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2 **Effects of transport on a biomass burning plume from Indochina**
3 **during EMeRGe-Asia identified by WRF-Chem**

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37 **Abstract.**

38 The Indochina biomass burning (BB) season in springtime has a substantial
39 environmental impact on the surrounding areas in Asia. In this study, we evaluated the
40 environmental impact of a major long-range BB transport event on 19 March 2018 (a
41 flight of the HALO research aircraft, flight F0319) preceded by a minor event on 17
42 March 2018 (flight F0317). Aircraft data obtained during the campaign in Asia of the
43 Effect of Megacities on the transport and transformation of pollutants on the Regional
44 to Global scales (EMeRGe) were available between 12 March and 7 April 2018. In the
45 F0319, results of 1-min mean carbon monoxide (CO), ozone (O₃), acetone (ACE),
46 acetonitrile (ACN), organic aerosol (OA) and black carbon aerosol (BC) concentrations
47 were up to 312.0 ppb, 79.0 ppb, 3.0 ppb, 0.6 ppb, 6.4 μg m⁻³, 2.5 μg m⁻³ respectively,
48 during the flight, which passed through the BB plume transport layer (BPTL) between
49 the elevation of 2000–4000 m over the East China Sea (ECS). During F0319, CO, O₃,
50 ACE, ACN, OA and BC maximum of the 1 minute average concentrations were higher
51 in the BPTL by 109.0 ppb, 8.0 ppb, 1.0 ppb, 0.3 ppb, 3.0 μg m⁻³ and 1.3 μg m⁻³
52 compared to flight F0317, respectively. Sulfate aerosol, rather than OA, showed the
53 highest concentration at low altitudes (<1000 m) in both flights F0317 and F0319
54 resulting from the continental outflow in the ECS.

55 The transport of BB aerosols from Indochina and its impacts on the downstream
56 area was evaluated using a WRF-Chem model. The modeling results tended to
57 overestimate the concentration of the species, with examples being CO (64 ppb), OA
58 (0.3 μg m⁻³), BC (0.2 μg m⁻³) and O₃ (12.5 ppb) in the BPTL. Over the ECS, the
59 simulated BB contribution demonstrated an increasing trend from the lowest values on
60 17 March 2018 to the highest values on 18 and 19 March 2018 for CO, fine particulate
61 matter (PM_{2.5}), OA, BC, hydroxyl radicals (OH), nitrogen oxides (NO_x), total reactive
62 nitrogen (NO_y), and O₃; by contrast, the variation of J(O¹D) decreased as the BB

63 plume's contribution increased over the ECS. In the low boundary layer (<1000 m), the
64 BB plume's contribution to most species in the remote downstream areas was <20 %.
65 However, at the BPTL, the contribution of the long-range transported BB plume was as
66 high as 30–80 % for most of the species (NO_y , NO_x , $\text{PM}_{2.5}$, BC, OH, O_3 , and CO) over
67 South China (SC), Taiwan, and the ECS. BB aerosols were identified as a potential
68 source of cloud condensation nuclei, and the simulation results indicated that the
69 transported BB plume had an effect on cloud water formation over SC and the ECS on
70 19 March 2018. The combination of BB aerosol enhancement with cloud water resulted
71 in a reduction of incoming shortwave radiation at the surface in SC and the ECS which
72 potentially has significant regional climate implications.

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75 **1 Introduction**

76 Biomass burning (BB) is one of the main sources of aerosols, greenhouse gases, and air
77 pollutants (e.g. Ramanathan et al., 2007; Lin et al., 2009; 2014; Tang, 2003; Carmichael
78 et al., 2003; Chi et al., 2010; Fu et al., 2012; Lin N.H. et al., 2012; Chuang et al., 2016).
79 Reid et al. (2013) and Giglio et al. (2013) investigated the seasonal aerosol optical depth
80 over Southeast Asia and have indicated that Indochina is a major contributor of carbon
81 emissions in springtime. Galanter et al. (2000) estimated that BB accounts for 15–30 %
82 of the entire tropospheric CO background. Huang et al. (2013) indicated that the
83 contribution of BB in Southeast Asia to the aerosol optical depth (AOD) in Hong Kong
84 and Taiwan could be in the range of 26–62 %. Moreover, BB emissions over Indochina
85 are a significant contributor to black carbon (BC), organic carbon (OC), and O_3 in East
86 Asia (Lin et al., 2014). In their BB modeling study, Lin et al. (2014) identified a
87 northeast (NE) to southwest (SW) zone stretching from South China (SC) to Taiwan
88 with a reduction in shortwave radiation of approximately 20 W m^{-2} at the ground

89 surface. In addition, the total carbon emission from BB in Southeast Asia is
90 approximately 91 Tg C yr⁻¹, accounting for 4.9 % of the global total (Yadav et al.,
91 2017). According to Xu et al. (2018), BB in Indochina leads to BC production at high
92 concentrations of up to 2–6 μg m⁻³ in spring. The authors reported that BC particles
93 were transported to the glaciers in the Tibetan Plateau, where it significantly affected
94 the melting of the snow, causing some severe environmental problems, such as water
95 resource depletion. Ding et al. (2021) indicated that BB aloft aerosols strongly increase
96 the low cloud coverage over both land and ocean and affect the monsoon in the
97 subtropical Southeast Asia.

98 Although many researchers have indicated the importance of BB emissions, their
99 precise estimation and applying in the modeling study remains challenging (Fu et al.
100 2012; Huang et al. 2013; Pimonstree et al. 2018; Marvin et al. 2021). For example,
101 Heald et al. (2003) conducted an emission inventory in Southeast Asia and reported that
102 the uncertainties of BB emission estimations could be a factor of three or even higher.
103 Following an inverse model analysis, Palmer et al. (2003) also indicated the
104 overestimation of regional BB emissions over Indochina. Shi and Yamaguchi (2014)
105 pointed out BB emissions exhibited strong temporal interannual variability between
106 2001 and 2010 over southeast Asia. Satellite data can be used to easily locate hotspots
107 such as those where agricultural residuals burning and forest wildfires are occurring
108 worldwide. However, accurately quantifying the amount of BB emission from satellite
109 data is difficult because anthropogenic pollutants and BB emissions are typically mixed
110 in the atmosphere. During the NASA Transport and Chemical Evolution over the
111 Pacific (TRACE-P) aircraft mission in spring 2001, Jacob et al. (2003) observed that
112 warm conveyor belts (WCBs) lift both anthropogenic and BB (from SE Asia) air
113 pollution to the free troposphere, resulting in complex chemical signatures.
114 Wiedinmyer et al. (2011) demonstrated that the uncertainty of emission estimation

115 could be as high as a factor of 2 because of the error introduced by estimates in fire
116 hotspots, area burned, land cover maps, biomass consumption, and emission factors in
117 the model. In this context, Lin et al. (2014) highlighted the uncertainty of emission
118 estimation in the first version of Fire Inventory from NCAR (Wiedinmyer et al., 2011).

119 The transport of BB pollution is strongly dependent on the atmospheric structure
120 and weather conditions. Tang et al. (2003) noted that most BB aerosols, having their
121 source in Indochina (mainly south of 25 °N and be alofted to an altitude of 2000–4000
122 m) during the TRACE-P campaign were associated with outflow in the WCB region
123 after frontal passage. Lin et al. (2009) suggested a mountain lee-side troughs as an
124 important mechanism, resulting in BB product transport from the surface to >3000 m.
125 BB pollution is often transported from its sources to the East China Sea (ECS), Taiwan,
126 and the western North Pacific within a few days.

