2	tral Amazonia: Seasonal and diel variations and impact of anthropogenic emissions
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CCN activity and organic hygroscopicity of aerosols downwind of an urban region in cen-

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- 33 Abstract
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35 During the Observations and Modeling of the Green Ocean Amazon (GoAmazon2014/5) 36 campaign, size-resolved cloud condensation nuclei (CCN) spectra were characterized at a research site (T3) 60 km downwind of the city of Manaus, Brazil, in central Amazonia for one year 37 38 (12 March 2014 to 3 March 2015). Particle hygroscopicity (κ_{CCN}) and mixing state were derived 39 from the size-resolved CCN spectra, and the hygroscopicity of the organic component of the aer-40 osol (κ_{org}) was then calculated from κ_{CCN} and concurrent chemical composition measurements. 41 The annual average $\kappa_{\rm CCN}$ increased from 0.13 at 75 nm to 0.17 at 171 nm, and the increase was 42 largely due to an increase in sulfate volume fraction. During both wet and dry seasons, $\kappa_{\rm CCN}$, $\kappa_{\rm org}$, 43 and particle composition under background conditions exhibited essentially no diel variations. 44 The constant κ_{org} of ~ 0.15 is consistent with the largely uniform and high O:C value (~0.8), in-45 dicating that the aerosols under background conditions are dominated by the aged regional aero-46 sol particles consisting of highly oxygenated organic compounds. For air masses strongly influ-47 enced by urban pollution and/or local biomass burning, lower values of κ_{org} and organic O:C 48 atomic ratio were observed during night, due to accumulation of freshly emitted particles, domi-49 nated by primary organic aerosol (POA) with low hygroscopicity, within a shallow nocturnal 50 boundary layer. The O:C, κ_{org} , and κ_{CCN} increased from the early morning hours and peaked 51 around noon, driven by the formation and aging of secondary organic aerosol (SOA) and dilution 52 of POA emissions into a deeper boundary layer, while the development of the boundary layer, 53 which leads to mixing with aged particles from the residual layer aloft, likely also contributed to 54 the increases. The hygroscopicities associated with individual organic factors, derived from PMF 55 analysis of AMS spectra, were estimated through multi-variable linear regression. For the SOA 56 factors, the variation of the κ value with O:C agrees well with the linear relationship reported 57 from earlier laboratory studies of SOA hygroscopicity. On the other hand, the variation in O:C of 58 ambient aerosol organics is largely driven by the variation in the volume fractions of POA and 59 SOA factors, which have very different O:C values. As POA factors have hygroscopicity values 60 well below the linear relationship between SOA hygroscopicity and O:C, mixtures with different 61 POA and SOA fractions exhibit a steeper slope for the increase of κ_{org} with O:C, as observed 62 during this and earlier field studies. This finding helps better understand and reconcile the differ-63 ences in the relationships between κ_{org} and O:C observed in laboratory and field studies, therefore providing a basis for improved parameterization in global models, especially in a tropical 64 65 context.

67 **1 Introduction**

Atmospheric aerosols have a major impact on the radiative balance of the Earth's climate 68 69 system by changing the microphysical structure, lifetime, and coverage of clouds. For the same 70 liquid water content, high aerosol concentration leads to more, smaller cloud droplets, and there-71 fore higher cloud albedo (Twomey, 1977). The smaller droplet size also delays or inhibits warm 72 precipitation, leading to increases in both cloud lifetime and coverage (Albrecht, 1989), and ul-73 timately invigoration of convective clouds (Rosenfeld et al., 2008). Currently, the effects of aer-74 osol on clouds remain one of the largest uncertainties in simulated climate change during indus-75 trial era, and a large portion of this uncertainty is due to the natural aerosol properties and pro-76 cesses represented in models (Carslaw et al., 2013; Ghan et al., 2013). The Amazon represents 77 more than half of the planet's rainforest and is a rapidly changing region where deforestation, 78 human activity and natural resource needs are all at play in changing the ecosystem (Andreae et 79 al., 2015; Batistella et al., 2013; Davidson et al., 2012). The Amazon basin also represents at 80 times one of the cleanest continental regions on the planet where it is still possible to find ex-81 tended periods of little or no impact of anthropogenic activity, although the long-distance 82 transport of pollution is occasionally observed (Andreae et al., 2015; Hamilton et al., 2014; 83 Martin et al., 2010b; Wang et al., 2016a; Wang et al., 2016b; Williams et al., 2002). This makes 84 the Amazon Basin an ideal location to characterize aerosol under near natural conditions and as-85 sess the impact due to urban emissions and biomass burning (Kuhn et al., 2010). The biogenic 86 activity of this region makes it a major source of organic carbon released into the atmosphere via 87 isoprene and monoterpenes (Guenther et al., 2006; Guenther et al., 2012; Kesselmeier et al., 88 2002; Kuhn et al., 2007) which are mediated by biotic stress through heat, sunlight and changes 89 in CO_2 (Heald et al., 2009).

90 To understand the impact of aerosol on clouds and climate requires knowledge of the concen-91 tration of cloud condensation nuclei, which are particles that are able to form cloud droplets un-92 der relevant atmospheric conditions. The minimum supersaturation required to activate a particle 93 into a cloud droplet can be predicted using κ -Köhler theory based on particle size and the single 94 hygroscopicity parameter κ , which combines a number of thermodynamic properties required for 95 the description of water activity of the growing droplets (Petters and Kreidenweis, 2007). The 96 value of κ is determined by the physicochemical properties of the solutes, including their molar 97 volume, activity coefficient, and the effect on surface tension. For multi-component particles, κ 98 is the volume average of participating species. Hygroscopicity also describes particle growth un-99 der sub-saturated conditions and can be derived from the particle growth factor (GF). However, 100 particles sometime exhibit larger κ values for droplet activation (derived from CCN measure-101 ments under supersaturated conditions) than for particle growth (derived from particle GF under 102 sub-saturated conditions) (e.g., Duplissy et al., 2008; Good et al., 2010; Mikhailov et al., 2013; 103 Pajunoja et al., 2015; Wex et al., 2009). In this paper, "hygroscopicity" represents κ associated 104 with droplet activation derived from CCN measurements unless noted otherwise.

105 The hygroscopicities of typical inorganic species in ambient particles are relatively well 106 known (Petters and Kreidenweis, 2007). However, atmospheric aerosols consist of a large num-107 ber of organic compounds, which often dominate the total fine aerosol mass, especially in forest-108 ed areas (e.g., de Sá et al., 2016; Jimenez et al., 2009; Zhang et al., 2007). The hygroscopicity of 109 aerosol organics (κ_{org}) have been examined in both laboratory (e.g., Asa-Awuku et al., 2009; 110 Duplissy et al., 2011; King et al., 2009; Lambe et al., 2011; Massoli et al., 2010; Prenni et al., 111 2007; Raymond and Pandis, 2003) and field studies (e.g., Cerully et al., 2015; Chang et al., 2010; 112 Dusek et al., 2010; Gunthe et al., 2009; Jimenez et al., 2009; Lathem et al., 2013a; Mei et al.,

113 2013a; Mei et al., 2013b; Moore et al., 2011; Moore et al., 2012; Pöhlker et al., 2016; Rose et al., 114 2010; Shantz et al., 2008; Wang et al., 2008). Overall, these studies show that aerosol organics 115 exhibit a wide range of κ values from 0 to ~0.3, and κ_{org} often increases substantially during aer-116 osol aging in the atmosphere (e.g., Duplissy et al., 2011; Jimenez et al., 2009; Lambe et al., 117 2011; Massoli et al., 2010; Mei et al., 2013a).

118 A number of recent studies examined the sensitivity of predicted CCN concentration and 119 cloud droplet number concentration to aerosol properties (e.g., Ervens et al., 2010; Kammermann 120 et al., 2010; McFiggans et al., 2006; Mei et al., 2013a; Reutter et al., 2009; Rissman et al., 2004; 121 Roberts et al., 2002; Wang, 2007; Wang et al., 2008). These studies show that the predicted CCN 122 concentration is often sensitive to $\kappa_{\rm org}$, especially for aerosol under background conditions where 123 organics tend to dominate submicron aerosol mass (Liu and Wang, 2010; Mei et al., 2013a). Us-124 ing a constant κ_{org} may lead to large biases in predicted CCN concentrations and aerosol indirect 125 forcing (Liu and Wang, 2010). Therefore, it is imperative to understand organic hygroscopicity 126 under background conditions, such as in the Amazon forest, as well as the variation of organic 127 hygroscopicity due to anthropogenic emissions.

