, 1997; Seinfeld et al., 2016). In the present study, meteorological parameters such as ambient pressure, temperature, and lapse rate are obtained from MACAWS aircraft measurements and are specified before model execution. The below-cloud dry size distribution is assumed to contain Aitken and accumulation modes, the characteristics of which (i.e., number concentration, geometric mean diameter, hygroscopicity) are set by the user. Particles within each mode can be specified as either internally or externally mixed. Each compositional class, 1 per size mode if internally mixed or 2 per size mode if externally mixed, contains 300 lognormally spaced bins ranging from 1 nm to 3 μm. Droplet activation is assumed to occur when the ambient supersaturation of the parcel exceeds the critical supersaturation of the particles in a given size bin, as determined from κ-Kohler theory (Petters & Kreidenweis, 2007). Following activation, the growth of individual cloud droplet bins due to water vapor diffusion is explicitly represented. Additional physical processes such as droplet coagulation, coalescence, and deposition are not included, as previous parcel model studies have demonstrated that these processes have little influence on model predictions for typical marine stratocumulus conditions (Sanchez et al., 2016). Model execution proceeds until a user-specified liquid water content (0.4 g m⁻³ in this study) has been reached. Activated particle size bins larger than 1 μm are considered cloud droplets; however, using an alternative size threshold of 2 μm or 0.75 μm has a negligible influence on the results.

2.5. Air Mass Backward Trajectories

Air mass backward trajectories (120 hr) were calculated in the MBL for each flight using the NOAA HYSPLIT v4.2 model with the global data assimilation system (GDAS) 1° × 1° meteorological data set (Draxler & Hess, 1997, 1998; Stein et al., 2015). The higher spatial resolution EDAS 40 km × 40 km meteorological data set was not used due to its limited spatial range over the Pacific Ocean. The ending altitude of each trajectory was the approximate midpoint of the MBL during each flight.
Table 1
Median Aerosol Number (N) and Cloud Condensation Nuclei (CCN) Concentrations Measured in the Marine Boundary Layer (MBL), Above-Cloud Organic Aerosol Layer (AC-OAL), and Free Troposphere (FT)

| Location | N (cm$^{-3}$) | CCN: 0.1% (cm$^{-3}$) | CCN: 0.3% (cm$^{-3}$) | CCN: 0.43% (cm$^{-3}$) | CCN: 0.57% (cm$^{-3}$) |
|----------|--------------|------------------------|------------------------|------------------------|------------------------|
| MBL      | 754 (509–978) | 75 (33–106)            | 194 (146–285)          | 302 (187–410)          | 410 (229–522)          |
| AC-OAL   | 1,662 (1,303–1,959) | 58 (41–84)            | 363 (260–537)          | 574 (403–876)          | 781 (539–1,051)        |
| FT       | 333 (296–555)  | 21 (14–35)             | 115 (89–145)           | 144 (102–194)          | 162 (118–240)          |

Note. Values in parentheses represent the interquartile range. CCN concentrations are provided as a function of the instrument supersaturation (%).

3. Results and Discussion

3.1. Aerosol Characteristics Over the N.E. Pacific

Results from the seven flights analyzed in this study are summarized in Figure 2 and Tables 1–3. In the subsequent analyses, “all flights” refers to these seven. Typical flight patterns included sampling within the MBL, FT, and, when present, the above cloud. The AC-OAL is operationally defined as the narrow altitude band (generally <200 m) directly above the marine stratocumulus cloud decks where OA mass loadings were relatively large (>1.5 μg m$^{-3}$) and a distinct AC-OAL PMF factor contributed >80% of total OA mass (Figure S6). This region occupies a similar location as the commonly referenced entrainment interface layer (EIL) above cloud decks (Dadashazar et al., 2018; Wood, 2012), but is defined by the aerosol characteristics described above rather than by turbulence and buoyancy characteristics, as is common for the EIL (Carman et al., 2012). Median aerosol properties are reported in Tables 1–3 for each of these three regions, while Figure 2 displays vertical profiles of aerosol and meteorological properties.

Distinct differences in particle properties were observed within each vertical region. Median aerosol number concentrations observed in the MBL (754 cm$^{-3}$) exceeded those in the FT (333 cm$^{-3}$), as expected. Observed particle concentrations were maximized within the AC-OAL (1,662 cm$^{-3}$), where intense actinic fluxes and elevated concentrations of the hydroxyl radical may drive new particle formation (Dadashazar et al., 2018; Mauldin et al., 1999). For all measured SS > 0.1%, observed CCN concentrations were also largest within the AC-OAL, rather than the MBL or FT, underscoring the importance of understanding the hygroscopicity of above-cloud CCN-active particles (Coggon et al., 2014; Sorooshian, Lu, et al., 2007; Sorooshian et al., 2007; Wang et al., 2008).

Observed aerosol composition in the MBL was relatively evenly divided between organic aerosol (OA) (43%) and sulfate (SO$_4$) (48%), with a minor contribution from ammonium (NH$_4$) (~10%) and negligible nitrate (NO$_3$) (~1%). Prabhakar et al. (2014) have demonstrated that nitrate is preferentially distributed in super-micron particles in this marine environment, in agreement with the minor contribution observed with the AMS in this study. Using the “clean” versus “perturbed” threshold introduced by Coggon et al. (2012) for this region (where “clean” is defined by aerosol mass concentrations <1 μg m$^{-3}$), average MBL conditions were “perturbed” by shipping vessel emissions or other anthropogenic sources such as continental outflow. A distinct, highly oxidized MBL PMF factor was extracted from the data set (Figure S6). The oxidized nature of the MBL factor (O:C = 0.91) precludes the use of marker ions to distinguish individual sources; however, potential sources include shipping and biogenic emissions, as well as oxidized continental outflow aerosol (Coggon et al., 2012; Hegg et al., 2010; Sorooshian et al., 2009). In the AC-OAL, observed aerosol composition was dominated by organics (80%), as has been previously reported (Coggon et al., 2014; Coggon et al., 2012).
Table 3