127 The airborne field experiment EMeRGe (Effect of Megacities on the transport and
128 transformation of pollutants on the Regional to Global scales) over Asia was led by the
129 University of Bremen, Germany and conducted in collaboration with Academia Sinica,
130 during the inter-monsoon period in 2018 ([http://www.iup.uni-](http://www.iup.uni-bremen.de/emerge/home/home.html)
131 [bremen.de/emerge/home/home.html](http://www.iup.uni-bremen.de/emerge/home/home.html)). The EMeRGe aircraft mission consists of two
132 parts. The first mission phase was conducted in Germany in July 2017 and the second
133 phase was conducted from Taiwan in 2018 (Andrés Hernández et al. 2022).EMeRGe in
134 Asia aimed at the investigation of the long range transport (LRT) of local and regional
135 pollution originating in Asian major population centers (MPCs) from the Asian
136 continent into the Pacific. A central part of the project was the airborne measurement
137 of pollution plumes on-board of the High Altitude and Long Range Research Aircraft
138 (HALO). The HALO platform was based in Tainan, Taiwan (Fig. 1a-b), and made
139 optimized transects and vertical profiling in regions north or south of Taiwan,
140 dependent on the relevant weather and emission conditions. HALO measurements

141 additionally provide important information for the evaluation of the LRT of BB
142 emissions and its potential environmental impact in East Asia between 12 March and 7
143 April 2018. During the EMERGE-Asia campaign, HALO carried out 12 mission flights
144 in Asia and 4 transfer flights from Europe to Asia with a total of 110 flight hours.

145 This paper is organized as follows: the model configuration and BB emission
146 analysis employed in the model simulation are described in Section 2, and the weather
147 conditions and HALO measurement results are presented in Section 3. The model
148 performance, as well as the evaluation of BB product transport and effects on East Asia
149 selected regions are discussed in Sections 4 and 5, respectively.

150

151 **2 Aircraft data and Model configuration**

152 **2.1 HALO aircraft data**

153 The HALO aircraft was equipped with a number of instruments and a detailed
154 description of the measurement systems onboard the HALO was presented in Andrés
155 Hernández et al.(2022). In this study, aerosol data (OA, BC, SO_4^{2-} , NO_3^- , NH_4^+), and
156 trace gases such as CO, SO_2 , O_3 , NO_x , NO_y , acetone (ACE), acetonitrile (ACN), HCHO,
157 HONO, and photolysis rate $J(\text{O}^1\text{D})$, $J(\text{NO}_2)$ were employed in the analysis.

158 **2.2 WRF-Chem Model and model configuration**

159 We used the Weather Research Forecasting with Chemistry (WRF-Chem) model (Ver.
160 4.1.1) (Grell et al., 2005; Powers et al. 2017) to study the LRT of air masses associated
161 with BB pollutants in Indochina. The initial and boundary meteorological conditions
162 for WRF-Chem were obtained from National Centers for Environmental Prediction
163 (NCEP)-GDAS Global Analysis data sets at 6-h intervals. The Mellor–Yamada–Janjic
164 planetary boundary layer scheme (Janjic, 1994) was applied. The horizontal resolution
165 for the simulations performed was 10 km, and the grid box had 442×391 points in the
166 east–west and north–south directions (Fig. 1a). A total of 41 vertical levels were

167 included, with the lowest level at an elevation of approximately 50 m. To improve the
168 accuracy of the meteorological fields, a grid nudging four-dimensional data
169 assimilation scheme was applied using the NCEP-GDAS Global Analysis data.

170 The cloud microphysics used followed the Lin scheme (Morrison et al., 2005). The
171 rapid radiative transfer model (Zhao et al., 2011) was used for both longwave and
172 shortwave radiation schemes. Moreover, land surface processes are simulated using the
173 Noah-LSM scheme (Hong et al., 2009). In terms of transport processes, we considered
174 advection by winds, convection by clouds, and diffusion by turbulent mixing. The
175 removal processes in this study were gravitational settling, surface deposition, and wet
176 deposition (scavenging in convective updrafts and rainout or washout in large-scale
177 precipitation). The kinetic preprocessor (KPP) interface was used in both of the
178 chemistry schemes of the Regional Atmospheric Chemistry Mechanism (RACM,
179 Stockwell et al., 1990). The secondary organic aerosol formation module, the Modal
180 Aerosol Dynamics Model for Europe (Ackermann et al., 1998)/Volatility Basis Set
181 (Ahmadov et al., 2012), was also employed in the WRF-Chem model. In RACM, the
182 “KET” represents acetone and higher saturated ketones (KET) (Stockwell et al. 1997).
183 According to Singh et al. (1994), BB and the primary anthropogenic emissions could
184 contribute 26% and 3%, respectively, to the atmospheric acetone sources. The model
185 configuration and physics and chemistry options are listed in Table 1.

186

187 **2.3 Emission Inventories**

188 Anthropogenic emissions, such as NO_x, CO, SO₂, nonmethane volatile organic
189 compounds, sulfate, nitrate, PM₁₀, and PM_{2.5}, were adopted on the basis of the emission
190 inventory in Asia – MICS-Asia III which is the year in 2010 (Li et al., 2020; Kong et
191 al., 2020). For BB emissions FINNv1.5 (<https://www.acom.ucar.edu/Data/fire/>) was
192 employed. FINN provided daily, 1000 m resolution, global estimates of the trace gas

193 and particle emissions from open BB, which included wildfires, agricultural fires, and
194 prescribed burning but not biofuel use and trash burning (Wiedinmyer et al., 2011). The
195 anthropogenic emissions in Taiwan were obtained from the Taiwan Emission Data
196 System (TEDS) which is the emission inventory of the air-pollutant monitoring
197 database of the Taiwan Environmental Protection Administration. The TEDS version
198 used for this study was V9.0 (2013) and contained data on eight primary atmospheric
199 pollutants: CO, NO, NO₂, NO_x, O₃, PM₁₀, PM_{2.5}, and SO₂.

200

201 **3 Characteristics of the field experiment**

202 **3.1 MODIS Aerosol optical depth and Weather conditions**

203 Figures 2a and b visualizes the numerous fire hotspots and high aerosol optical depth
204 on 17 March 2018 registered by the MODIS satellite. Indeed, a large number of BB fire
205 hotspots frequently occurred over Indochina during the springtime (Lin et al. 2009;
206 2014) and EMeRGe-Asia campaign (Supplementary Figure S1). On 17 March 2018 at
207 06:00 UTC (14:00 LT; LT = UTC+8:00) the weather data indicated a series of high-
208 pressure systems in northern China and a separate high-pressure system over the Japan
209 sea (Fig. 2c). At 1000 hPa, a strong northerly continental outflow was identified over
210 southern Japan, the ECS, and Taiwan (Fig. 2d). On 19 March 2018, a new frontal
211 system was located from Korea to the Guangdong province in SC (Fig. 2e). On the
212 same day at 06:00 UTC, a discontinued flow was identified at the frontal zone to the
213 north of Taiwan in the ECS (Fig. 2f). In other words, Taiwan was located at the
214 prefrontal and warm conveyor area due to the surrounding southerly flow on 19 March
215 2018 at 06:00 UTC (Figs. 2e and 2f, respectively). The southerly wind was gradually
216 replaced by the northeasterly after another frontal passage on 20 March 2018 at 00:00
217 UTC (data not shown).