128 There have been several studies of aerosol hygroscopicity in the Amazon Basin over the past 129 20 years (Gunthe et al., 2009; Mikhailov et al., 2013; Pöhlker et al., 2016; Rissler et al., 2006b; 130 Roberts et al., 2001; Vestin et al., 2007; Whitehead et al., 2016; Zhou et al., 2002). Gunthe et al. 131 (2009) performed size resolved CCN measurements during the wet season in February and 132 March 2008 as part of the AMAZE-08 campaign (Martin et al., 2010a). That study reported no 133 diel cycle in the CCN concentration during periods with little or no influence of pollution. 134 Pöhlker et al. (2016) measured size-resolved CCN spectra at a remote background site (Amazon Tall Tower Observatory ATTO) over a one-year period from March 2014 to February 2015 and 135

136 observed no diel cycle and only weak seasonal trends in derived particle hygroscopicity, while 137 CCN concentrations had a pronounced seasonal cycle as the background aerosol concentration 138 was strongly influenced by regional biomass burning during the dry season. During the SMOCC-139 2002 campaign (Large scale Biosphere atmosphere experiment in Amazonia – SMOke, aerosols, 140 Clouds, rainfall, and Climate) particle hygroscopicity was derived from HTDMA measurements 141 in the state of Rondônia in the southwest of the Amazon region during the dry season from Sep-142 tember to November of 2002 (Rissler et al., 2006b; Vestin et al., 2007). The study concluded that 143 the diel variation in the aerosol hygroscopicity could be linked to the structure and dynamics of 144 the boundary layer. Local sources dominated night-time aerosol properties with downward mix-145 ing from the residual layer aloft as the day progressed. All of these studies found that particle 146 hygroscopicity increased with particle size (from the Aitken to accumulation modes), consistent 147 with higher sulfate content at larger sizes (Gunthe et al., 2009). The same boundary layer evolu-148 tion has been found to influence particle number and CCN evolution in a number of other related 149 studies (Fisch et al., 2004; Martin et al., 2010a; Rissler et al., 2006b; Vestin et al., 2007; 150 Whitehead et al., 2010; Zhou et al., 2002).

151 In this study we present measurements of size resolved CCN spectra at five particle diame-152 ters ranging from 75 to 171 nm downwind of Manaus, Brazil, in central Amazonia for a period 153 of one year from March 12, 2014 to March 3, 2015. Particle hygroscopicity, mixing state, and 154 organic hygroscopicity are derived from the size-resolved CCN activated fraction and concurrent 155 aerosol composition measurements. The diel variations of these properties are examined for dif-156 ferent seasons (i.e., wet season, dry season, and transition seasons) and for different types of rep-157 resentative air masses, including background conditions, as well as influences of urban pollution 158 plumes and/or local biomass burning. During the wet season, the background air mass represents

near natural conditions, with occasional impact from anthropogenic emissions, while in the dry season, the background is dominated by regional biomass burning aerosol particles. The relationship between organic hygroscopicity and particle oxidation level (i.e., O:C atomic ratio) is examined for both dry and wet seasons. Hygroscopicities associated with organic factors from Aerosol Mass Spectrometry (AMS) Positive Matrix Factorization (PMF) analysis are derived, and their relationship with the O:C ratio is compared with those from previous laboratory studies.

165 2 Experimental Setup

166 2.1 Measurement Sites

167 Observations and Modeling of the Green Ocean Amazon (GoAmazon2014/5), sponsored by 168 the US Department of Energy (DOE) and several Brazilian and German agencies, took place at 169 multiple surface sites surrounding Manaus, Brazil, from January 2014 through December 2015 170 (Martin et al., 2016b). This work focuses on the measurements carried out at a downwind site 171 (T3, 3°12'47.82"S, 60°35'55.32"W, 60 km west of Manaus) from March 2014 to March 2015. 172 Depending on the wind direction, the T3 site experienced conditions ranging from nearly natural 173 to heavily polluted. More detailed characterizations of aerosol and gas phase chemical composi-174 tion were carried out at the T3 site during Intensive Operating Periods (IOPs) from February 1, 175 2014 to March 31, 2014 and from August 15, 2014 to October 15, 2014. In addition, data from 176 two sites normally upwind of Manaus are also used in this study. These background sites include 177 the T0a site, (ATTO, 2° 8'47.88"S, 59° 0'18.00"W) (Andreae et al., 2015) and the T0t site, 178 (2.6091°S, 60.2093°W) (Martin et al., 2010a).

179 **2.2 Activated Fraction of size-selected Particles**

180 The CCN activation fraction of size-selected particles was measured using a Differential 181 Mobility Analyzer (DMA) coupled to a condensation particle counter (CPC, TSI Inc., 3010) and 182 a cloud condensation nuclei counter (CCNC, Droplet Measurement Technologies, Boulder, CO) 183 (Frank et al., 2006; Mei et al., 2013a; Moore et al., 2010; Petters et al., 2007). Aerosol particles were sampled with a total flow rate (Q_a) of 1.53 L min⁻¹ from a height of 5 m above ground level 184 185 and were dried to RH below 20% by a Nafion dryer immediately upon entering the instrument 186 container. The dried aerosol particles then reached steady state charge distribution inside a Kr-85 187 aerosol charger (TSI, model 3077A), prior to being introduced into the DMA operated with a sheath flow rate (Q_{sh}) of 15.3 L min⁻¹ to maintain a 10:1 sheath to aerosol flow ratio (Q_{sh}/Q_a) . 188 189 The aerosol particles were size-selected by the DMA and the size-selected particles were simultaneously characterized by a CPC ($Q_{CPC} = 0.53 \text{ Lmin}^{-1}$) and a CCNC (see Fig. S1 in the Sup-190 191 plementary Information for further details). This system had been operated in previous field 192 campaigns by scanning the particle size while CCNC supersaturation was held constant (Mei et 193 al., 2013a; Mei et al., 2013b). During GoAmazon2014/5 the particle size classified by the DMA 194 was stepped through seven particle diameters (51, 75, 94, 112, 142, 171, and 222 nm), while the CCNC supersaturation was also changed at each diameter by stepping the flow rates (Q_{CCN} rang-195 196 ing from 0.2 to 1.0 L min⁻¹) and temperature gradient ($\Delta T = 4.5, 5.5, 6.5, 8.0$ and 10.0°C). At a 197 given supersaturation, data were acquired for a minimum of 30 s and until 1500 particles were 198 counted by the CPC or up to a maximum time of 120 s. Depending on the aerosol number size 199 distribution, the measurement cycled through the seven particle sizes in 1 - 2 h (see Fig. S1 and 200 S2 for further details of the measurement setup and sampling protocol). The sampling sequence 201 was designed so that the change of CCNC supersaturation was mostly accomplished by stepping

202 flow rates, as the CCNC reaches steady state faster following flow changes than temperature 203 changes. Change of the temperature gradient was kept at a minimum frequency, but was neces-204 sary given the wide range of supersaturation explored. Given the low particle number concentration (e.g., $\sim 200 \text{ cm}^{-3}$ under background conditions during the wet season), these approaches were 205 206 important to achieve adequate counting statistics with good time resolution to capture changes of 207 air mass within 10 - 20 minutes (Liu et al., 2016). The supersaturation of the CCNC was cali-208 brated using ammonium sulfate aerosol, as described previously in the literature (e.g., Mei et al., 209 2013a), at each operational set point (Q_{CCN} and ΔT), ranging from 0.075 – 1.1 %. Fluctuation of 210 the temperature inside the instrument container, ranging from 20 to 30°C over the course of a 211 day, led to substantial variation in the absolute temperature inside the CCNC growth chamber. 212 Calibrations were therefore repeated under a range of container and associated growth chamber 213 temperatures. The dependence of the supersaturation on the temperature at the top of CCNC col-214 umn (instrument temperature T_1) was derived for each Q_{CCN} and ΔT pair and used to retrieve the 215 supersaturation over the range of the instrument operating conditions (see Fig. S3 and further 216 description in the Supplementary Information).

217 **2.3 Aerosol Chemical Composition**

Non-refractory sub-micron aerosol composition (organics, sulfate, nitrate, ammonium, and chloride) was measured by a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS, Aerodyne Research Inc. (DeCarlo et al., 2006) during the two IOPs, and by an Aerosol Chemical Speciation Monitor (ACSM, Aerodyne Research, Inc., Ng et al., 2011) from July 2014 to March 2015. The AMS sampled from an inlet equipped with a PM2.5 cyclone located at 5 m above ground level. The ambient sample was first dried outside the container by a poly-tube Nafion dryer (Perma Pure, model PD-100T). Once inside the container the sample was further dried by a mono-tube Nafion dryer (Perma Pure, model MD-110) to achieve RH < 40% and was split between the AMS and a Scanning Mobility Particle Sizer (SMPS, TSI, model 3081). Ambient measurements were obtained every 4 of 8 minutes. Further details of the AMS set up and operation are described in de Sá et al. (2016).