| Location       | \( \kappa_{\text{AMS}} \) | \( \kappa_{\text{CCN}: 0.3\%} \) | \( \kappa_{\text{CCN}: 0.43\%} \) | \( \kappa_{\text{CCN}: 0.57\%} \) |
|----------------|-----------------|-----------------|-----------------|-----------------|
| MBL            | 0.45 (0.35–0.52) | 0.39 (0.20–0.61) | 0.35 (0.24–0.50) | 0.40 (0.27–0.54) |
| AC-OAL         | 0.19 (0.17–0.25) | 0.13 (0.08–0.20) | 0.19 (0.14–0.25) | 0.17 (0.12–0.27) |
| FT             | 0.37 (0.30–0.43) | 0.32 (0.18–0.65) | 0.50 (0.29–0.88) | 0.37 (0.21–0.72) |

Note: Values in parentheses represent the interquartile range. \( \kappa_{\text{CCN}} \) are provided as a function of the instrument supersaturation (%).

Hersey et al., 2009; Sorooshian et al., 2007; Sorooshian, Ng, et al., 2007; Wang et al., 2008). A second, distinct factor displayed large mass loadings (up to 8 \( \mu g \) m\(^{-3}\)) within the AC-OAL (Figure S6) (O:C = 0.76), and the mass ratio of the AC-OAL to the MBL PMF factor is used as a tracer of AC-OAL entrainment into the MBL, as discussed in section 3.3.2. Possible aerosol production mechanisms in the AC-OAL include oxidation and transport of biogenic volatile organic compounds emitted by forested regions in the northwest United States, cloud droplet evaporation, and oxidation of sparingly soluble organics vented through the stratuscumulus layer (Coggan et al., 2014; Heald et al., 2005; Sorooshian, Lu, et al., 2007). While large eddy simulations (LES) have demonstrated that the altitude of the top of the stratuscumulus cloud deck can undergo diurnal variations of 10–100 m, providing a potential mechanism for AC-OAL aerosol production through droplet evaporation (Chen et al., 2011; Sorooshian, Lu, et al., 2007), the substantially larger mass fraction of organic aerosol in the AC-OAL than the MBL suggests that particle production is primarily a result of continental biogenic sources (Coggan et al., 2014). Observed aerosol mass loadings in the FT were the lowest sampled (1.5 \( \mu g \) m\(^{-3}\)) but agree well with previous aircraft measurements by Wang et al. (2008) off the coast of Pt. Reyes, CA, at a similar time of year (June–July).

3.2. Overview of Observed Aerosol Hygroscopicity

Figure 3 displays median aerosol number size distributions, \( \kappa_{\text{AMS}} \), and \( \kappa_{\text{CCN}} \) values observed within the MBL, AC-OAL, and FT during each flight. For these comparisons, \( \kappa_{\text{AMS}} \) values are calculated assuming \( \kappa_{\text{org}} = 0.1 \), as is typical for non-urban regions (Mei et al., 2013; Moore et al., 2011, 2012). However, we note that using the parameterization developed by Lambe et al. (2011), the calculated \( \kappa_{\text{org}} \) values for the MBL and AC-OAL PMF factors are 0.19 and 0.17, respectively, due to their highly oxidized nature (Figure S6), suggesting the true \( \kappa_{\text{org}} \) values for large particles may be greater than 0.1.

Within the MBL, observed hygroscopicity values appear to cluster into three relatively distinct groups that span the range of values previously observed in this environment (Roberts et al., 2010; Royalty et al., 2017; Yakobi-Hancock et al., 2014). The strong temporal variation observed in both particle number size distributions and hygroscopicities underscores the complexity involved in accurately modeling CCN in coastal environments influenced by continental and marine sources. This is further demonstrated in Table 4, which depicts estimated organic and inorganic volume fractions of Aitken mode particles derived from MBL \( \kappa_{\text{CCN}} \) values. Assuming inorganic aerosol is entirely ammonium sulfate for these calculations, estimated organic fractions vary from effectively zero, as median \( \kappa_{\text{CCN}} \) during RF13 are larger than that of ammonium sulfate (\( \kappa = 0.61 \)) to as high as 84%. The low hygroscopicities and subsequently large estimated organic fractions observed during flights RF9 and RF15 are uncharacteristic of remote marine environments and imply a continental influence on particle characteristics. \( \kappa_{\text{AMS}} \) values calculated during these flights are ~50–100% larger than \( \kappa_{\text{CCN}} \) values, implying addition of particle mass during growth that is more hygroscopic than the Aitken mode particles. While the difference between \( \kappa_{\text{AMS}} \) and \( \kappa_{\text{CCN}} \) values during these flights are nearly within the uncertainty range of the \( \kappa_{\text{CCN}} \) calculation, these observations align with those in many continental locations, where addition of inorganic mass to organic-rich Aitken mode particles growth is thought to lead to a positive relationship between particle hygroscopicity and size (Ervens et al., 2010; Kawana et al., 2016; Levin et al., 2014; Moore et al., 2012; Rose et al., 2011). On the other hand, \( \kappa_{\text{AMS}} \) and \( \kappa_{\text{CCN}} \) are quite similar during the other five flights, with relative deviations on the order of ~25% or less, which is well within the uncertainty of the \( \kappa_{\text{CCN}} \) measurements. A compilation of data reported by Royalty et al. (2017) suggests that minor variation of particle hygroscopicity with size is a common feature of remote marine aerosol, which generally exhibits elevated Aitken mode hygroscopicity. Four individual flights (RF4, RF5, RF13, and RF15) provide
specific insight into the combined roles of aerosol sources and meteorological processes in determining aerosol hygroscopicity in the MBL, and these are discussed in further detail in section 3.3.2.