218 In the upper layer (700 hPa; Figs. 2g–2j), the flow pattern differed from that at the

219 near-ground surface (1000 hPa; Figs. 2d and 2f). A southwesterly strong wind, coming
220 from the east side of the Tibetan Plateau in SC, moving to the North Eats i.e. Korea, is
221 converted to a polar front wave flow in northeastern China and Korea on 17 March
222 2018 (Fig. 2g). This high-elevation northward strong wind belt distribution at 700 hPa
223 was associated with a corresponding lee-side trough at the east of the Tibetan Plateau,
224 whereas a ridge was noted over the east coast of China on the same day (Fig. 2h).
225 Consistent with the mechanism reported by Lin et al. (2009), once a significant lee-side
226 trough formed, it provided favorable conditions for the upward motion over the lee-side
227 of the Tibetan Plateau and brought BB emission to the free troposphere layer following
228 the strong wind belt transport to the downwind area. After the weather system moved
229 to the east, the north–south trough turned to SW–NE such that the strong wind belt was
230 in an approximately SW–NE direction and located between 20 and 30 °N on 19 March
231 2018 (Figs. 2i and 2j). In conclusion, the Indochina BB pollutants were driven by the
232 strong wind belt from Indochina, northward to SC on 17 March 2018 and then eastward
233 passing over Taiwan between 20 and 30 °N to the south of Japan on 19 March 2018.

234 **3.2 Characteristics of LRT BB to the ECS by WRF-Chem model**

235 Figure 3 shows latitude longitude plots of the simulated CO concentration
236 differences with and without BB emission at an elevation of 1000 m (Fig. 3a), mainly
237 in Indochina, SC, and the South China Sea on 17 March 2018. The ambient flow was
238 easterly and then northward from the South China Sea to SC at 1000 m elevation
239 between 00:00 and 12:00 UTC on 17 March 2018 (Fig. 3a-b). The BB plume
240 accumulated and persisted for an extended period in the lower part of the boundary
241 layer on 17 and 19 March 2018 (Figs. 3a-b, and 3e-f). In contrast, the high CO
242 concentration followed the southwesterly or westerly strong wind belt (Figs. 3c-d, and
243 3g-h) and its weather conditions (Fig. 2) at an elevation of 3000-m (700 hPa). Following
244 the movement of the ridge and trough at the 700 hPa geopotential height (Fig. 2h and

245 2j), the associated strong wind belt turned to move eastward in the SW–NE direction
246 between 17 and 19 March 2018. The BB plume transport over Indochina was affected
247 by a fast-moving strong flow at 700 hPa (Fig. 2g and 2i), shifting the plume toward
248 Taiwan and the ECS, during 17–19 March 2018. The highest CO concentration
249 contributed by the BB plume was >150 ppb, originally sourced from Indochina, and it
250 was mainly transported northward on 17 March 2018 (Figs. 3c-d) and then covered a
251 large area in East Asia at a CO concentration of >100 ppb on 19 March 2018 (Figs. 3g-
252 h). Figure 4 indicates simulation differences for the contribution of BB along an E–W
253 cross-section at 30 °N at 16:00 UTC on 18 March 2018 (Fig. 4a) and 06:00 UTC on 19
254 March 2018 (Fig. 4b). We noted that a strong wind at 2000 m elevation and a high CO
255 concentration (>70 ppb) due to BB at the BPTL. Moreover, the CO concentration
256 attributed to BB was low at the elevation of >4000 m on 19 March at 06:00 UTC (Fig.
257 4b), showing that the BB pollutants mainly affect altitudes below 4000 m.

258 **3.3 Aircraft measurements**

259 Two HALO flights were scheduled to the ECS to measure the pollutants following the
260 continental outflow; the flights departed on 17 (Fig. 5a) and 19 (Fig. 6a) March 2018
261 and followed similar tracks. To indicate the measurement results along the flight path,
262 the 1-min average data is shown in Figures 5b and 6b. On 17 March 2018, the flight
263 departed from Tainan (Fig. 1b) at 01:09 UTC (09:09 LT) first southbound and then
264 northward to the ECS (Fig. 5a). The elevation for sample collection was mainly <4000
265 m, where the CO concentration was found to be <200 ppb in most cases on that day
266 (Fig. 5b). At elevations between 2000 and 4000 m, the concentration of the major
267 aerosol components (i.e., OA, BC, SO_4^{2-} , NO_3^- , and NH_4^+) was mostly <2 $\mu\text{g m}^{-3}$,
268 except just above western Taiwan after 08:00 UTC (Figs. 5a–5d). The peak
269 concentrations for OA, BC, SO_4^{2-} , NH_4^+ , and NO_3^- were 3.4, 1.2, 2.1, and 0.7 $\mu\text{g m}^{-3}$,
270 respectively, at the altitude between 2000 and 4000 m. SO_4^{2-} demonstrated the highest

271 concentration among the aerosol components, especially during 04:00–04:37 and
272 05:48–06:15 UTC (peaking at $5.1 \mu\text{g m}^{-3}$) when the flight was north of 30°N and an
273 elevation of $<1000 \text{ m}$ (Figs. 5a–5c). This result could be attributed to anthropogenic
274 pollution from the continental outflow (Lin et al. 2012) or probably part from Japan
275 contributed to the high sulfate concentration in the boundary layer over the ECS. As for
276 the trace gases such as ACE, ACN and O_3 , their concentrations between 2000 and 4000
277 m were ranging between 1–2 ppb, 0.1–0.3 ppb, and 60–70 ppb (Fig. 5b), respectively,
278 implying minor influence over the ECS by the BB plume in this flight. Figure 5e
279 illustrates the HYSPLIT (Stein et al., 2021) 96-h backward trajectories, which
280 identified the air mass origin starting at 02:00 UTC, followed by 04:00, 06:00, and
281 09:00 UTC. The continental outflow contributed to higher sulfate concentrations ($3\text{--}5$
282 $\mu\text{g m}^{-3}$ at 33°N) at 04:00 and 06:00 UTC (Figs. 5b, 5c, and 5e) at $<1000 \text{ m}$ along the
283 flight path. In contrast, south of 25°N and above Taiwan, the local pollution and
284 continental outflow are dominating sources on 17 March 2018.

285 The HALO flight on 19 March 2018 departed at 00:19 UTC (08:19 LT). It was
286 bound northward and sampled air at an altitude of $<4000 \text{ m}$ most of the time, as shown
287 in Figures 6a and 6b. Figures 6c and 6d indicate the latitude-height variation of SO_4^{2-}
288 and OA mass concentrations along the flight path on 19 March 2018. As the flight left
289 Taiwan, it maintained an elevation of 3000 m during 01:00–02:00 UTC (Fig. 6a, 121--
290 126°E) and then descended to $<1000 \text{ m}$ during 02:00–02:40 UTC (Fig. 6b). The OA
291 mass concentration was higher at 3000 m than at the low altitude during 01:00–03:00
292 UTC (Figs. 6b and 6d). In particular, CO, OA and BC exhibited a substantial peak
293 concentration of 312 ppb, $6.4 \mu\text{g m}^{-3}$ and $2.5 \mu\text{g m}^{-3}$ at 01:54 and 02:51 UTC at 26°N ,
294 $125\text{--}126^\circ\text{E}$, and an altitude of $2000\text{--}4000 \text{ m}$, where a BPTL was observed. The trace
295 gases such as ACE, ACN, and even O_3 (Fig. 6b) have consistent peak times in the BPTL
296 with concentrations of 3.0 ppb, 0.6 ppb, and 79 ppb, respectively. In this flight, SO_4^{2-}

297 had the second-highest concentration among the aerosol components ($1\text{--}2.4\ \mu\text{g m}^{-3}$;
298 Figs. 6b and 6c) upstream of Taiwan ($25\text{--}27\ ^\circ\text{N}$) during 1:00–3:00 UTC.