229 The ACSM was a part of the Aerosol Radiation Measurement (ARM) Mobile Facility-1 230 (AMF-1) Mobile Aerosol Observation System (MAOS). Aerosol was sampled through an inlet 231 located 10 m above the ground. The aerosol sample was first dried through five large (40 x 1.75 232 cm I.D.) Nafion dryers before being distributed among various instruments including the ACSM. 233 The ACSM sampling alternated between with and without an in-line filter using a 3-way valve, 234 such that aerosol-free background could be subtracted from the ambient measurement. Twenty-235 eight ambient and background scans of the quadrupole mass spectrometer (unit mass resolution) 236 were averaged to give one measurement every 30 min. The mass concentrations of organic spe-237 cies, sulfate, nitrate, ammonium, and chloride were derived from measurements using approach-238 es described in Ng et al. (2011).

239 Refractory black carbon (rBC) was measured using both a Single Particle Soot Photometer 240 (SP2, Droplet Measurement Technologies, Boulder, CO) and an aethalometer (Magee Scientific) 241 co-located with the AMS and ACSM. The SP2 measures rBC using laser-induced incandescence, 242 whereas the aethalometer measures equivalent black carbon (BCe) (Andreae and Gelencsér, 243 2006) using light absorption from particles collected onto a filter. While these are fundamentally 244 different aerosol properties, both species (rBC and BC_e) were treated as equivalent in this study, 245 and BC_e concentration was adjusted to match that of rBC using the approach detailed in Section 246 2.2 of the Supplementary Information.

247 **2.4 Additional Relevant Measurements**

248 Additional measurements of aerosol microphysics, trace gas concentrations, and atmospheric 249 conditions used in this study are briefly described here. These measurements were part of the de-250 ployment of the ARM AMF-1 facility during GoAmazon 2014/5 (Martin et al., 2016b). Relevant 251 aerosol measurements include dry particle number-size distributions from 10 to 480 nm by a 252 SMPS (TSI Inc. Model 3081) and the number concentration of particles with diameters greater 253 than 10 nm by a CPC (TSI Inc. Model 3772). Mixing ratios of CO and O_3 were characterized by 254 an Off-Axis Integrated Cavity Output Spectroscopy CO, N₂O and H₂O analyzer (model number 255 908-0014, Los Gatos) and UV Photometry O₃ analyzer (model 49i, Thermo Scientific Inc.), re-256 spectively. Oxides of nitrogen (NO, NO_x, NO₂, NO_y) were measured using a chemiluminescence 257 technique (details given in Section S2.1 of the Supplementary Information). Meteorological data 258 included relative humidity (RH), ambient temperature, wind speed and direction, and rain accu-259 mulation. The vertical profile of atmospheric backscatter (clouds and aerosol) and boundary lay-260 er heights were estimated from ceilometer (model CL31, Vaisala) measurements.

261 3 Methods

262 **3.1 Derivation of Particle Hygroscopicity and Mixing State**

263 The particle hygroscopicity parameter, κ (Petters and Kreidenweis, 2007), was derived from 264 the activation spectrum (i.e., activated fraction as a function of supersaturation *S*) at the individu-265 al particle sizes using approaches detailed in the literature (Bougiatioti et al., 2011; Cerully et al., 266 2011; Lance et al., 2013; Mei et al., 2013a; Rose et al., 2008b). The activation spectrum of size-267 selected particles was first corrected for the influence of multiply charged particles, which is es-268 timated using the size distribution measured by the SMPS in MAOS and the activation spectrum 269 measured at the sizes of the doubly and triply charged particles (See Section S3.1). The corrected 270 activation spectrum of size selected particles was then fit with a cumulative lognormal (Mei et 271 al., 2013a; Rose et al., 2008a) functional form (See Fig. S6 for examples):

272
$$R_{a}(S) = \frac{E}{2} \left[1 + erf\left(\frac{\ln S - \ln S^{*}}{\sqrt{2\sigma_{s}^{2}}}\right) \right]$$
(1)

273 where R_a is the activated fraction as a function of supersaturation S, E is maximum activated 274 fraction, and (1-E) represents the number fraction of particles consisting of only non-hygroscopic 275 species (e.g., uncoated rBC) that cannot serve as CCN under typical atmospheric supersatura-276 tions. S* is the supersaturation at which R_a reaches 50% of E, and represents the median critical 277 supersaturation of size-selected particles that serve as CCN. The value of σ_s is related to the slope of the increasing R_a with S near S^{*} and reflects the heterogeneity of critical supersaturation, 278 279 which to a large degree arises from the heterogeneity of the hygroscopicity among size-selected 280 particles (Cerully et al., 2011; Mei et al., 2013a). The probability density function of hygroscopicity for size-selected particles is derived from the $R_a(S)$. The average hygroscopicity $\overline{\kappa_{\text{CCN}}}$ and 281 dispersion of the hygroscopicity $\sigma(\kappa)/\overline{\kappa_{_{\rm CCN}}}$ for the size-selected CCN were then derived from 282 283 the probability density function of hygroscopicity using the approach detailed in Section S3.4 in 284 Supplementary Information. The dispersion of the hygroscopicity reflects the composition heter-285 ogeneity (i.e., mixing state) among size-selected particles (Mei et al., 2013a). For simplicity, we use $\kappa_{\rm CCN}$ to represent the average hygroscopicity of size-selected CCN in the following sections. 286 287 As hygroscopicities reported in this study were derived from particle dry diameter and critical 288 supersaturation, they represent "apparent hygroscopicity", which includes the potential impact due to the limited solubility of organics and the reduction of surface tension by surface activespecies (Sullivan et al., 2009).

291

292 **3.2 Derivation of Organic Aerosol Hygroscopicity**

293 The average particle hygroscopicity was then combined with the chemical composition data to derive the hygroscopicity of the organic component of the size-selected particles, $\kappa_{\rm org}$. Collec-294 295 tively, the AMS, ACSM, SP2, and aethalometer provided mass concentrations of organic spe-296 cies, sulfate, nitrate, ammonium, and rBC. The concentration of chloride was negligible (<<1% 297 of aerosol mass) and was not included in the analysis. Given the low concentrations during 298 GoAmazon 2014/5, size-resolved mass concentrations at the time resolution of the CCN meas-299 urement were not directly derived from AMS particle time of flight (PToF) mode data. For IOP 300 1, measurements were classified into three groups based on bulk organic mass fraction and the 301 characteristic mass size distribution of each species was averaged from measurements in each 302 group. For the dry season, the measurements were classified into three groups each for day and 303 night periods based on the bulk aerosol organic mass fraction, and the mass size distribution of 304 each species was averaged from measurements in each of the six groups. The size-resolved mass 305 concentrations of sulfate, nitrate, and organics at the time resolution of the CCN measurement 306 were then derived by scaling the total mass concentration using the average mass size distribu-307 tions for the corresponding group (based on the bulk organic mass fraction) of either wet or dry season (de Sá, 2017). The shape of the NH₄⁺ mass size distribution was assumed to be the same 308 309 as that of sulfate, as ammonium cations were primarily associated with sulfate. rBC was as-310 sumed to have the same size distribution shape as the total aerosol mass (i.e., mass fraction of 311 rBC was independent of particle size); though this assumption may not always be appropriate, 312 the effect is expected to be very small as the monthly average volume fraction of rBC was al-313 ways less than 4% (Fig. 1). A detailed description of the derivation of the size-resolved mass 314 concentrations is given in Section S4.1.

In most cases, NH_4^+ was insufficient to completely neutralize SO_4^{2-} . The concentrations of 315 both organonitrate and inorganic nitrate during the two IOPs were retrieved from AMS data 316 based on the ratio of ions NO^+ and NO_2^+ (de Sá, 2017; Fry et al., 2009). When the inorganic ni-317 trate mass concentration was negligible (i.e., less than 30 ng m⁻³), as in most of the cases, the 318 319 contributions of ammonium sulfate and ammonium bisulfate were calculated based on the mass concentrations of SO_4^{2-} and NH_4^+ (Nenes et al., 1998). In rare cases when the mass concentra-320 tion of inorganic nitrate was greater than 30 ng m⁻³, sulfate was assumed to be ammonium sul-321 322 fate. During non-IOP periods, only the total nitrate mass concentration is available, all nitrate 323 was assumed to be organonitrate (Cerully et al., 2015; Lathem et al., 2013b; Nenes et al., 1998; 324 Zhang et al., 2005), and sulfate fully neutralized by ammonium (see Section S4.2 in the Supple-325 ment for a sensitivity study of these assumptions for non-IOP periods). The -ONO₂ portion of the organonitrate was added back to the organic mass. We note that the amount of -ONO₂ added 326 327 back was typically small given the low mass fraction of nitrate in the aerosol. Cloud condensa-328 tion nuclei were assumed to be internal mixtures of (NH₄)₂SO₄, NH₄HSO₄, NH₄NO₃, organics, 329 and rBC, and the volume fractions of the species were derived from the mass concentrations and 330 densities. Densities of organics were estimated from the ratios of O:C and H:C measured by the AMS (Kuwata et al., 2012) and were on average 1450 ± 100 and 1470 ± 80 kg m⁻³ for IOP1 and 331 IOP2, respectively. Densities of 1770, 1790, 1730, 1800 kg m⁻³ were used for (NH₄)₂SO₄, 332 NH₄HSO₄, NH₄NO₃, and rBC (Bond and Bergstrom, 2006; Park et al., 2004), respectively. In 333