Within the AC-OAL, observed aerosol hygroscopicity is remarkably similar from flight-to-flight, and little difference is observed between \( \kappa_{\text{CCN}} \) and \( \kappa_{\text{AMS}} \) values. The combination of reduced hygroscopicity (i.e., \( \kappa \sim 0.2 \)) and little variation with particle size suggests that within the AC-OAL, Aitken mode particles are organic-rich and grow through condensation of additional organic vapors, rather than addition of inorganic mass. Even under the assumption that the organic species in Aitken mode AC-OAL particles are entirely

Figure 3. Median aerosol size distributions (a–c) and hygroscopicities (\( \kappa_{\text{CCN}} \) and \( \kappa_{\text{AMS}} \)) (d–f) measured in the marine boundary layer (MBL), above-cloud organic aerosol layer (AC-OAL), and free troposphere (FT), during the seven RFs. \( \kappa_{\text{AMS}} \) values are calculated assuming \( \kappa_{\text{org}} = 0.1 \) and are plotted at the median of the cumulative aerosol volume distribution. Vertical bars represent the interquartile range of hygroscopicity measurements. Previously observed values in the MBL are included for reference in (d), as are typical values for continental and marine environments from Pringle et al. (2010).
Table 4
Calculated Aitken Mode Organic (\(f_{\text{org}}\)) and Inorganic (\(f_{\text{finorg}}\)) Volume Fractions Based on Median \(\kappa_{\text{CCN}}\) Values Derived From CCN Measurements at \(SS = 0.43\%\) for MBL Measurements During Each Flight

| Flight | \(\kappa_{\text{CCN}} - SS = 0.43\%\) | \(f_{\text{org}}\) | \(f_{\text{finorg}}\) | \(f_{\text{org}}\) | \(f_{\text{finorg}}\) |
|--------|---------------------------------|----------------|----------------|----------------|----------------|
| RF4    | 0.41                           | 0.39           | 0.61           | 0.61           | 0.39           |
| RF5    | 0.46                           | 0.29           | 0.71           | 0.55           | 0.45           |
| RF9    | 0.18                           | 0.84           | 0.16           | 0.90           | 0.10           |
| RF12   | 0.50                           | 0.22           | 0.78           | 0.50           | 0.50           |
| RF13   | 0.76                           | –              | –              | 0.18           | 0.82           |
| RF15   | 0.18                           | 0.84           | 0.16           | 0.90           | 0.10           |
| RF16   | 0.28                           | 0.65           | 0.35           | 0.78           | 0.22           |

Note. Values of \(f_{\text{org}}\) and \(f_{\text{finorg}}\) are calculated assuming the inorganic aerosol component is either ammonium sulfate ((NH\(_4\))\(_2\)SO\(_4\)) or sulfuric acid (H\(_2\)SO\(_4\)). Note that the hygroscopicity measured during RF13 cannot be reproduced assuming the inorganic component is entirely (NH\(_4\))\(_2\)SO\(_4\).

Due to the low aerosol number concentrations in the FT, observed \(\kappa_{\text{CCN}}\) values vary widely between flights and exhibit large variability within individual flights. As a result, we hesitate to draw definitive conclusions based on these data. Other than RF13, average \(\kappa_{\text{AMS}}\) values from each flight are near or below 0.4, implying a substantial organic contribution to free tropospheric aerosol. In the absence of continental influence, observation of aerosols of such low hygroscopicity is unexpected, given that particle formation in the upper FT over tropical oceans is driven primarily by sulfuric acid nucleation and growth (Clarke, 1993; Clarke et al., 1998, 1999, 2013). Long range transport of organic aerosol layers from the Asian continent have been noted previously (Roberts et al., 2006, 2010), but estimates of aerosol hygroscopicity in such layers have varied dramatically. For instance, during the CIFEX experiments (Roberts et al., 2006), average \(x\) attributed to aged aerosol layers were only ~0.04, whereas our measurements suggest a more moderate value of ~0.4, while observations by Roberts et al. (2010) indicated a value of 0.93 was more appropriate. While the substantial difference in particle concentrations in the MBL and FT observed during this campaign suggests FT aerosol plays a minor role in dictating MBL CCN activity on average, in remote marine environments, entrainment from the FT is the dominant source of MBL particles (Clarke, 1993; Clarke et al., 1996, 1998, 2013; Raes, 1995), and as such further research into the variability of FT aerosol composition is warranted.

3.3. Observation of Distinct Influences on MBL Particle Characteristics

Observations shown in Figure 3 indicate highly variable flight-averaged hygroscopicities in the MBL, suggesting that temporal variations in regional meteorology and/or particle source strengths can strongly influence CCN characteristics in this environment. Further analysis suggests that in four of the seven flights discussed in this study, specific meteorological patterns and emissions sources influencing particle characteristics can be identified. We discuss these observations to provide insight into the level of physicochemical detail (both in terms of emissions and atmospheric dynamics) required for atmospheric models to simulate MBL CCN concentrations with high fidelity.

3.3.1. Shipping Emissions

Aerosol properties measured during RF4 and RF5 suggest a prominent influence of regional shipping emissions on particle characteristics and hygroscopicity in this environment. During these flights, the dominance of an Aitken mode near ~50–60 nm with much larger concentrations than in the FT suggests relatively...
recent formation from an MBL-based particle source. While such size distributions could hypothetically result from continental outflow (Moore et al., 2012), air mass backward trajectories remained over the ocean and near or within the MBL (<1,000 m) for the previous 5 days (Figure 4). Furthermore, trajectories transited primarily within the major shipping corridor along the coast, as observed for flights “perturbed” by shipping vessel emissions by Coggon et al. (2012), rather than recently arriving from the remote ocean (e.g., RF13). Downward mixing of AC-OAL particles is also ruled out as an Aitken mode particle source during these flights due to the distinctly different hygroscopicities observed in the MBL and AC-OAL (Figure 3). Finally, average wind speeds within the MBL were ~12 m s\(^{-1}\) and ~9 m s\(^{-1}\) during RF4 and RF5, respectively. Modini et al. (2015) previously noted that primary sea spray emissions produced particle concentrations of only 12 cm\(^{-3}\) during periods with similar windspeeds (12 m s\(^{-1}\)) in the same marine environment (equivalent to ~2% of particle number concentrations in the MBL during RF4 and RF5).