299 In the northern part of the flight between 03:00 and 05:00 UTC at an elevation of
300 $>3000\ \text{m}$, the aerosol component concentrations were all at their lowest level (Figs. 6b–
301 6d). During 05:00–07:00 UTC, the HALO aircraft flew back southward to $25\ ^\circ\text{N}$, where
302 high OA mass concentrations appeared again between 2000 and 4000 m (Figs. 6a, 6b,
303 and 6d). Sulfate was the species with the highest concentration between 05:30 and
304 06:30 UTC (Figs. 6b and 6c) when the flight's elevation was $<1000\ \text{m}$ in the lower
305 boundary between 25 and $27\ ^\circ\text{N}$ (upstream of Taiwan). The reason explaining this
306 observation is that the transport of anthropogenic pollutants of continental origin takes
307 place mainly in the boundary layer (Figs. 6b–6d). Other aerosol species, such as NO_3^-
308 and NH_4^+ , demonstrated low concentrations, except when the elevation was $<1000\ \text{m}$,
309 where they ranged up to $1\ \mu\text{g m}^{-3}$ (Fig. 6b).

310 The 96-h HYSPLIT backward trajectory starting from the flight locations at
311 02:00–07:00 UTC (Fig. 6e) indicated that the air masses at elevations between 2000
312 and 4000 m were potentially transported from Indochina. North of $30\ ^\circ\text{N}$ and at altitudes
313 of $>3000\ \text{m}$ at 04:00 UTC, the concentrations of air pollutants (including OA, SO_4^{2-} ,
314 NO_3^- , and NH_4^+) were low (Figs. 6b and 6e) even though the air mass in the low
315 boundary was sourced from SC and the Taiwan Strait. In general, the BPTL was mainly
316 located south of $30\ ^\circ\text{N}$ as presented by Carmichael et al. (2003), and Tang et al. (2003).
317 However, the ACN still could be around 300ppt or less as the flight at the north of $30\ ^\circ\text{N}$
318 (during 3:30–4:30 UTC) and could be recognized as the contribution of BB (Förster et al.
319 2022). In other words, it might still have BB products being transported to the north of 30
320 N under favorable weather conditions although the ACN concentration was low compared
321 to the south of it at the layer of BPTL (between 2000 and 4000 m). The fact that higher
322 OA was observed rather in the higher altitudes than in the lower boundary also

323 demonstrated the vertical distribution over the ECS.

324 Figure 7 displays the vertical distribution of the gases and major aerosol
325 components found on the flights on 17 (blue) and 19 (green) March 2018 as well as the
326 mean concentrations noted in the seven flights (on 17, 19, 22, 24, 26, and 30 March and
327 4 April 2018; red) to the ECS during EMERGe-Asia. Figure 7 illustrates all profiles
328 calculated as 1-min mean and every 500-m interval with one standard deviation ($\pm\sigma$).
329 The number of the data points is displayed on the right side of each figure. The mean
330 CO concentration profile demonstrated a decreasing trend from 240 ppb near the
331 ground to 150 ppb at an altitude of 2500 m and 140–160 ppb at altitudes >6000 m (Fig.
332 7a). The concentration for 17 March 2018 (flight F0317) was similar to the mean
333 concentration profile, except for that at the <1500 m elevation in the lower boundary.
334 However, a higher CO concentration (40–80 ppb) enhancement was noted on 19 March
335 2018 (flight F0319) than the mean profile and flight F0317. The mean difference in CO
336 concentration between flights F0319 and F0317 was as high as 80 ppb at an elevation
337 of 3000-3500 m (Fig. 7a). Similarly, OA concentration was significantly higher in the
338 BPTL vertical distribution in flight F0319 than in the mean profile and flight F0317
339 (Fig. 7b). The mean OA concentration for the flight F0319 peaked at an elevation of
340 2000–2500 m, increasing to $2 \mu\text{g m}^{-3}$ more than in the mean profile and flight F0317.
341 Other aerosol components such as SO_4^{2-} , NH_4^+ , and NO_3^- (Supplementary Fig. S2a-c)
342 also had a similar vertical distribution trend, but the concentration differences were
343 minor compared with OA concentrations. The magnitude of the maximum differences
344 between the flights F0319 and F0317 in the BPTL was 1.3, 0.7, and $0.4 \mu\text{g m}^{-3}$ for
345 SO_4^{2-} , NH_4^+ , and NO_3^- , respectively. The maximum difference concentration of BC can
346 be as high as $1.2 \mu\text{g m}^{-3}$ at 2000-2500 m between the flights F0319 and F0317 (Fig.7c).
347 Regarding the variations in ACN (Fig. 7d) and ACE (Fig. 7e) in the BPTL, their
348 maximum mean concentrations in the flight F0319 were higher than those in the profile

349 of the flight F0317 by 0.18 and 0.9 ppb, respectively. In other words, flight F0319 had
350 a more significant impact on the CO, OA, BC, and volatile organic compound (VOC)
351 species such as ACN and ACE in the BPTL, which might account for the effect of BB
352 emission transport from Indochina. The ozone concentration was lower in both flights
353 F0317 and F0319 than in the mean profile at the elevations <2000 m (Fig. 7f). The
354 ozone titration by NO_x in the low boundary might also play a role. However, it was
355 approximately 5–7 ppb higher in the flight F0319 than in the flight F0317 between the
356 elevations of 1500 and 3000 m. In their downwind area, LRT of BB emissions might
357 increase this concentration further at the BPTL (Tang et al., 2003; Lin et al., 2014) and
358 also discussed in section 4. By contrast, the J value [J(O¹D)] (Fig. 7g) was higher for
359 flight F0317 than for F0319 in the elevation range 1000–3000 m, in line with high aerosol
360 concentrations and associated cloud enhancement that typically lead to decreased
361 photolysis frequencies [i.e., J(O¹D)] (Tang et al., 2003). Figure S3 (Supplementary)
362 indicated the aircraft measurement for the J value (J(O¹D)) and CCN (Cloud
363 Condensation Nuclei; at a constant instrument supersaturation of 0.38 %) along the
364 flight on 19 March 2018. The CCN number concentration (per cm³), was consistently
365 increased with the aerosol species (such as OA) as the flight passed through the BPTL
366 (2000–4000 m). Consistently, at altitudes >4000 m the presence of clouds below the aircraft
367 led to greater J values.

368 The concentrations of other species such as NO_y (Fig.7h) and HONO
369 (Supplementary Fig. S2d) were also greater in flight F0317 than in flight F0319 by 0.4–
370 1.2 ppb and 10–34 ppt, respectively, in the low boundary (<1500 m). At the BPTL, the
371 concentration of NO_y (1–2 ppb) in the flight F0319 was higher than in the flight F0317,
372 but the difference was less than 0.6 ppb. The results from the TRACE-P campaign,
373 which examined the Asian outflow of NO_y, also demonstrated large increases in NO_y
374 concentrations (0.5–1 ppb) downwind from Asia. The NO_y consisted mainly of HNO₃

375 and peroxyacetyl nitrate (Miyazaki et al., 2003; Talbot et al., 2003).