334 very rare cases, E was less than 100%, suggesting some of the size-selected particles were non-335 hygroscopic. The non-hygroscopic particles were assumed to consist entirely of rBC (Mei et al., 336 2013a). The volume concentration of the non-hygroscopic particles was derived as the product of 337 (1-E) and the total volume concentration (i.e., the sum of volume concentrations of organics, 338 (NH₄)₂SO₄, NH₄HSO₄, NH₄NO₃ and rBCs) at the size classified by the DMA. The volume con-339 centration of rBC internally mixed within the CCN-active particles was then calculated as the 340 difference between the total rBC volume concentration and the volume concentration of the non-341 hygroscopic particles. (Mei et al., 2013a; Mei et al., 2013b). Assuming a κ value of zero for rBC, 342 we can derive the hygroscopicity of the organic component of the CCN κ_{org} as:

343
$$\kappa_{\rm org} = \frac{1}{x_{\rm org}} \left(\kappa_{\rm CCN} - x_{(\rm NH_4)_2 SO_4} \kappa_{(\rm NH_4)_2 SO_4} - x_{\rm NH_4 HSO_4} \kappa_{\rm NH_4 HSO_4} - x_{\rm NH_4 NO_3} \kappa_{\rm NH_4 NO_3} \right)$$
(2)

where x_i is the volume fraction of the respective species. The κ values are 0.61, 0.7 and 0.67 for (NH₄)₂SO₄, NH₄HSO₄, and NH₄NO₃, respectively (Petters and Kreidenweis, 2007). The uncertainty in κ_{org} using these calculations has been derived using the approach detailed in earlier studies (Mei et al., 2013a; Mei et al., 2013b) and is on the order of 0.01 – 0.02 (which was generally between 10 and 20%) for this data set.

349 **3.2.1 Derivation of κ for AMS PMF Factors**

PMF was applied to the AMS mass spectra (Lanz et al., 2008; Ulbrich et al., 2009), and six organic factors were identified for each of the two IOPs (de Sá, 2017). For IOP2 the PMF analysis included data from 24 August to 15 October 15, 2014, excluding a major regional biomass burning event from 16 to 23 August, which was treated separately in the PMF analysis. For IOP1 (wet season), the six factors were isoprene-epoxydiol-derived secondary organic aerosol (IE- 355 POX-SOA), more-oxidized oxygenated organic aerosol (MO-OOA, i.e., highly oxidized organ-356 ics), less-oxidized oxygenated organic aerosol (LO-OOA), biomass burning organic aerosol 357 (BBOA) with characteristic peaks at m/z = 60 and 73 and correlated with the concentrations of 358 levoglucosan and vanillin, a factor with high contribution from m/z = 91 (Fac91) and correlated 359 with anthropogenic emissions of aromatics, and hydrocarbon-like organic aerosol (HOA). The 360 six factors for IOP2 included IEPOX-SOA, MO-OOA, LO-OOA, an aged biomass burning or-361 ganic aerosol factor (aged-BBOA), a fresh biomass burning organic aerosol factor (fresh-362 BBOA), and HOA. Further details of PMF analysis and the characteristics of the factors can be 363 found in de Sá (2017). The O:C ratio and calculated density for each factor are presented in Ta-364 ble 1. In this study, the O:C ratio was derived using Improved-Ambient Method (Canagaratna et 365 al., 2015).

For each IOP, hygroscopicities associated with the six factors were attributed based on multilinear regression of κ_{org} with respect to the volume fractions of the factors (Levenberg-Marquardt algorithm, IGOR Pro, Wavemetrics):

$$369 \qquad \kappa_{\rm org} = \sum_{i}^{n} \kappa_{i} x_{i} \tag{3}$$

where κ_i and x_i are the hygroscopicity and volume fraction of the individual organic PMF factors. The volume fraction was derived from mass concentrations and the densities of the factors. κ_{org} represents the average organic hygroscopicity at particle diameters (D_p) of 142 and 171 nm. As the PMF analysis is based on the mass spectra of the bulk sub-micrometer aerosol (i.e., MS mode measurements), an implicit assumption of Eq. (3) is that the bulk volume fractions of the factors represented those over the sizes at which κ_{org} was derived (i.e., $D_p = 142$ and 171 nm). The va376 lidity of this assumption is discussed in the results section. The robustness of the factor hygro-377 scopicity derived through linear regression depends on the variation of the factor volume fraction 378 during the measurement period. The HOA hygroscopicity was assumed as zero based on the re-379 sults from previous studies (Cappa et al., 2009; Cappa et al., 2011; Jimenez et al., 2009), and the 380 hygroscopicity of the other five factors were derived by multilinear regression as described 381 above.

382 **3.3 Classification of Seasons and Air Masses**

383 The one-year sampling period was divided into different seasons by grouping months accord-384 ing to the similarity of the aerosol properties and trace gas concentrations measured at the two 385 background sites, T0a and T0t, as well as monthly accumulated rainfall. In this study, the sea-386 sons were defined as follows: the first wet season: March, April and May of 2014; the first tran-387 sition season: June and July 2014; the dry season: August and September of 2014; the second 388 transition season: October, November and the first half of December 2014; and the second wet 389 season: the second half of December 2014 and January, February and the first few days of March 390 2015.

For each season, the air masses arriving at the T3 site were classified into three different types: background, urban pollution, and local biomass burning based on trace gas and aerosol measurements at all sites. During the wet season, the background air mass represented near natural conditions, with occasional impact from anthropogenic emissions, while in the dry season, the background was dominated by regional biomass burning aerosol particles. Polluted air masses represent those with strong influence from urban emissions, which were mostly from Manaus. The local-biomass-burning type describes those air masses strongly influenced by local (i.e.,

398 fresh) biomass burning activities, which dominated over the impact from urban pollution, if any. 399 For each season, background conditions were identified when CO and CN concentrations were 400 below the thresholds derived from measurements at the background TOa and TOt sites, and the 401 NO_v mixing ratio was below 1.5 ppb. Non-background conditions were identified by condensa-402 tion nuclei (CN) and CO concentrations above the respective threshold levels. As biomass burn-403 ing aerosol typically has a higher fraction of accumulation mode particles, and the emissions 404 from Manaus were more dominated by Aitken mode particles, the fraction of particles with di-405 ameter less than 70 nm was used to differentiate air masses strongly influenced by local biomass 406 burning from those with more impact from urban pollution (see Table S2 in Supplementary In-407 formation for details). Contamination by the emissions from an on-site diesel generator, grass 408 cutting activities, tractors, and other vehicles were evidenced by rBC concentrations above 1.0 µg m⁻³ or CN concentration above 10,000 cm⁻³. Over the one-year measurement period, back-409 410 ground, urban pollution, and local biomass burning represented 12.4%, 38.5%, and 28.4% of the 411 CCN measurements, respectively (Table S3). We note that the air masses arriving at the T3 site 412 often included contributions from different sources. The classification of the air masses using the 413 above three types clearly represents a simplification, but is very helpful for understanding the 414 properties of aerosols influenced by the various major sources.

415 **4 Results**

416 **4.1 Seasonal Trend and Size Dependence of Hygroscopicity and Chemical Composition**

417 The monthly average κ_{CCN} at the T3 site varied from 0.1 to 0.2 at five particle diameters 418 ranging from 75 nm to 171 nm (Fig. 1a), and was substantially lower than the value of 0.3 ± 0.1 419 suggested for continental sites (Andreae and Rosenfeld, 2008). This was due to the large organic volume fraction, up to 95% observed at the T3 site. In this study, measurements at 51 and 222 nm were not included, because the range of supersaturation sampled inside the CCN counter only adequately captured the activation spectrum for 51/222 nm particles with relatively high/low $\kappa_{\rm CCN}$ values, leading to a positive/negative bias of the average $\kappa_{\rm CCN}$. The value of $\kappa_{\rm CCN}$ exhibited similar seasonal variations at all five sizes. During the transition from wet to dry season, $\kappa_{\rm CCN}$ decreased by 20-30% with the absolute minimum of 0.116 occurring at 75 nm in September and October (Fig. 1a).