Shipping emissions have been previously noted as major contributors to aerosol and cloud properties in the N.E. Pacific environment (Cappa et al., 2014; Coggon et al., 2012; Lack et al., 2011; Murphy et al., 2009). Coggon et al. (2012) demonstrated that 70% of cloud residual particles measured in the California shipping lanes were impacted by nearby shipping emissions. Available compositional data further suggest that shipping emissions could be expected to produce Aitken mode hygroscopicities observed during RF4 and RF5. For instance, Lack et al. (2011) observed an effective \(\kappa\) value of 0.68–0.73 from exhaust produced by a
large (96,500 ton) container vessel, while the smaller Research Vessel Atlantis sampled during the same study produced a value of ~0.2. Hygroscopic growth factor measurements of shipping exhaust emitted by another large (90,000 ton) container vessel by Murphy et al. (2009) suggest an effective $\kappa = 0.1$ – 0.5.

Direct measurements of a large container vessel exhaust plume during RF7 provide further support for the attribution of aerosol characteristics to shipping emissions in RF4 and RF5. As shown in Figure 5, the strong Aitken mode peak in the size distribution measured directly within the plume aligns well with those measured in RF4 and RF5, while the total magnitude of the flight-median size distributions agree well with those measured in the diluted plume more than 20 km downwind. As the plume was relatively narrow directly behind the ship, $\kappa_{CCN}$ values are not available, but $\kappa_{AMS}$ measurements agree well with those in RF4 and RF5 (Figure 5c). However, given the variability in the measured $\kappa$ values of particulate shipping exhaust just discussed, this agreement cannot be viewed as definitive. Ultimately, while the insights provided by the size distributions, backward trajectories, and $\kappa_{AMS}$ values would not be definitive on their own, taken together they support a shipping emission signature on aerosol characteristics during these flights. This influence highlights the importance of accurate physicochemical representation of shipping vessel emissions within the California coastal zone. As an example, the implementation of recent regulations on the sulfur content of shipping fuel within coastal waters of the United States (up to 200 miles off the coast) should increase the organic:inorganic ratio of particulate shipping emissions in major shipping lanes over time (Cappa et al., 2014; Lack et al., 2011). Assuming, as a strictly upper limit estimate, that all Aitken mode particles observed during RF4 and RF5 are derived from shipping vessel emissions,
changing the assumed hygroscopicity of these emissions from the value observed during ambient measurements in this study (~0.4–0.5) to a value of 0.1 (purely organic, partially hygroscopic), would change the CCN concentration at SS = 0.3% by 15–36%.

### 3.3.2. Entrainment From the AC-OAL

The observation of a single, dominant Aitken mode with reduced hygroscopicity during RF15 suggests an influence of the AC-OAL on MBL particle properties. According to Figure 4, the air mass sampled during RF15 had not recently transited over the continent or within the FT, which has previously shown to occasionally contain distinct layers of reduced hygroscopicity aerosol (Roberts et al., 2006, 2010). Clear evidence of entrainment from the AC-OAL is provided in Figure 6, which contrasts size distributions and $\kappa_{\text{AMS}}$ values observed during RF15 and RF4, another flight with a prominent Aitken particle mode and relatively similar backward trajectory. During RF15, the MBL and AC-OAL size distributions are remarkably similar, exhibiting peak diameters at ~55 nm and lacking a larger accumulation mode. Liquid water contents measured within the MBL during RF15 demonstrate a fully developed stratocumulus layer encompassing roughly half of the MBL. $x_{\text{AMS}}$ values vary linearly with altitude from ~0.4 near the ocean surface to ~0.15–0.2 at the top of the cloud layer, aligning with the hypothesis of downward mixing of AC-OAL particles into the MBL. These observations are in stark contrast to those from RF4, where the Aitken mode diameter of the MBL and AC-OAL aerosol differ by ~20–25 nm, and importantly, the Aitken mode diameter in the MBL is smaller...
than the AC-OAL, suggesting a distinct particle source in each location. Finally, as the AC-OAL and MBL PMF factors are clearly distinguished in each flight where the AC-OAL layer was observed, the AC-OAL:MBL PMF factor mass ratio acts as a tracer for entrainment mixing. During RF15, the median AC-OAL:MBL PMF factor mass ratio was 0.81 in the MBL, in contrast to a value of 0.36 measured during RF4 and a median value of 0.42 in all flights other than RF15 where the AC-OAL was observed. The information obtained from the aerosol size distribution (no accumulation mode) and hygroscopicity (similar to the AC-OAL) in the MBL suggests that the distinct AC-OAL signature may result from entrainment following precipitation scavenging of the preexisting MBL aerosol. As typical AC-OAL particle concentrations are ~5 times as large as those in the overlying FT, failure to simulate this layer will result in underprediction of MBL particle concentrations during such distinct precipitation/entrainment events.