376

377 **4 Simulation results and discussion**

378 **4.1 Model performance and BB transport identification**

379 Tables 2 and 3 and Fig. 8 plot the Pearson correlation coefficients between 5-min
380 merged observations on board the HALO and the simulation for flights F0317 and
381 F0319. Meteorological parameters such as potential temperature (θ), relative
382 humidity (RH), and wind speed (WS) were all captured well by the model along the
383 HALO flight path on the 2 days. The correlation coefficient (R) for meteorological
384 parameters was high, ranging from 0.92 to 0.99 (Table 2). The strong correlation
385 indicates the high representativeness of the reanalysis of meteorological data used in
386 the simulation. Among the trace species and aerosol components, toluene (TOL), NO_x ,
387 BC, OA, ketones (KET), HONO, SO_2 , and HCHO demonstrated an R of >0.5 (good
388 correlation) and CO and O_3 showed an R of nearly 0.5 (Table 2). The simulation
389 performance was investigated in the BL (<1000 m; Fig. 8), at 2000–4000 m altitude
390 (Table 3 and Fig. 8) and for the whole period of both flights (Table 2 and Fig. 8; blue
391 dot). Even in the BPTL, the simulated meteorological parameters presented a good
392 correlation ($R > 0.93$), followed by OA, BC, KET, CO, O_3 , NO_x as well as NH_4^+ and
393 NO_y ($R > 0.5$) (Table 3). In other words, at the BPTL, the R for the simulation
394 significantly increased for OA, BC, CO, O_3 , NO_y and KET (Tables 2 and 3 and Fig. 8),
395 which are indicators for BB being a source of pollution in the model. In contrast, SO_4^{2-} ,
396 NO_3^- , SO_2 , NO_x , TOL, XYL, HCHO and HONO had better correlation in the lower part
397 of the boundary layer, at altitudes <1000 m (see Fig. 8) than in the BPTL. We explain
398 this by the transport of anthropogenic pollutants in the continental outflow in the lower
399 part of the boundary layer in ECS.

400 The modeling results tended to overestimate the concentration of the species, with

401 examples being CO (64 ppb), OA ($0.3 \mu\text{g m}^{-3}$), BC ($0.2 \mu\text{g m}^{-3}$) and O₃ (12.5 ppb;
402 Table 3) in the BPTL. Because high concentrations of CO, BC and OA in BPTL are
403 accurate indicators of BB in the model, the BB emission from the source of FINN data
404 are probably also overestimated (Lin et al., 2014). Except for OA and BC, the
405 correlations for other aerosol components such as NO₃⁻, and SO₄²⁻ were poor (0.13 and
406 0.2, respectively). The poor correlation for SO₄²⁻ may result from the large uncertainty
407 in the emission of SO₂.

408 Because the meteorological parameters were simulated well, the simulation
409 discrepancies for chemical species are either caused by the emission estimation
410 uncertainties or by inaccuracies in the simulation of chemical oxidation processes
411 during LRT. Because CO, OA, and BC are accurate indicators of simulated BB
412 transport from Indochina (Carmical et al., 2003), the airborne measurements on board
413 the HALO are used as reference to evaluate the performance of the model for the flight
414 F0319 (Fig. 9). The 5-min merged simulation of CO concentration with (blue line) and
415 without (green line) BB was compared to that measured on board the HALO (red line);
416 the concentration was mostly in the range of 100–200 ppb, with its peak approaching
417 300 ppb (at 01:50, 02:50, and 04:00 UTC) at the BPTL (Fig. 9a). In general, the
418 simulation captured the CO variation along the flight path. However, it overestimated
419 the observations by nearly 100 ppb for the simulation with BB at the BPTL during
420 01:00–02:00, 03:40–04:20, 05:00–05:40, and 06:30–07:20 UTC (Fig. 9a). Notably, the
421 simulation difference was minor when the flight was in the lower part of the boundary
422 layer (02:30 and 06:00 UTC) i.e. < 1000m or at elevations of >4000 m (03:00–03:30
423 and 04:20–05:00 UTC). The model underestimated CO concentration in the lower part
424 of the boundary (<1000 m) (02:30 and 05:50–06:30 UTC) over the ECS. In conclusion,
425 our model simulation overestimates BB emissions but underestimates continental CO
426 emissions from China due to the underestimation of the emission inventory of the

427 MICS-Asia III (Kong et al.,2020) was adopted in this study.

428 OA and BC are also important BB indicators and were reasonably captured by the

429 model before 03:00 UTC when the flight was south of 28 °N at elevations of <4000 m

430 (Fig. 9 b-c). The time series of simulated OA and BC has peak concentrations of nearly

431 4-5.5 $\mu\text{g m}^{-3}$ and 2 $\mu\text{g m}^{-3}$, respectively, during HALO shuttle flights passing through

432 the BPTL (2000–4000 m) around 01:50 and 02:50 UTC. When BB emission was not

433 included in the simulation, the concentration peaks were not observed (see Fig. 9b-c,

434 green plot). Similar to the simulated CO results, the simulated OA and BC overestimate

435 the amounts of these species to the north of 28 °N during 03:30-04:20 UTC (Fig. 6a

436 and 9). Furthermore, when the simulation only considered direct effect (case ROCD,

437 purple), the overestimations were increased as shown in Figure 9b-c. As mentioned

438 earlier, a frontal system was just located from the ECS to SC (Fig. 2e) on 19 March

439 2018. In other words, the effect of wet scavenging reduced the aerosol concentration

440 bias in the ECS and SC, as for the frontal system providing the moist air mass in the

441 event flight F0319. The model after 07:30 UTC, which was related to local emissions

442 before HALO landed over western Taiwan on 19 March 2018. In general, our model

443 simulation captured reasonably well OA and BC with an R of 0.61 and 0.74,

444 respectively. A minor mean bias for OA (BC) is 0.3 $\mu\text{g m}^{-3}$ (0.1 $\mu\text{g m}^{-3}$) and the root

445 mean square error (RMSE) of OA (BC) is 1.1 $\mu\text{g m}^{-3}$ (0.4 $\mu\text{g m}^{-3}$) (Table 2). The R for

446 OA (BC) reached 0.85(0.79), with an RMSE of $0.7 \mu\text{g m}^{-3}$ ($0.5 \mu\text{g m}^{-3}$) when we
447 calculated the BB transport layer only between 2000 and 4000 m (Table 3 and Fig. 8).
448 In addition to OA and BC, simulated aerosol species such as SO_4^{2-} was overestimated,
449 whereas NO_3^- was underestimated although their concentrations were low (Table 3).
450 Because the BPTL was mainly between altitudes of 2000 and 4000 m, the subsequent
451 discussion focuses on the influence of the BPTL from Indochina on the downstream
452 areas, particularly the ECS and Taiwan.