427 The seasonal trend of $\kappa_{\rm CCN}$ was mainly driven by the variation of $\kappa_{\rm org}$, which shows the low-428 est value in September during the dry season (Fig. 1b). Despite a strong increase of aerosol mass 429 concentration from wet to dry season due to biomass burning emissions, the organic volume 430 fraction exhibited little season variation, and was ~90% or higher at the four sizes from 94 to 431 171 nm (Fig. 1c). A minor increase in organic volume fraction in October might have also con-432 tributed to the lower κ_{CCN} value observed. The species volume fractions at 75 nm are not shown 433 due to the very low signal-to-noise ratio of the AMS PToF data in the small particle diameter 434 range. No clear seasonal trend was observed for sulfate volume fraction, which ranged from 3% 435 to 9% at the four sizes. The lack of clear seasonal trends of sulfate and organic fractions are con-436 sistent with observations at the T0a site (Andreae et al., 2015). Nitrate and rBC represented a 437 small fraction of aerosol volume, and were less than 1% and ~4%, respectively.

The average κ_{CCN} increased with increasing particle size for all three air mass types and during all the seasons (Fig. 2), consistent with decreasing organic volume fraction with increasing particle size (Fig. 1c). The κ_{CCN} at 75, 94, 112, 142, and 171 nm averaged for the one-year measurement period were 0.130 ± 0.028, 0.144 ± 0.039, 0.148 ± 0.043, 0.164 ± 0.046, and 0.167 ± 0.042 respectively. The value of κ_{CCN} and its size dependence under background conditions were

443 largely consistent among different seasons, and were in good agreement with those observed un-444 der near natural conditions during the AMAZE-08 campaign at T0t in the wet season (Gunthe et 445 al., 2009) and during the one-year period from March 2014 to February 2015 at the background 446 T0a site (Pöhlker et al., 2016). For the air masses with strong influence from local biomass burn-447 ing, the value of $\kappa_{\rm CCN}$ and its size dependence are consistent with the κ value derived from parti-448 cle growth factor measurements in July 2001, during a "recent biomass burning period" of the 449 CLAIRE-2001 study (Rissler et al., 2004), which took place at Balbina, about 125 km northeast 450 of Manaus. In contrast, κ values derived from particle growth factor measurements from 11 Sep-451 tember to 8 October 2002, during the dry period of the LBA-SMOCC (Rissler et al., 2006) are 452 substantially lower than $\kappa_{\rm CCN}$ observed at the T3 site for local biomass burning air masses at all 453 sizes. As LBA-SMOCC took place in the state of Rondônia in southwestern Amazonia with ex-454 tensive biomass burning activities during the dry season, the difference in κ could be due to the 455 differences in fire condition and the type of vegetation burned. Previous studies show particles 456 sometime exhibit larger κ values for droplet activation (derived from CCN measurements under 457 supersaturated conditions) than for particle growth (derived from particle growth factor under 458 sub-saturated conditions), this could also contribute to the differences in κ values. Compared to 459 $\kappa_{\rm CCN}$, $\kappa_{\rm org}$ was largely independent of particle size for all three air mass types, indicating that the 460 size dependence of κ_{CCN} was mainly due to the size dependence of the organic volume fraction 461 and particle composition (Fig. 1c - f). During the dry season, aerosols classified as urban pollu-462 tion and local biomass burning exhibited lower κ_{org} values compared to background aerosols, 463 contributing to the lower values of overall $\kappa_{\rm CCN}$.

464 **4.2 Diel Trends of Particle and Organic Hygroscopicities**

The diel variations of aerosol properties are presented in Figs. 3-7 for different air masses during the two IOPs. Aerosol properties derived from the size-resolved CCN measurements, including κ_{CCN} , $\sigma_{\kappa_{\text{CCN}}}$, $\overline{\kappa}_{\text{CCN}}$, and κ_{org} , and the volume fractions of different species were averaged at the three largest sizes ($D_p = 112$, 142, and 171 nm). The fraction of organic mass at m/z = 44 (f_{44}) and O:C were derived from the AMS bulk measurements. Also shown are diel variations of planetary boundary layer height, CN, and aerosol volume concentrations based on 5-minute average data.

472 **4.2.1 Wet Season Aerosol**

473 Background conditions

474 Figures 3 and 4 show the diel variations of aerosol properties during the wet season of 2014 475 for background and urban pollution air masses, respectively. There were only 0.7% of data clas-476 sified as local biomass burning (see Table S3), which is insufficient to evaluate the diel trends. 477 During the wet season, the background air mass represents near-natural conditions, with mini-478 mum impact from anthropogenic emissions, although long-distance transport of African biomass 479 burning may contribute to the aerosols observed (Chen et al., 2009; Wang et al., 2016b). Back-480 ground aerosol constantly exhibited relatively high hygroscopicity of ~0.19 throughout the day. 481 The lack of a diel trend in $\kappa_{\rm CCN}$ is also in agreement with the results from the T0a site (ATTO), 482 which is upwind of Manaus and served as a background site (Fig. 8). The particle composition 483 averaged for the three particles diameters was dominated by organics, representing ~ 90% of the 484 aerosol volume. The lack of a diel trend in $\kappa_{\rm CCN}$ and $\kappa_{\rm org}$ suggest little variation in particle com-485 position throughout the day. The nearly constant κ_{org} of ~0.16 is also consistent with the lack of a 486 diel trend in f_{44} and O:C. The values of f_{44} and O:C are ~0.2 and ~0.8, respectively, indicating 487 that the aerosol under background condition during the wet season was dominated by the aged 488 regional aerosol particles consisting of highly oxygenated organic compounds.

Aerosol number and volume concentrations exhibited a minimum at \sim 310 cm⁻³ and \sim 0.3 μ m³ 489 cm⁻³, respectively, just before sunrise. The number and volume concentrations started increasing 490 after sun rise, and peaking at 400 cm⁻³ and 0.8 um³ cm⁻³ in the afternoon. These diel variations 491 492 are partially attributed to the wet scavenging of accumulation mode particles, which dominate 493 the submicron particle concentrations under background conditions, and the mixing of the parti-494 cles from the residual layer aloft down to the surface as the boundary layer develops in the morn-495 ing. During the night, the radiative cooling at the surface leads to a shallow nocturnal boundary 496 layer with low and variable winds. RH near surface was near 100%, and fog or mist was identi-497 fied by the weather station (Present Weather Detector, Visalia) 62% of the time during the one-498 year measurement period. The gradual decrease of particle number and volume concentration 499 during these fog events were due to the wet deposition of the accumulation mode particles acti-500 vated into droplets. Similar decreases of particle number concentration were previously reported 501 during night fog events in the tropical rainforest in Borneo (Whitehead et al., 2010). After sun-502 rise, the boundary layer deepened on average from less than 200 up to 800 m as a result of solar 503 heating (Fig. 3g). Consequently, particles in the residual layer aloft (Fisch et al., 2004; Rissler et 504 al., 2006b), which were not impacted by the fog, were mixed down to the surface, leading in part 505 to the observed increases in both number and volume concentrations. Such mixing of particles 506 from the residual layer in the morning had been observed previously in the Amazon basin during 507 the dry season (Rissler et al., 2006a). The formation of secondary organic aerosol (SOA) as a 508 result of photochemical oxidation (in both gas and particle phases) likely contributed to the increase in volume concentration (Chen et al., 2015; Chen et al., 2009; Martin et al., 2010a; Pöschl
et al., 2010).

511 Air masses impacted by urban pollution

512 Air masses arriving at the T3 site frequently had passed over urban and industrial areas up-513 wind. When the air mass was influenced by the urban pollution, $\kappa_{\rm CCN}$ and its dispersion exhibited 514 clear diel variations (Fig. 4). The value of $\kappa_{\rm CCN}$ was lower during the night at 0.15, and it in-515 creased from the early morning hours, peaking at a value of 0.19 around noon (local time, UTC – 516 4 h). The dispersion was anti-correlated with $\kappa_{\rm CCN}$, exhibiting higher values (i.e., increased het-517 erogeneity in particle chemical composition) during night and a minimum value around noon. To a large degree, the diel trend of $\kappa_{\rm CCN}$ was due to the variation of $\kappa_{\rm org}$. The value of $\kappa_{\rm org}$ was lower 518 519 during night at 0.10 and increased to 0.16 at noon. The increase of κ_{org} is consistent with the var-520 iation of O:C, which increased during the early morning and reached the highest value of 0.8 521 around noon time. The pollution strongly affected the particle number and volume concentra-522 tions, both exhibiting similar diel trends. Under polluted conditions, particle number concentration ranged on average from 1500 to 2300 cm⁻³, an increase by a factor of ~5 from that under 523 524 background conditions. In comparison, the increase of the volume concentration was only about a factor of 2 (i.e., from a range of 0.3 - 0.75 to 1.0 to 1.3 μ m³ cm⁻³), as the urban pollution is 525 526 dominated by Aitken mode particles that make a relatively small contribution to aerosol mass 527 and volume concentration.