3.3.3. Transport From the Remote Pacific Ocean

Hygroscopicity measurements made during RF13 are notably larger than those from the other six flights, indicating a lack of organic aerosol across the particle size distribution. As expected, backward trajectories calculated within the MBL during this flight indicate recent arrival from the remote Pacific Ocean, rather than extended transport through the major shipping lanes along the coast. The boundary layer was substantially compressed (<300 m) and cloud-free during the flight, suggesting ongoing subsidence of free tropospheric air masses (Figure 7a). As new particle formation through sulfuric acid nucleation is known to be a notable source of CCN throughout the marine boundary layer (Clarke, 1993; Clarke et al., 1998, 2013), downwelling and entrainment of such nucleated particles is a possible explanation for the elevated Aitken mode hygroscopicities observed. While low number concentrations in the FT make $\kappa_{CCN}$ estimates less reliable, the values observed in RF13 are relatively similar to those in the MBL, supporting entrainment. While aerosol size distribution measurements in the FT suggest such entrainment was not responsible for increases in Aitken mode particles locally, as concentrations directly above the MBL are substantially lower than those in the MBL, the elevated aerosol concentrations at ~1,000 m suggest entrainment may have produced MBL Aitken mode particles during transport (Figure 7b). Furthermore, the vertical profile of the aerosol size distribution in the FT is consistent with past observations of growth of nucleation-produced Aitken mode particles during large-scale subsidence (Clarke et al., 1999).

Due to the compressed height of the MBL during RF13, the potential contribution of primary sea spray aerosol to MBL particle characteristics is also enhanced. However, using the size distribution fitting technique established by Modini et al. (2015), the calculated concentration of primary sea spray aerosol is only 18 cm$^{-3}$ or ~4% of the average MBL particle concentration during the flight, suggesting sea spray provides at most a minor contribution.
3.4. CCN Closure Analysis

Figure 8 shows CCN closure results for the three sampled environments using six different assumptions regarding aerosol composition and mixing state. Three cases assume internally mixed aerosol components with composition determined by AMS measurements. These cases are differentiated by their assumptions regarding organic aerosol hygroscopicity, with $\kappa_{\text{org}}$ increasing from 0 (first case), to 0.1 (second case), and finally to values predicted from time-varying measured OA O:C ratios according to the relationship developed by Lambe et al. (2011) (third case). The final three cases are similar to the internally mixed cases in their treatment of $\kappa_{\text{org}}$; however, the organic and inorganic aerosol components are assumed to be externally mixed. Bulk aerosol mass loadings were too low to obtain robust estimates of size-resolved composition, precluding more detailed treatment of composition in CCN closure calculations. Closure was assessed in terms of the normalized mean bias ($\text{NMB} = \sum (\text{CCN}_{\text{pred},i} - \text{CCN}_{\text{meas},i}) / \sum \text{CCN}_{\text{meas},i}$), similarly to Asa-Awuku et al. (2011), which provides a representation of the average CCN prediction error observed for each flight. Data for the MBL and FT are shown for individual flights, while data from the AC-OAL are aggregated from all flights where the layer was observed, as fewer size distributions were obtained from the AC-OAL during each flight (and the AC-OAL was not observed at all during three flights).

For the majority of analyzed flights (five out of seven), closure is obtained within 20% using AMS-measured bulk composition and an assumption of either insoluble ($\kappa_{\text{org}} = 0$) or slightly hygroscopic organics ($\kappa_{\text{org}} = 0.1$). While the assumption of insoluble organics disagrees with observed O:C ratios (e.g., the O:C ratio of the MBL PMF factor is 0.85), CCN closure studies often find this assumption is ideal when assuming internal mixing (Chang et al., 2007; Lance et al., 2009; Moore et al., 2011; Wang et al., 2008). The lack of strong dependence on $\kappa_{\text{org}}$ suggests that in non-urban areas, regional models may be able to assign a single value to organic aerosol rather than attempt to dynamically model changes in organic aerosol hygroscopicity with aging (Wang et al., 2008). This is further highlighted by the fact that closure results assuming a constant $\kappa_{\text{org}}$ value (0.1) are generally more accurate than those produced by parameterizing $\kappa_{\text{org}}$ based on the observed O:C ratio (Lambe et al., 2011). As larger aerosols are more likely to have undergone cloud-processing, parameterizing organic hygroscopicity based on bulk measurements of the organic O:C ratio, which is biased by the largest particles, may also overpredict the oxidation state of particles near the
critical diameter of CCN activation. Without size-resolved compositional data, it is difficult to definitively conclude whether the overprediction observed when $\kappa_{\text{org}}$ is parameterized based on the organic O:C ratio is due to such variability with size or is the result of a different relationship between O:C and $\kappa_{\text{org}}$ for organic aerosols in this environment. However, other published parameterizations between O:C and $\kappa_{\text{org}}$ in the literature either agree well with the Lambe parameterization (Chang et al., 2010; Massoli et al., 2010; Thalman et al., 2017) or predict more hygroscopic particles at the same O:C ratio (and as a result would lead to further overprediction if implemented in the CCN closure analysis) (Mei et al., 2013). The overprediction in CCN observed here when incorporating the Lambe parameterization therefore suggests that small particles near the critical activation diameter are less hygroscopic than larger particles that dominate the mass size distribution and thereby dictate AMS-measured composition.

Overall, generally good closure is expected in a semi-remote environment such as the California coastal zone, as previous studies have noted that closure is likely to be achieved within 20% when the bulk aerosol $\kappa$ exceeds 0.1 (Wang et al., 2010). Furthermore, it is expected that aerosol in this coastal environment can be modeled as internally mixed, regardless of its true mixing state, due to the substantial contribution of inorganic constituents and distance from emission sources (Ervens et al., 2010; Fierce et al., 2016; Moore et al., 2012). Fierce et al. (2016) have demonstrated that in semi-remote environments (i.e., non-urban locations), initially externally mixed aerosol becomes internally mixed on a time scale of about 1 day, while the conversion is even faster (on the order of hours) in urban environments, in agreement with the results of Wang et al. (2010). Notable underpredictions (i.e., >20%) of CCN concentrations are produced when assuming externally mixed aerosol with insoluble organics, in agreement with the aged nature of the aerosol in this environment, which should lead to both oxidized organic aerosol and an appreciable amount of internal mixing.