453 **4.2 Effects of LRT BB plume from Indochina on East Asia**

454 To investigate the regional impacts of BB plume transport from Indochina, we
455 compared the simulation with and without BB emission for the events on 17 and 19
456 March 2018. The analysis of the calculations focused on the impact over SC, Taiwan
457 and ECS. These three selected regions are SCA (in South China), TWA (covered the
458 whole Taiwan), and ECSA (in the ECS) as shown in Figure 1a. After being emitted the
459 BB pollutants from Indochina were then transported northward to China and
460 subsequently northeastward. The exact flow pattern depended on the weather
461 conditions and flow types (ridge or trough) at 700 hPa (3000 m) between 17 and 19
462 March 2018 (see Fig. 2). Consequently, we investigated the hourly variation in the area
463 mean concentrations or mixing ratios of air pollutant trace constituents to assess the
464 importance of BB emissions from Indochina on the selected downstream region e.g. the
465 ECSA (Fig. 10), SCA, TWA and ECSA (Table 4). The contribution of CO (or others
466 species) due to BB was estimated by the difference between simulations with and
467 without the BB emission. These differences are then expressed as a fraction in
468 percentage shown in Figure 10 (blue line). The mean concentration of CO (red line)

469 over the ECSA (Fig. 10a) was at its lowest (115 ppb) on 17 March 2018; it gradually
470 increased to a peak concentration of 280 ppb on 18 March 2018 and then remained
471 stable at 260 ppb on 19 March 2018. The contribution of CO from BB (blue line)
472 ranged from 19 % (<22 ppb) on 17 March 2018 to a peak of 42 % (~113 ppb) on 18
473 March 2018 and then gradually declined to 26 % on 19 March 2018 (Fig. 10a). As
474 for OA (BC), the lowest percent contribution by BB was 14-16% (<5%) between 16
475 and 17 March 2018 while the highest could be more than 40% (80%) during 18 and 19
476 March 2018 (Fig. 10b and c). The BB contributed to PM_{2.5} was 19 % (0.39 $\mu\text{g m}^{-3}$) on
477 17 March 2018 (Fig. 10d), increasing to 45 % (3.6 $\mu\text{g m}^{-3}$) on 18-19 March 2018
478 because the BB plume spread by the strong wind to the ECSA.

479 The variation of O₃ (Fig. 10e) depends on transport and photochemistry, which
480 involves the precursors NO_x and VOC and the photolysis frequency of NO₂, J(NO₂).
481 For the elevations between 2000–4000 m, O₃ changes are similar to those of CO, NO_x
482 and KET, which were mainly contributed by the LRT BB plume and related to the
483 ozone precursor after 18 March 2018. The lowest and highest O₃ concentrations on 17
484 and 18 March 2018 were 56 and 75 ppb, respectively, of which we estimate that 5.6
485 ppb (10 %) and 34 ppb (45 %) were BB's contributions, respectively. Although the
486 mean NO_x concentration was relatively small (0.06–0.18 ppb), the BB contributed 35–
487 70 % (0.02–0.13 ppb) during 17–19 March 2018 (Supplementary Fig. S4a). The KET
488 concentration was in the range 0.4 to 2.7 ppb, with BB contributing nearly 20–26 %
489 (0.08–0.7 ppb) during 17–19 March 2018 (Supplementary Fig. S4b).

490 The area-mean OH contributed by BB increased from its lowest level (<30 %) on
491 17 March 2018 to its highest (nearly 70 %) on 19 March 2018 (Fig. 10f). HO₂ was also
492 observed to increase trend from 10 % to 40 % during daytime over the period 17–19
493 March 2018 (Supplementary Fig. S4c). The amounts of the oxidizing agent, OH, and
494 the free radical HO₂ depend on the amounts of trace gases, which produce and remove

495 these radicals, (eg. NO_x , water vapor, ozone, hydrocarbons, etc.) and the relevant
496 photolysis frequencies $J(\text{O}_3 \rightarrow \text{O}^1\text{D})$, $J(\text{NO}_2)$ etc.. Thus trace constituents from BB were
497 expected to increase OH and HO_2 . However, BB's contribution to photolysis
498 frequencies $J(\text{O}_3 \rightarrow \text{O}^1\text{D})$ (Fig. 10g), $J(\text{NO}_2)$ (Supplementary Fig. S4d) etc. decreased
499 as the mean BB aerosol concentration increased over the ECS during 17–19 March
500 2018. This is because photolysis calculation results used simulated aerosol and cloud
501 formation, which increased over the ECSA (Fig. 12).

502 The NO_y , mean concentration ranged from 1.0 to 4.5 ppb, of which BB's
503 contribution was from 55 to 82 % (Supplementary Fig. S4e). Such a high contribution
504 from BB also demonstrated the effects of long-distance transport. Figure 10h indicates
505 an increasing trend of HCHO concentration from 17 to 19 March 2018. HCHO
506 formation and destruction depend on the rate of reaction of OH with HCHO precursors
507 and the rate of reaction of HCHO with OH and the photolysis frequency of HCHO. As
508 a result, HCHO production varied with OH concentration. The lowest and highest
509 concentrations of HCHO were on 17 and 19 March 2018, respectively. In summary,
510 the consistent variations in BB contributions to CO, OA, BC, $\text{PM}_{2.5}$, OH, HCHO, NO_x ,
511 NO_y , and O_3 peaked on 18 or 19 March 2018, whereas $J(\text{O}^1\text{D})$ decreased between 17
512 and 19 March 2018.

513 Figure 11 displays the fraction in % that the long-range transported BB emission
514 contributes to the amounts of NO_x , NO_y , $\text{PM}_{2.5}$, OA, BC, OH, O_3 , CO, KET, HO_2 ,
515 HCHO and $J(\text{O}^1\text{D})$, over the ECSA on 17 and 19 March 2018. Except for NO_y , BB
516 contribution was generally <11 % at elevations of <1000 m over the ECSA. The scatter
517 distribution of the simulation results indicates that the effect of BB emission at
518 elevations of <1000 m (Fig. 11a) was significantly lower than that between the
519 elevations of 2000 and 4000 m (Fig. 11b). For NO_y , NO_x , $\text{PM}_{2.5}$, BC, OH, O_3 , and CO,
520 the BB contribution was >30 % at the elevation of 2000–4000 m over the ECSA (Fig.

521 11b). Table 4 further summarizes the effect of BB emission on the downwind areas
522 (SCA, TWA, and the ECSA) at the <1000 m and 2000–4000 m elevations. The
523 contribution of BB to NO_y, NO_x, PM_{2.5}, BC, OH, O₃ and CO was at least 30–80 % at
524 the elevation of 2000–4000 m over the regions SCA, TWA and ECSA (Table 4). In the
525 lower boundary layer (i.e. <1000 m), the BB contribution for most species at the remote
526 downstream areas was <20 %, except for TWA. Because of the high mountains (Lin et
527 al. 2021) present in TWA, the BB plume passing over Taiwan was potentially
528 transported downward through mountain–valley circulation to the lower boundary layer
529 (Ooi et al., 2021). The influence of BB over TWA was the highest among these three
530 downstream regions (see Table 4) as its location was directly on the transport pathway
531 for the BB plume on the major event day (flight F0319).