528 The diel variations of κ_{CCN} , its dispersion, κ_{org} , and O:C are explained as follows. During 529 night, particles in freshly emitted pollution, which are dominated by primary organic aerosol 530 (POA) and have low hygroscopicity, are mixed with more aged particles within a shallow noc-

531 turnal boundary layer (Bateman et al., 2016). In the absence of photochemical oxidation and ag-532 ing, this external mixture leads to higher dispersion of particle hygroscopicity, as well as overall 533 lower O:C and κ_{org} . As the pollution aerosols are mainly from isolated point sources, they are 534 confined in the shallow nocturnal boundary layer during night, and the residual layer above the 535 T3 site is expected to consist of aged background aerosols. Therefore, unlike under background 536 conditions, the mixing of aerosol aloft in the residual layer down to the surface cannot by itself 537 explain the increase in particle number and volume concentration during the day, both of which 538 were substantially above the background values. These increases under polluted conditions might 539 be due to the stronger urban influence at T3 during the day. The strong increase of CN concen-540 tration at 16:00 UTC (local time 12:00) could be caused by the arrival at the T3 site of the Ma-541 naus plume emitted during early morning traffic hours. To a large degree, the increases in O:C 542 and $\kappa_{\rm org}$ are due to the formation and aging of SOA in the pollution, while the development of the 543 daytime boundary layer, which leads to dilution of pollution and mixing with aged particles from 544 the residual layer, can also contribute to the increases. The condensation of secondary species 545 and photochemical aging also lead to more homogenous composition among particles (Mei et al., 546 2013a), and therefore lower dispersion of κ_{CCN} , as was observed. The O:C reached a maximum 547 average value of 0.8, similar to that under the background conditions. This suggests that the for-548 mation and photochemical aging of SOA quickly led to highly oxygenated organic compounds 549 (i.e. within several hours) (de Sá, 2017).

550 4.2.2 Dry Season Aerosol

551 Background conditions

552 Figures 5-7 show the diel variations of the aerosol properties observed during the dry season 553 for background, urban pollution, and local biomass burning air masses, respectively. During the 554 dry season, the background aerosol (Fig. 5) is strongly influenced by regional biomass burning, 555 and air masses arriving at the T3 site often pass through urban and industrial areas along the 556 Amazon River and in northeast Brazil (Andreae et al., 2015), indicating that the background aer-557 osol is also impacted by more aged urban and industrial emissions (Martin et al., 2016a). Despite different aerosol sources and processes, the particle hygroscopicity, dispersion, and κ_{org} exhibited 558 559 similar values as those of background aerosol during the wet seasons, and a lack of obvious diel 560 variations. This is also consistent with the absence of a significant diel trend of κ_{CCN} observed at 561 T0a (ATTO) during the dry season (Fig. 8). The O:C value increased slightly from 0.8 during 562 night to 0.9 in the afternoon, possibly due to further oxidation and aging of background aerosols 563 during the day time. The high value of O:C is consistent with the relatively high value of κ_{org} 564 (0.15) and is close to that observed under background condition during the wet season, indicating 565 highly oxygenated organic aerosol. The number and volume concentrations were lower just be-566 fore dawn and increased during the early morning, again a result of wet scavenging of particles 567 by fog followed by the mixing of aerosol aloft in the residual layer down to the surface as the 568 boundary layer develops in the morning.

569 Air masses impacted by urban pollution and local biomass burning

570 For urban pollution (Fig. 6) and local biomass burning (Fig. 7) air masses during the dry sea-571 son, κ_{CCN} (urban pollution: 0.12 – 0.20, local biomass burning: 0.10 – 0.17), its dispersion (urban

pollution: 0.4 – 1.0, local biomass burning: 0.4 – 0.9), κ_{org} (urban pollution: 0.10 – 0.15; local 572 573 biomass burning: 0.08 - 0.14), and O:C (polluted: 0.7 - 0.85; biomass burning: 0.7 - 0.86) 574 showed similar values and diel variations as those under polluted conditions during the wet sea-575 son. This is consistent with the picture that freshly emitted particles (in either the case of urban 576 pollution or local biomass burning) lead to overall lower O:C, κ_{org} , and higher κ_{CCN} dispersion 577 during night, followed by increases in O:C, κ_{org} , and a decrease in the dispersion during day time, 578 which are mainly driven by the formation and photochemical aging of SOA, with contributions 579 from the mixing of background aerosol aloft in the residual layer down to the surface and dilu-580 tion of fresh emission as the boundary layer develops. Compared to urban pollution, local bio-581 mass burning air masses exhibited lower κ_{CCN} and κ_{org} values during night and stronger diel vari-582 ations. In the afternoon, κ_{org} and O:C reached high values of 0.14 and 0.86, respectively, as ob-583 served for background aerosols.

584 For urban pollution air masses, aerosol number and volume concentrations showed similar 585 trends for both seasons. The increases of number and volume concentration from early morning to noon were similar, about 1000 cm⁻³ and 0.5 µm³/cm³, respectively, for both wet and dry sea-586 587 sons. The percentage increases were less pronounced in the dry season due to the higher back-588 ground values. In contrast to urban pollution, local biomass burning showed higher aerosol num-589 ber and volume concentrations at night, and decreased during the morning. Local biomass burn-590 ing activities typically peaked during evening hours, consistent with frequent classification of the 591 night time aerosol as local biomass burning (Fig. 7i) (Vestin et al., 2007). Despite the wet re-592 moval of particles by fog, the strong emission from local biomass burning, largely confined with-593 in the shallow nocturnal boundary layer, led to higher surface aerosol concentrations than those 594 in the residue layer aloft, which likely represented the regional background. As the boundary layer deepened in the morning, the mixing with aerosol from the residual layer led to decreasesin both aerosol number and volume concentration observed at the surface (Fig. 7e,f).

4.3 Hygroscopicity of PMF Factors and the Variation of Organic Hygroscopicity with Oxi-

598 **dation Level**

597

599 The hygroscopicities associated with the AMS PMF factors were estimated through multi-600 variable linear regression using different subsets of the data, as well as the entire dataset for each 601 of the two IOPs (IOP1 in Fig. 9 and IOP2 in Fig. 10). The different subsets included measure-602 ments during day, night, certain sampling periods, and ranges in particle hygroscopicity disper-603 sion. Comparison of the hygroscopicities derived from the different subsets of data allowed us to 604 examine the robustness of this approach. Uncertainty in the derived κ for individual factors was 605 determined by the number of points available to fit in the time series, with greater data coverage 606 and therefore lower uncertainty during the dry season. For the wet season (IOP1), the hygrosco-607 picity associated with PMF factors derived using different subsets of the data are largely in 608 agreement with those derived from the entire dataset. There are notable differences between the 609 hygroscopicities of MO-OOA and Fac91 factors derived using data under background conditions 610 only and those derived using the entire dataset. Such difference could be partially due to the lim-611 ited data under the background conditions during IOP1 (Fig. 3). For the dry season (IOP2), the 612 hygroscopicities of PMF factors derived using measurements under background conditions or 613 data with hygroscopicity dispersion less than 0.4 are quite different from those derived using 614 other data subsets and the entire dry season dataset. The agreement among the PMF factor hy-615 groscopicities derived using different sub datasets during the wet season and the disagreements 616 for the dry season are attributed to the applicability of the underlying assumption that the bulk 617 volume fractions of PMF factors (i.e., derived from MS mode data) represented those at the sizes 618 of CCN measurements. For the wet season, the average f_{44} was largely independent of particle 619 size from 130 to 400 nm (Fig. S10), the size range that dominated bulk aerosol mass concentra-620 tion measured by AMS. This is consistent with the assumption that the bulk volume fractions of 621 the PMF factors represent those at the two CCN sizes (142 and 171 nm). For the dry season, the 622 f_{44} averaged over local biomass burning air masses and the entire IOP2 exhibited an increase 623 with particle diameter from 100 to 300 nm (Fig. S10). For periods with hygroscopicity dispersion 624 less than 0.4 or under background conditions, the average size distribution of f_{44} was noisier due to fewer data points. Nevertheless, the size distribution shows f_{44} was largely independent of the 625 626 particle size under these conditions, consistent with the assumption described above. In the fol-627 lowing discussion, we focus on the PMF factor hygroscopicities derived using all data during the 628 wet season, and under background conditions in the dry season.