CCN are strongly overpredicted in the MBL during RF9 (37%) and RF15 (57%) when assuming an internal mixture with hygroscopic organics. Aerosol composition during these flights was dominated by organic species in the MBL (59% and 58% of AMS-derived aerosol mass, respectively), indicative of a continental influence on aerosol properties. AMS-derived hygroscopicities are substantially larger than those derived from CCN measurements (Figure 3), suggesting that size-dependent composition may lead to the observed overprediction of CCN concentrations when using bulk AMS measurements of aerosol composition. Comparison of CCN closure results when assuming internal versus external mixing suggests that organic and inorganic components are externally mixed, implying either distinct particle sources or a lack of significant aging prior to measurement. In the case of RF15, this external mixing aligns with the hypothesis of downward mixing from the organic-rich AC-OAL. Figure S7 depicts the CCN closure normalized mean bias resulting from an assumption of internally mixed aerosols with hygroscopic organics as a function of the CCN-derived hygroscopicity. In general, CCN closure error increases rapidly as $\kappa_{\text{CCN}}$ decreases past ~0.25, suggesting that detailed mixing state and/or size resolved compositional information is critical for accurate CCN prediction in this coastal environment during periods of intense organic aerosol intrusion into the MBL. As the aerosol hygroscopicity calculation used in this study relies on an assumption of internal mixing of organic and inorganic aerosol components, it is difficult to determine whether CCN closure error when assuming internal mixing during this flights is a result of externally mixed organic and inorganic aerosol or a result of variable composition with size. Ultimately, as these atypical organic aerosol-dominated marine conditions are the least likely to be accurately reproduced by regional models, further investigation of their frequency, particle characteristics, and resulting impact on cloud properties is warranted.

The analysis presented in Figure 8 implies that for typical conditions in the MBL (5 out of 7 flights in this study), mixing state and organic hygroscopicity have relatively little influence on CCN number concentrations. Additional closure analyses were performed assuming a constant $\kappa$ equivalent to values attributed to average continental ($\kappa = 0.27$) and marine ($\kappa = 0.72$) environments (Pringle et al., 2010) (Figure 9). These results highlight the fact that assuming coastal aerosols have a strictly marine character leads to substantial errors in CCN prediction (>20% for 8 out of 9 flights) even if size distribution parameters are well characterized. Furthermore, for five out of the seven analyzed flights (RF4, RF5, RF9, RF13, RF16), assuming a constant marine $\kappa$ (0.72) results in CCN prediction error similar to or larger than the error produced by assuming a constant aerosol size distribution derived from the median value measured in the MBL during this study. This underscores the importance of capturing organic contributions to coastal MBL aerosol, whether due to continental outflow, downwelling from the AC-OAL, shipping emissions, or marine biota.
Figure 9. Normalized mean bias resulting from additional CCN closure analyses performed on data from each flight. $\kappa_{\text{Cont}}$ and $\kappa_{\text{Marine}}$ refer to analyses assuming a constant $\kappa$ equivalent to values representative of continental (0.27) and marine (0.72) environments (Pringle et al., 2010). The Constant S.D. case assumes a constant aerosol number size distribution equivalent to the median value observed in the MBL during the campaign. Blacked dashed lines indicate closure error of ±20%. Marker size corresponds to the $R^2$ value computed from a linear fit of observed and predicted CCN from each flight. Note the split in the y-axis.

3.5. Sensitivity of Stratocumulus CDNC to Below-Cloud Aerosol Hygroscopicity

In order to directly investigate the sensitivity of N.E. Pacific stratocumulus CDNC to below-cloud aerosol properties, droplet activation was simulated using an aerosol-cloud parcel model constrained with detailed below-cloud aerosol measurements obtained from three cloud sampling passes performed during the campaign. While a number of previous cloud parcel modeling studies have assumed unimodal size distributions (Chen et al., 2016; Reutter et al., 2009; Ward et al., 2010), observed aerosol size distributions over the N.E. Pacific were frequently bimodal (Figure 3). As many current aerosol modules incorporated within global atmospheric chemistry models involve multiple aerosol size modes (Liu & Wang, 2010; Pringle et al., 2010; Rothenberg et al., 2018), we carried out parcel model runs to analyze CDNC sensitivity to properties of the Aitken and accumulation modes separately. Sensitivities were calculated following McFiggans et al. (2006), where $S(X_i) = \delta\ln N_{\text{CDNC}}/\delta\ln X_i$ and $X_i$ is the parameter under investigation. Standard linear regressions of $\ln N_{\text{CDNC}}$ versus $\ln X_i$ were used to determine $S(X_i)$ values, as is convention (Reutter et al., 2009; Sánchez-Gáñita et al., 2017; Ward et al., 2010). Measured aerosol and meteorological properties utilized as model constraints are summarized in Table 5. Sensitivity to hygroscopicity was computed across the range of $\kappa = 0.2$–0.6. Initial results confirmed that for observed MSc updraft velocities ($u = 0.15$–0.3 m s$^{-1}$), below-cloud particle number concentrations (~500–800 cm$^{-3}$), and typical hygroscopicities ($\kappa \sim 0.2$–0.4), properties of the Aitken mode have a minor impact on stratocumulus properties ($S(X_i) < 0.05$), as minimum simulated activation diameters exceed 100 nm. Therefore, Figure 10 depicts the sensitivity of stratocumulus CDNC to properties of the accumulation mode and the simulated updraft velocity.