532 Figure 12a displays the simulated cloud water difference with and without BB
533 emission over different regions on 17 and 19 March 2018. BB aerosols are a potential
534 source of cloud nuclei. The simulations show the impact of BB on cloud water
535 enhancement (Fig. 12a) in the vertical distribution. Cloud water enhancement over SCA
536 was associated with aerosol enhancement from the BB in the altitude range 1000–4000
537 m: the peak being 1.8–2.0 mg kg⁻¹ at 2000 m on these 2 days (Fig. 12a). The abundance
538 of BB emissions transported from Indochina to SCA (Fig. 3) is expected to contribute
539 to the high cloud water formation over SCA. Furthermore, the southerly flow (Fig. 3)
540 that transports warm and moist air mass from the South China Sea may have favored
541 cloud formation in flights F0317 and F0319. High cloud water related to BB can be
542 seen in the simulations of these two days. In the remote ECSA regions, the cloud water
543 substantially increased on 19 March 2018 (Fig. 12a) compared to 17 March 2018
544 because of a significant difference in BB emissions transported to the ECSA between
545 17 and 19 March 2018 (Fig. 3). Similarly, the cloud water enhancement over Taiwan
546 also only appeared on 19 March 2018 (Fig. 12a). Furthermore, nearly no difference in

547 the cloud water vertical distribution over the region IDCA (Fig. 1a) in Indochina was
548 noted because in the Indochina region, spring is the dry season (Lin et al., 2009) and
549 thus unfavorable for cloud water formation. Figure 12b shows the cloud water
550 difference when the aerosol indirect effect turned off in the simulation over different
551 regions on 19 March 2018. The significant cloud water shortage over ECSA, and SCA
552 could be as high as 2.4 mg/kg and 1.5 mg/kg, respectively (Fig.12b). In other words,
553 the role of the chemistry-microphysics interactions (indirect effect) plays an important
554 role in the cloud water enhancement in the SCA and ECSA in this study.

555 The simulated downward shortwave flux at the noontime at ground surface due
556 to BB was 2-4% and 5-7% reduction over the regions ECSA and SCA, respectively,
557 (supplementary Fig. S5a-b, blue line) during 18-19 March 2018. However, a significant
558 shortwave flux reduction at the noontime at ground surface could be 15-20% due to
559 aerosol indirect effect in the region SCA during 18-19 March 2018 (supplementary Fig.
560 S5a-b blue dashed line). The combination of BB aerosols enhancement and increased
561 cloud water results in shortwave radiation reduction, implying the possibility of
562 regional climate change in East Asia driven by BB aerosols.

563

564 **5. Summary**

565 The BB during spring in Indochina has a significant impact on the chemistry and
566 composition of the troposphere in the surrounding regions of East Asia. During the
567 EMERGE campaign in Asia, atmospheric pollutants were measured on board the HALO
568 aircraft. In this study, a minor long-range BB transport event was observed from
569 Indochina on 17 March 2018 (flight F0317), followed by a major long-range BB
570 transport event on 19 March 2018 (flight F0319). The impact on tropospheric trace
571 constituent composition and the environment has been investigated.

572 During the major BB transport event F0319, the 1-min mean of the peak

573 concentrations of the trace constituents CO, O₃, ACE, ACN, OA and BC between the
574 altitudes of 2000 and 4000 m over the ECS were 312.0 ppb, 79.0 ppb, 3.0 ppb, 0.6 ppb,
575 6.4 $\mu\text{g m}^{-3}$, 2.5 $\mu\text{g m}^{-3}$ respectively. In comparison during the F0317 event CO, O₃,
576 ACE, ACN, OA and BC were 203.0 ppb, 71.0 ppb, 2.0 ppb, 0.3 ppb, 3.4 $\mu\text{g m}^{-3}$, 1.2
577 $\mu\text{g m}^{-3}$ respectively.

578 When the elevation was <1000 m for both the F0317 and F0319 events, the sulfates,
579 rather than OA, had the highest concentrations. The peak concentration could be as high
580 as 5.1 $\mu\text{g m}^{-3}$ in the low boundary for the event F0317 in the ECS. This observation is
581 most likely explained by a continental outflow from regions having fossil fuel
582 combustion in the lower boundary layer over the ECS.

583 In this study, the WRF-Chem model was employed to evaluate the BB plume
584 transported from Indochina and its influence on the downstream areas including South
585 China, Taiwan, and the ECS. The contribution of the BB plume for most species in the
586 remote downstream areas was <20 % in the lower boundary layer (altitude <1000 m).
587 In comparison, the contribution of long-range transported BB plume was 30–80 %, or
588 even higher, for many of the trace constituents (NO_y, NO_x, CO, OH, O₃, BC and PM_{2.5})
589 in the altitude range between 2000 and 4000 m for SC, Taiwan, and the ECS. The large
590 influence of BB over Taiwan is most probably because the BB transport passes directly
591 over Taiwan.

592 BB aerosols are potential sources of cloud nuclei. The WRF simulations estimate
593 the effect of the BB plume on cloud water formation over SC and the ECS. We observe
594 in the simulations cloud water enhancement over SC at elevations of 1000–4000 m.
595 This increase of cloud water is consistent with an increase in aerosol, caused by BB
596 emissions, transported from Indochina to SC. In remote regions of the ECS, the
597 simulated cloud water was significantly larger during the major BB event on 19 March
598 2018 than the minor BB event on 17 March 2018. The simulated decrease of the

599 photolysis frequency ($J(\text{O}^1\text{D})$ and $J(\text{NO}_2)$) is attributed to the difference in aerosol
600 concentrations and associated cloud enhancement between the two events over the ECS.
601 This we explain by the significant differences in BB emissions transported to the ECS
602 between the two events. The combination of increased BB aerosol concentration and
603 increased amounts of cloud water led to reductions in the amount of incoming
604 shortwave radiation at the surface over the ECS and SC. This influences tropospheric
605 chemistry and composition, regional climate, precipitation, ocean biogeochemistry,
606 agriculture, and human health.

607

608 ***Data availability***

609 The EMERGe data are available at the HALO database
610 (<https://doi.org/10.17616/R39Q0T>, DLR, 2022) and can be accessed upon registration.
611 Modeling data can be made available upon request to the corresponding author.

612 ***Author contribution***

613 CYL conceived the idea, analyzed the data, writing and editing of the manuscript. WNC
614 and YYC run the model and analyzed the data. CKC joined the manuscript
615 discussion. CYLiu provided the MODIS data. HZ and HS provided trace gases data. EF
616 provided acetonitrile data. FO performed the ozone measurement. OOK, BAH and
617 MLP were responsible for the BC measurement. KK and JS were responsible for C-
618 ToF-MS measurements. KP and BW provided HONO data. JPB and MDAH led the
619 EMERGe-Asia experiment. All authors have read and agree to the published version of
620 the manuscript.

621 ***Competing interests***

622 The authors declare that they have no conflict of interest.

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820 Table 1: WRF-Chem model configuration and physics and chemistry options in this
 821 study. (RRTMG=Rapid Radiative Transfer Model for General Circulation Models;
 822 FINN=Fire Inventory from National Center for Atmospheric Research)
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Resolution	10km
Microphysics	Lin
Cumulus parameterization	Grell 3D ensemble scheme
Planetary Boundary Layer	Mellor-Yamada-Janjic TKE scheme
Longwave radiation	RRTMG
Shortwave radiation	RRTMG
Fire emissions	FINN V1.5
Anthropogenic emissions	MICS-Asia III(2010) + Taiwan Emission Data System ver 9.0 (2013)
Biogenic emissions	MEGAN V2.04
Chemistry option	RACM Chemistry with MADE/VBS aerosols using KPP library along with the volatility basis set (VBS) used for Secondary Organic Aerosols
Photolysis option	Madronich
wet scavenging	On , (Neu and Prather, 2012)
Cloud chemistry	On,
feedback from the aerosols to the radiation schemes	On
the time interval for calling the biomass-burning plume rise subroutine	180 min
feedback from the parameterized convection to the atmospheric radiation and the photolysis schemes	On
Subgrid-scale wet scavenging	on
Subgrid aqueous chemistry	on

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833 Table 2 Observed and simulated mean values for bias (BIAS), root mean square error
 834 (RMSE), and correlation coefficients (R) for EMerGe HALO flights on 17 and 19
 835 March 2018. KET*: the observed Acetone is applied to compare with simulated ketones
 836 (KET).