629 The MO-OOA factors for the two IOPs exhibit very similar O:C and κ values. The O:C val-630 ues were 1.19 and 1.24, and κ values were 0.20 and 0.21 for IOP1 and 2, respectively (Table 1). 631 The O:C and κ values are consistent with those of some typical SOA compounds, such as malo-632 nic acid, which has an O:C value of 1.33 and a κ value of 0.23 (Kumar et al., 2003), and succinic 633 acid, which has an O:C value of 1 and a κ value of 0.23 (Hori et al., 2003). For the LO-OOA and 634 IEPOX-SOA factors, the hygroscopicities vary between the two IOPs. The κ of IEPOX-SOA were 0.18 and 0.08 during IOP1 and IOP2, respectively, and the κ of LO-OOA factor varied 635 636 from 0.12 to 0.20 between IOP 1 and IOP 2. The difference in κ may be partially due to the 637 change of O:C values of the factors derived for the two IOPs. Difference in SOA precursors and 638 therefore composition in LO-OOA (Ng et al., 2010) may also contribute to the difference in its κ 639 values between the two IOPs. The variation of IEPOX-SOA κ between the two IOPs could be a result of the different RH conditions, which may strongly influence the composition of IEPOX-SOA (Riva et al., 2016).

During the wet season, a factor with high contribution from m/z = 91 (Fac91) was identified. The Fac91 factor correlates with several tracers for anthropogenic emissions, including NO_x, benzene, toluene, trimethylbenzene (TMB) and xylenes, but not high NO_x isoprene products (e.g. methylglyceric acid) (de Sá, 2017). This factor likely represents SOA formed from aromatics emitted from urban areas, possibly combined with a mixture of freshly oxidized biogenic compounds within the urban-influenced air (de Sá, 2017). The Fac91 factor has a hygroscopicity value of 0.10, and a much lower O:C ratio of 0.328 compared to those of MO-OOA and LO-OOA.

649 Less oxidized organic factors identified by the PMF analysis were HOA of both IOPs, 650 BBOA of IOP1 and fresh and aged BBOA of IOP2. These factors represent primary OA (POA), 651 except that the aged BBOA of IOP2 likely included contributions from oxidized POA or SOA. 652 The hygroscopicity of the HOA factors was fixed as zero in the multivariate regressions. All 653 BBOA factors have a distinctive m/z = 60 peak and correlate with biomass burning traces includ-654 ing levoglucosan and vanillin (de Sá, 2017). The retrieved hygroscopicity values for the BBOA 655 factors are substantially lower than those of SOA factors, especially for the fresh BBOA factor 656 during IOP2. The extremely low hygroscopicity suggests that the fresh BBOA, likely produced 657 by local fires, behaves very similar to HOA in term of CCN activation despite a substantially 658 higher O:C.

Figures 11 and 12 show that for SOA factors, including IEPOX-SOA, LO-OOA, MO-OOA for both IOPs and Fac91 for IOP1, the κ value increases with increasing O:C, and the variation of κ with O:C agrees with the linear relationship derived from laboratory studies of SOA CCN ac-

662 tivities (Lambe et al., 2011). The low hygroscopicity of the HOA and the BBOA factors, which 663 are below the linear relationship for SOAs, are also consistent with laboratory results of POA 664 and oxidized POA (Lambe et al., 2011). Cerully et al. (2015) derived κ of LO-OOA, MO-OOA, 665 and IEPOX-SOA factors from data collected in the southeast US during Southern Oxidant and 666 Aerosol Study (SOAS). A different name, Isoprene-OA, was used for the IEPOX-SOA factor in 667 Cerully et al. (2015), as while this factor is mainly attributed to SOA formed from IEPOX up-668 take, it might not be entirely due to IEPOX (Schwantes et al., 2015; Xu et al., 2015a; Xu et al., 669 2015b). The O:C values calculated using the Improved-Ambient method (Canagaratna et al., 670 2015) are 0.59, 0.61, and 0.92 for the IEPOX-SOA, LO-OOA, and MO-OOA factors reported in 671 Cerully et al. (2015), respectively (personal communication, L. Xu and N.L. Ng). For the LO-672 OOA, MO-OOA factors reported in Cerully et al. (2015), κ and O:C values are largely consistent 673 with the linear relationship between κ and O:C derived from Lambe et al. (2011). Cerully et al. 674 (2015) also reported IEPOX-SOA (i.e., called Isoprene-OA in their study) factor κ of 0.2, simi-675 lar to 0.18 derived for IOP1. The O:C value of the IEPOX-SOA factor during the SOAS study 676 was 0.59, somewhat lower than those derived from both IOPs of GoAmazon 2014/5. While the 677 IEPOX-SOA factors identified using different datasets share many similar features, they are not 678 identical and can consist of different groups of compounds. Such differences may be due to vary-679 ing degrees of oxidation in different environment between the two field campaigns.

For comparison with earlier field studies, the values of κ_{org} and O:C were averaged according to the hours of the day over particle diameters of 142 and 171 nm for data under polluted conditions during IOP1 and all data during IOP2 (Figs. 11 and 12). For the one-hour diel averages, the slope of κ_{org} vs. O:C, derived through a bivariate least squares fit (i.e., Orthogonal distance regression), is steeper than that derived from laboratory studies of SOA hygroscopicity, especially

685 during IOP2. This steep slope during IOP2 is consistent with the results from earlier field studies 686 (Mei et al., 2013b), although there is a clear offset between the two relationships. The O:C ratios 687 from Mei et al. (2013b) and Lambe et al. (2011) were scaled by a factor of 1.27 to account for 688 changes in the method of calculating the O:C ratio (Improved-Ambient, Canagaratna et al., 2015) 689 while all O:C values from this work were calculated using the Improved-Ambient method. This 690 offset between the field studies may be partially due to the different precursors of the SOA for 691 the campaigns, with higher anthropogenic VOC fraction expected for CalNex and CARES, 692 which took place near Los Angeles and Sacramento, respectively. In addition, biomass burning 693 represented a much smaller fraction of the organics during CalNex and CARES (Mei et al., 694 2013a; Mei et al., 2013b). The factors associated with secondary processes (e.g., MO-OOA, LO-695 OOA, and IEPOX-SOA), which have higher O:C values, exhibited higher volume fraction dur-696 ing the day, whereas the factors associated with primary emissions (e.g. HOA and BBOA), 697 which have lower O:C, had higher volume fractions during the night (de Sá, 2017). As a result, 698 the diel trend of overall O:C was to a large degree driven by the variations in volume fractions of 699 the POA and SOA factors with very different O:C values. This is in contrast to laboratory stud-700 ies, in which the increase of O:C was mainly driven by oxidation. As POA exhibits hygrosco-701 picity values well below the linear fit between SOA hygroscopicity and O:C, mixtures with dif-702 ferent POA and SOA fractions lead to a steeper slope for the increase of κ_{org} with O:C, as shown 703 by the results from this and previous field studies (Mei et al., 2013b).

704 **5 Conclusions**

Size-resolved CCN spectra at five particle diameters ranging from 75 to 171 nm were
characterized down-wind of Manaus, Brazil, in central Amazonia for a period of one year from
March 12, 2014 to March 3, 2015 during GoAmazon2014/5. For each season, the air masses ar-

708 riving at the site were classified into different types, including background, urban pollution, and 709 local biomass burning. During the wet season, the background air mass represented near natural 710 conditions, at times with impact from anthropogenic emissions, while in the dry season, the 711 background was dominated by regional and long-distance biomass burning aerosol particles. Pol-712 luted air masses represented those with strong influence from urban emissions, which were most-713 ly from Manaus. The local-biomass-burning type describes those strongly influenced by local 714 (i.e., fresh) biomass mass burning activities that dominate over the impact from urban pollution, 715 if any.

716 Particle hygroscopicity (κ_{CCN}), mixing state, and organic hygroscopicity (κ_{org}) were derived 717 from size-resolved CCN activation fraction and concurrent aerosol composition measurements. 718 The monthly mean κ_{CCN} exhibits the lowest values during the dry season, largely due to lower $\kappa_{\rm org}$ when aerosol was often strongly influenced by local biomass burning. The $\kappa_{\rm CCN}$ increased 719 720 with particle size during all seasons, consistent with decreasing organic volume fraction with in-721 creasing particle size. Under background conditions, the value of κ_{CCN} and its size dependence 722 were largely consistent among different seasons, despite the very different aerosol sources. Dur-723 ing the dry season, aerosols classified as urban pollution and local biomass burning exhibited 724 lower κ_{org} values compared to background aerosols, contributing to the lower values of overall 725 $\kappa_{\rm CCN}$.