The average sensitivity of CDNC to aerosol hygroscopicity (0.19), while smaller than the sensitivity to size distribution parameters, is 39% as large as the sensitivity to the geometric mean diameter of the accumulation mode. This agrees with the consensus that particle size distribution properties have a larger influence on CCN concentration than particle composition (Dusek et al., 2006; McFiggans et al., 2006; Reutter et al., 2009), but also suggests accurate hygroscopicity reproduction should be included in future model improvement efforts. Observed below-cloud particle number concentrations and updraft velocities suggest that CCN activation occurs in the transitional regime according to the designations defined by Reutter et al. (2009), and simulated sensitivity to hygroscopicity agrees well with those previously reported for the transition regime (0.17–0.2) (Reutter et al., 2009; Ward et al., 2010). Aging processes during transport likely lead to internally rather than externally mixed aerosol in the MBL. The simulated error in predicted CDNC when assuming fully externally mixed components is only 7.6–8.7% for the three modeled cases. This aligns with the observation of similarly accurate CCN closure results for the MBL when assuming internally or externally mixed components and a $\kappa_{\text{org}}$ of 0.1 or larger. As the volume fraction of inorganic aerosol in the accumulation mode is likely to increase with increasing distance from the coast, this predicted mixing-state-related error may be an upper bound for marine conditions in general.

Previous aerosol-cloud parcel modeling studies have demonstrated that the sensitivity of predicted CDNC to aerosol hygroscopicity tends to decrease as bulk hygroscopicity increases, especially for the aerosol-limited and transitional aerosol activation regimes (Reutter et al., 2009; Sánchez-Gáñita et al., 2017). If this is the case, accurate hygroscopicity characterization in marine regions subject to organic

Table 5

| Parameter | RF5-1 | RF5-2 | RF16 |
|-----------|-------|-------|------|
| $N_{\text{CN}, \text{Aitken}}$ (cm$^{-3}$) | 296   | 301   | 128  |
| $D_{\text{Aitken}}$ (nm) | 55    | 57    | 70   |
| $\kappa_{\text{Aitken}}$ | 1.27  | 1.27  | 1.24 |
| $N_{\text{CN, Accum}}$ (cm$^{-3}$) | 0.36  | 0.42  | 0.21 |
| $D_{\text{Accum}}$ (nm) | 492   | 465   | 406  |
| $\kappa_{\text{Accum}}$ | 1.04  | 1.09  | 1.24 |
| $w$ (m s$^{-1}$) | 2.21  | 2.20  | 1.96 |
| $w/N_{\text{CN}}$ (m s$^{-1}$ cm$^{-3}$) | 0.37  | 0.34  | 0.28 |
| Activation Regime | Trans. | Trans. | Trans. |

Note. “Activation Regime” refers to the classifications of cloud droplet formation environments developed by Reutter et al. (2009). “Trans.” = transitional.
aerosol inputs, which contain aerosol with lower-than-average \( \kappa \) values, may be more important for global CDNC prediction accuracy than accurate hygroscopicity characterization in remote regions subject to aerosol sources with different, but elevated, hygroscopicities (e.g., ammonium sulfate \( [\kappa = 0.61] \) vs. sodium chloride \( [\kappa = 1.28] \)).

To investigate this possibility, we calculated local CDNC sensitivity to aerosol hygroscopicity for four hypothetical marine aerosol size distributions. Rather than performing a linear regression on data obtained from a broad range of hygroscopicities, as was done for the data shown in Figure 10, local sensitivities refer to calculations performed on incremental variations in \( \kappa \) (e.g., \( \kappa = 0.1 \) vs. \( 0.2 \)).

Figure 11 displays the size distributions used as well as the sensitivity results. In order to span the likely range of size distributions observed in marine environments, the “Coastal” distribution is similar to median distributions observed during RF4 and RF5. A “Remote” distribution was generated using reported size distribution parameters from measurements over the remote subtropical N. Pacific by Ueda et al. (2016). Two additional size distributions were produced by interpolating between the “Coastal” and “Remote” distributions.

Total particle concentrations in the simulations varied between 300 and 800 cm\(^{-3}\) depending on the size distribution used. Five different updraft velocities were simulated \( (w = 0.1–0.5 \text{ m s}^{-1}) \), corresponding to the range typically observed within MSc over the Pacific (Zheng et al., 2016).

A few notable trends are evident in the results shown in Figure 11. As has been previously reported, CDNC sensitivity to aerosol hygroscopicity tends to decrease as hygroscopicity increases. However, even at low hygroscopicities, calculated sensitivities never exceed 0.3, suggesting that at a maximum, a 50% error in marine aerosol hygroscopicity should lead to an error of only 15% in predicted CDNC. Sensitivity slightly increases as the assumed particle concentration increases, and therefore, hygroscopicity is slightly less important in remote marine environments than in more polluted, coastal locations, as expected. In typical remote marine conditions \( (\kappa \approx 0.6) \) for instance, a 50% error in hygroscopicity is associated with only a \( \sim 2.5–7.5\% \) error in predicted CDNC, while in coastal environments \( (\kappa \approx 0.35) \) the error is estimated to be \( \sim 7.5–15\% \).

Figure 10. Sensitivity of calculated CDNC to accumulation mode aerosol hygroscopicity \( (\kappa) \), below-cloud aerosol particle number concentration \( (N_{CN}) \), accumulation mode geometric mean diameter \( (D_{pg}) \), accumulation mode standard deviation \( (\sigma) \), and updraft velocity \( (w) \). Data obtained during three cloud sampling passes were used as model constraints and are listed in Table 5. Numbers near each group of symbols represent average values from simulations in this study. Green symbols correspond to values reported by Reutter et al. (2009) for the transitional activation regime, while those in black correspond to values reported by McFiggans et al. (2006).

Figure 11. (Left) Aerosol number size distributions used as aerosol-cloud-parcel model inputs and (right) local CDNC sensitivities to aerosol hygroscopicity calculated using five updraft velocities. \( N_{CN} \) refers to the aerosol number concentration represented by each aerosol size distribution.
When simulating certain combinations of updraft velocity and aerosol size distribution, the sensitivity of predicted CDNC to aerosol hygroscopicity does not decrease monotonically as hygroscopicity increases. Furthermore, at a given hygroscopicity value shown in Figure 11, CDNC sensitivity is a non-monotonic function of updraft velocity. Here, we demonstrate that these phenomena are a result of activation of the distinct Aitken aerosol mode. Variation in CDNC sensitivity to hygroscopicity with increasing updraft velocity is shown in Figure 12 for κ = 0.6–0.8. Local CDNC sensitivity to hygroscopicity initially decreases with increasing updraft velocity before increasing again at updraft velocities >0.2–0.3 m s\(^{-1}\). This trend is consistent regardless of κ range analyzed; however, the shape of the curve becomes “stretched” horizontally as κ values decrease (Figure 12). Using a unimodal size distribution, Reutter et al. (2009) demonstrated that moving from the transitional to the aerosol-limited regime caused CDNC sensitivity to hygroscopicity to decline for all κ > 0.05. For the four marine size distributions simulated in this study, increasing the updraft velocity from 0.1 to 1.0 m s\(^{-1}\) shifts activation from the transitional regime to the aerosol-limited regime, implying CDNC sensitivity to hygroscopicity should subsequently decline. Our observation of the opposite phenomenon is due to the fact that at low (w = 0.1 m s\(^{-1}\)) and high (w = 1–1.5 m s\(^{-1}\)) updraft velocities, critical diameters produced within the rising air parcel occur near the peak of the accumulation and Aitken aerosol modes, respectively (Figure 12b). As the size distribution is peaked at these locations, subtle changes in aerosol hygroscopicity that induce small changes in the critical diameter result in a relatively large change in computed CDNC—hence elevated sensitivity to hygroscopicity. In contrast, for moderate (w ~ 0.2–0.3 m s\(^{-1}\)) updraft velocities, minimum critical diameters occur between the peaks of the Aitken and accumulation modes, and for very strong updraft velocities (w > 1.5–2 m s\(^{-1}\)) minimum critical diameters occur at sizes smaller than the peak of the Aitken mode, leading to lowered sensitivity (Figure 12b). This implies that in aerosol-limited environments with bimodal aerosol size distributions, the sensitivity of CDNC to hygroscopicity cannot necessarily be assumed to be negligible based solely on the ratio of the updraft velocity to particle number concentration. Ultimately, our results suggest that the sensitivity of marine CDNC to hygroscopicity is maximized in weak updraft conditions occurring in MSc (w < 0.2 m s\(^{-1}\)), where hygroscopicity of the accumulation, rather than the Aitken, mode is most relevant to accurate CDNC prediction, and in relatively strong updraft conditions (0.5 < w < 2 m s\(^{-1}\)) in either MSc or marine cumulus (Clarke et al., 1996), where Aitken mode hygroscopicity has a larger influence on CDNC than that of the accumulation mode.

4. Summary and Conclusions

Measurements of aerosol properties obtained over the N.E. Pacific Ocean during the MACAWS campaign in June and July 2018 were combined with results from an aerosol-cloud-parcel model to gain insight into aerosol hygroscopicity and its influence on CCN and MSc CDNC prediction in this environment. Three characteristic vertical regions were characterized, corresponding to the MBL, FT, and AC-OAL. Within the MBL,
flight-averaged hygroscopicities varied from values typical of continental environments ($\kappa = 0.27$), to those representative of remote marine locations ($\kappa = 0.72$) (Pringle et al., 2010). Distinct influences on MBL particle characteristics, including shipping emissions, entrainment from the AC-OAL, and transport from the remote Pacific, were identified through analysis of hygroscopicity data. In the AC-OAL, observed hygroscopicity suggests a dominant contribution of organic aerosol in both the Aitken and accumulation mode size ranges.

For the majority of flights, measured CCN concentrations could be reproduced within 20% using measurements of the aerosol size distribution, bulk hygroscopicity, and an assumption of either internally or externally mixed organic and inorganic components, in agreement with past results in non-urban locations (e.g., Ervens et al., 2010). Notably, for five of the seven flights, MBL CCN were better predicted when assuming a constant aerosol number size distribution derived from the median value measured in the MBL than when assuming a constant $\kappa$ typical of remote marine locations (0.72).

Results from an aerosol-cloud-parcel model confirm that the sensitivity ($S(\kappa) = \delta\ln N_{CDNC}/\delta\ln \kappa$) of predicted CDNC to accumulation mode aerosol hygroscopicity (0.19) is substantially smaller than sensitivity to size distribution parameters, such as the accumulation mode geometric diameter (0.49) and standard deviation ($\sim 0.64$). Simulations using a variety of possible MBL aerosol size distributions and hygroscopicities suggest that a 50% error in predicted hygroscopicity should rarely produce a CDNC error greater than 15%. However, model results further suggest that CDNC sensitivity to hygroscopicity does not monotonically decrease with increasing updraft velocity. Rather, sensitivity appears to decrease or remain constant with increasing updraft velocities from low to moderate values (e.g., 0.1–0.3 m s$^{-1}$) and then increase as updraft velocities increase further (>0.4 m s$^{-1}$) due to activation of the distinct Aitken mode of the aerosol size distribution. This phenomenon is observed despite the fact that at large updraft velocities (>0.4–0.5 m s$^{-1}$), marine conditions generally occupy the aerosol-limited regime of cloud droplet activation. Ultimately, CDNC sensitivity to hygroscopicity is predicted to be maximized in weak updraft conditions occurring in MSc (<0.2 m s$^{-1}$) and in strong updraft conditions (>0.5 m s$^{-1}$) expected to occur in either MSc or marine cumulus.

Data Availability Statement

Airborne field data used in this work can be accessed on the Figshare database (Sorooshian et al., 2017: https://doi.org/10.6084/m9.figshare.5099983.v10).

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