Under background conditions during both wet and dry seasons, the largely constant diel trends of κ_{CCN} and κ_{org} suggest little variation in particle composition throughout the day. The constant κ_{org} of ~ 0.15 is consistent with the lack of a diel trend in f_{44} and O:C. The high values of f_{44} and O:C indicate that the aerosols under background conditions are dominated by the aged regional aerosol particles consisting of highly oxygenated organic compounds. When the air 731 mass is influenced by urban pollution or local biomass burning, κ_{CCN} , κ_{org} , f_{44} , and O:C exhibit 732 clear diel variations. The value of κ_{CCN} (0.1 - 0.2) is lower during the night, and increases from 733 the early morning hours, peaking around noon (local time, UTC - 4 h). This diel trend of κ_{CCN} is 734 largely driven by the variation in κ_{org} (0.08-0.15), consistent with the variation of O:C. The dis-735 persion of $\kappa_{\rm CCN}$ is anti-correlated with $\kappa_{\rm CCN}$, exhibiting higher values during night and a mini-736 mum value around noon, indicating an increased heterogeneity in particle chemical composition 737 during night time. These diel variations for air masses strongly influenced by urban pollution and 738 local biomass burning indicate that during night, freshly emitted particles, dominated by POA 739 and with low hygroscopicity, are mixed with more aged particles within a shallow nocturnal 740 boundary layer. In the absence of photochemical oxidation and aging, this external mixture leads 741 to higher dispersion of particle hygroscopicity, as well as overall lower O:C and κ_{org} . The in-742 creases in O:C and $\kappa_{\rm org}$ during daytime are driven by the formation and aging of SOA and dilu-743 tion of POA emissions into a deeper boundary layer, while the development of the boundary lay-744 er, which leads to mixing with aged particles from the residual layer, likley also contributes to 745 the increases.

746 The hygroscopicities associated with individual PMF organic factors were derived through 747 multi-variable linear regression. For the SOA factors, κ increases within increasing O:C, and the 748 variation of κ with O:C agrees well with the linear relationship derived from laboratory studies of 749 SOA hygroscopicity (Lambe et al., 2011). The low hygroscopicity of HOA and the BBOA fac-750 tors, which are below the linear relationship, are also consistent with laboratory results of POA 751 and oxidized POA (Lambe et al., 2011). In contrast, the slope of κ_{org} (i.e., overall organic hygro-752 scopicity) vs O:C is much steeper when compared to that derived from laboratory studies of 753 SOA hygroscopicity, especially for IOP2. Such difference is because the increase of O:C was

754	driven primarily by oxidation in laboratory SOA studies, while the variation in O:C of ambient
755	organics is to a large degree due to the variations in volume fractions of POA and SOA factors,
756	which have very different O:C values. As POA factors show hygroscopicity values well below
757	the linear fit between SOA hygroscopicity and O:C, mixtures with different POA and SOA frac-
758	tions lead to a steeper slope for the increase of κ_{org} with O:C, as shown by the results from this
759	and earlier field studies (Mei et al., 2013).

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1194 **Tables**

Table 1: Density, O:C ratio and hygroscopicity associated with organic factors derived from Positive Matrix Factorization (PMF) analysis. For IOP1 and IOP2, hygroscopicities of PMF organic
factors were derived from time series of particle hygroscopicity under all conditions and background conditions, respectively.

IOP1 (wet season)				IOP2 (dry season)			
PMF Factor	ρ (g cm ⁻³)	O:C	К	PMF Factor	ρ (g cm ⁻³)	O:C	κ (Bkgd)*
IEPOX-SOA	1.47	0.798	0.18±0.02	IEPOX-SOA	1.42	0.711	0.08±0.03
MO-OOA	1.80	1.19	0.20±0.02	MO-OOA	1.81	1.24	0.21±0.03
LO-00A	1.48	0.786	0.12±0.02	LO-OOA	1.52	0.883	0.20±0.03
BBOA	1.42	0.712	0.04±0.03	Aged BBOA	1.37	0.666	0.08±0.03
Fac91	1.14	0.328	0.10±0.03	Fresh BBOA	1.23	0.536	0.00 ± 0.07
НОА	0.95	0.163	0	НОА	1.02	0.223	0

1199 * κ (Bkgd) refers to κ values of PMF factors derived from the time series of particle hygroscopicity under background conditions 1200 (see Sections 4.2.2 and 4.3).

1201 Figures

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Figure 1: Seasonal variations of aerosol properties observed at the T3 site from March 2014 to March 2015, including (a) κ_{CCN} , (b) κ_{org} , and size resolved volume fraction of (c) organics, (d) sulfate, including (NH₄)₂SO₄ and NH₄HSO₄, (e) nitrate, and (f) refractory Black Carbon. Data points are the monthly mean; error bars represent the 25th and 75th percentiles.

1208



Figure 2: The variation of $\kappa_{\rm CCN}$ and $\kappa_{\rm org}$ with particle diameter during different seasons for each of three air mass types. Data points are the mean values; error bars are the 25 and 75 percentiles. The top left panel also includes κ_{CCN} observed under near natural conditions during the AMAZE-08 campaign at T0t in the wet season (Gunthe et al., 2009) and during the one-year period from March 2014 to February 2015 at the background T0a site (Pöhlker et al., 2016). The top right panel includes κ derived from particle growth factor measurements in July 2001, during a "recent biomass burning period" of the CLAIRE-2001 study (Rissler et al., 2004) and from 11 Septem-ber to 8 October 2002, during the dry period of the LBA-SMOCC (Rissler et al., 2006).



1221 1222 Figure 3: Diel variations of aerosol properties and meteorological parameter under background conditions during the wet season, including (a) $\kappa_{\rm CCN}$, (b) $\sigma_{\kappa_{\rm CCN}}$, (c) fraction of the organic 1223 1224 mass at m/z = 44 (f_{44}) and the elemental ratio O:C, (d) κ_{org} derived using size resolved particle 1225 composition, (e) the total number of condensation nuclei $(N_{\rm CN})$, (f) the total aerosol volume de-1226 rived from size distribution measured by the SMPS in MAOS, (g) planetary boundary layer 1227 height as estimated using the ceilometer data, (h) the volume fractions of aerosol species, and (i) 1228 the number of samples in each hour bin corresponding to the data by the same colors and symbols in their respective panel. The values of κ_{CCN} , $\sigma_{\kappa_{\text{CCN}}}$, κ_{org} , and volume fraction of aero-1229 sol species were averaged over three particle diameters of 112, 142 and 171 nm. The values of f_{44} 1230 1231 and O:C were derived from the AMS bulk measurements. Data include the last two weeks of March 2014 when valid data from both size-resolved CCN system and AMS were available. Da-1232 1233 ta points are hourly averaged mean values; error bars represent the 25 and 75 percentiles of the 1234 data. Yellow shading represents the local daytime (10:00 - 22:00 UTC).



1235Wet Season - Polluted Conditions1236Figure 4: Diel variations of aerosol properties and meteorological parameters for urban pollution

1237 air masses during the wet season (analogous to Fig. 3).



Figure 5: Diel variations of aerosol properties and meteorological parameters under backgroundconditions during the dry season (analogous to Fig. 3).





1246 Figure 6: Diel variations of aerosol properties and meteorological parameters for urban pollution

1247 air masses during the dry season (analogous to Fig. 3).



Figure 7: Diel variations of aerosol properties and meteorological parameters for local biomassburning air masses during the dry season (analogous to Fig. 3).



Figure 8: Comparison κ_{CCN} values derived from measurements at the T0a (ATTO) site (Pöhlker et al., 2016) and at the T3 site under background conditions (this study) during the (a) wet season (April and May 2014) and (b) dry season. The size resolved CCN data at T0a was collected by stepping the particle size at given CCNC supersaturations (Rose et al., 2008b). Data displayed for T0a is averaged over critical particle diameters ranging from 44 to 175 nm, while the T3 data is averaged from measurements at 112, 142, and 171 nm.

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Figure 9: Hygroscopicity of AMS PMF factors for IOP1 (i.e., wet season) retrieved by multilinear regressions using all data (red square), data from UTC 12:00-24:00 (cyan circle), data from UTC 0:00-12:00 (blue triangle), data under background conditions (red triangle), data under polluted conditions (red diamond), and data with hygroscopicity dispersion $\sigma_{\kappa_{CCN}} / \bar{\kappa}_{CCN}$ less than 0.6 (black diamond).



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Figure 10: Hygroscopicity of PMF factors for IOP2 (i.e., dry season) retrieved by multi-linear regressions using all data (red square), data from UTC 12:00-24:00 (cyan circle), data from UTC 0:00-12:00 (blue triangle), data with strong influence from local biomass burning (red right triangle), data under background conditions (red upside-down triangle), data from Aug 21 to Sep 14, 2014 only (red left pointing triangle), data from Sep 15 to Oct 15 only (brown diamond), and data with a dispersion ($\sigma_{\kappa_{CCN}} / \bar{\kappa}_{CCN}$) < 0.4 (black diamond).



Figure 11: The variation of PMF factor hygroscopicity, 1-hour diel average of organic hygrosco-picity and O:C ratio at 142 and 171 nm for urban pollution air masses. Also shown are the rela-

1281 tionships between κ_{org} and O:C reported by earlier field and laboratory studies.



Figure 12: The variation of PMF factor hygroscopicity, 1-hour diel average of organic hygroscopicity, and O:C ratio at 142 and 171 nm for local biomass burning air masses. Also shown are the relationships between κ_{org} and O:C reported by earlier field and laboratory studies.