	OBS_ave	SIM_ave	BIAS	RMSE	R
THETA(K)	304.8	304.2	-0.6	1.1	0.99
WS(m/s)	9.1	8.5	-0.6	2.0	0.94
RH(%)	63.6	62.9	-0.6	10.7	0.92
OA($\mu\text{g}/\text{m}^3$)	1.2	1.4	0.3	1.1	0.61
BC($\mu\text{g}/\text{m}^3$)	0.4	0.5	0.1	0.4	0.74
SO ₄ ²⁻ ($\mu\text{g}/\text{m}^3$)	1.1	2.5	1.4	2.3	0.42
NO ₃ ⁻ ($\mu\text{g}/\text{m}^3$)	0.2	0.6	0.5	2.1	0.31
NH ₄ ⁺ ($\mu\text{g}/\text{m}^3$)	0.4	0.7	0.3	1.2	0.49
CO(ppb)	170.8	191.8	20.9	72.8	0.45
SO ₂ (ppb)	0.2	0.7	0.4	1.2	0.55
O ₃ (ppb)	59.7	63.2	3.5	14.4	0.43
NO _x (ppb)	0.2	0.2	0.0	0.2	0.72
NO _y (ppb)	1.2	2.6	1.3	2.3	0.03
KET*(ppb)	1.4	1.6	0.1	0.9	0.59
TOL(ppb)	0.1	0.1	0.0	0.1	0.75
XYL(ppb)	0.1	0.0	0.0	0.1	0.40
HCHO(ppb)	0.1	0.7	0.5	0.7	0.51
HONO(ppt)	10.5	1.0	-9.4	15.3	0.56

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838 Table 3 Observed and simulated mean values at an elevation between 2 km and 4 km
 839 for bias (BIAS), root mean square error (RMSE), and correlation coefficients (R) during
 840 EMeRGe HALO flights on 17 and 19 March 2018. KET*: the observed Acetone is
 841 applied to compare with simulated ketones (KET).

	OBS_ave	SIM_ave	BIAS	RMSE	R
THETA(K)	307.5	306.7	-0.7	0.9	0.98
WS(m/s)	8.2	7.9	-0.3	1.7	0.93
RH(%)	55.8	56.0	0.2	7.6	0.96
OA($\mu\text{g}/\text{m}^3$)	1.3	1.6	0.3	0.7	0.85
BC($\mu\text{g}/\text{m}^3$)	0.4	0.7	0.2	0.5	0.79
SO ₄ ²⁻ ($\mu\text{g}/\text{m}^3$)	0.8	2.5	1.7	2.1	0.20
NO ₃ ⁻ ($\mu\text{g}/\text{m}^3$)	0.1	0.0	-0.1	0.3	0.13
NH ₄ ⁺ ($\mu\text{g}/\text{m}^3$)	0.4	0.4	0.0	0.2	0.52
CO(ppb)	164.4	228.7	64.2	85.4	0.58
SO ₂ (ppb)	0.0	0.7	0.6	0.9	0.07
O ₃ (ppb)	60.1	72.6	12.5	15.0	0.55
NO _x (ppb)	0.1	0.2	0.0	0.1	0.53
NO _y (ppb)	1.0	3.6	2.6	3.0	0.51
KET*(ppb)	1.5	2.0	0.5	1.0	0.70
TOL(ppb)	0.1	0.0	0.0	0.1	0.16
XYL(ppb)	0.0	0.0	0.0	0.0	-0.17
HCHO(ppb)	0.1	0.7	0.6	0.7	0.25
HONO(ppb)	6.0	0.6	-5.4	7.2	0.23

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862 Table 4: Simulated biomass burning contribution (with and without BB emission in
863 Indochina) in percentage (%) on 17 and 19 March, 2018 for different regions: SCA,
864 TWA, ECSA as shown in Figure 1a

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Average	SCA		TWA		ECSA	
	< 1KM	2-4KM	< 1KM	2-4KM	< 1KM	2-4KM
NO _y	13.6	72.2	39.7	83.3	14.8	69.9
NO _x	-1.3	58.1	2.9	71.1	1.4	51.0
PM _{2.5}	7.5	46.0	15.1	55.6	7.6	34.4
OA	5.3	41.4	7.5	48.1	4.4	28.5
BC	8.0	79.5	16.4	81.4	6.8	47.9
OH	14.7	43.8	24.1	67.4	9.2	48.3
O ₃	18.8	34.2	23.2	39.2	9.2	31.3
CO	9.8	31.7	21.9	38.4	11.1	32.2
KET	6.2	17.8	9.5	27.5	7.2	24.7
HCHO	-4.2	9.8	-4.8	20.6	-4.7	10.4
HO ₂	8.8	2.6	15.2	35.8	6.3	23.2
J(O ¹ D)	-1.5	-0.8	-1.1	0.5	-1.5	-1.0

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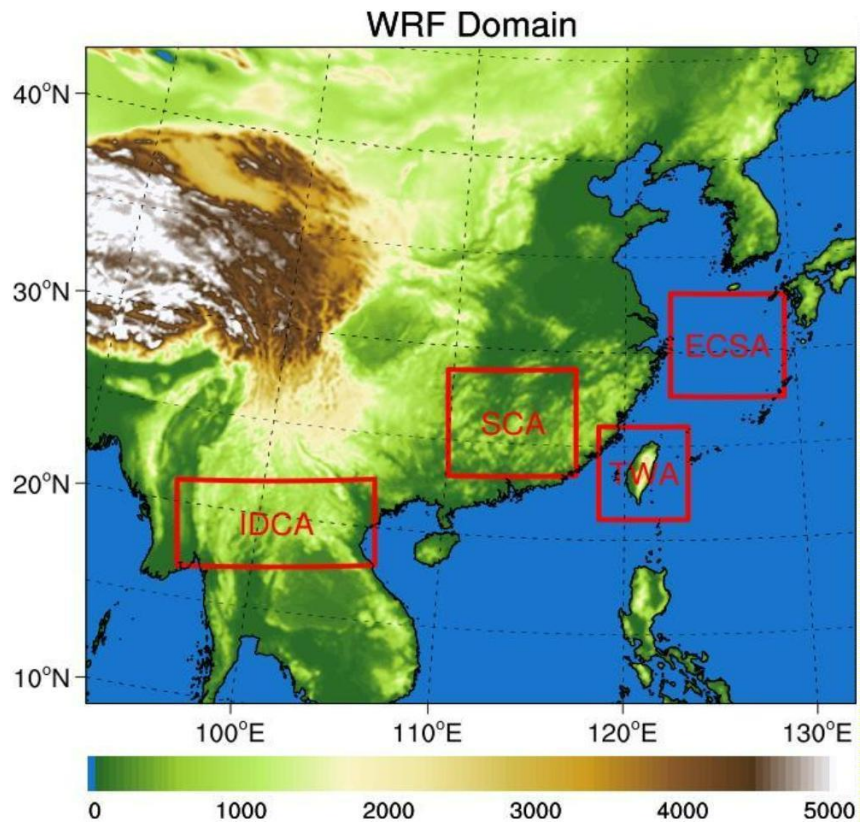
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874 (a)

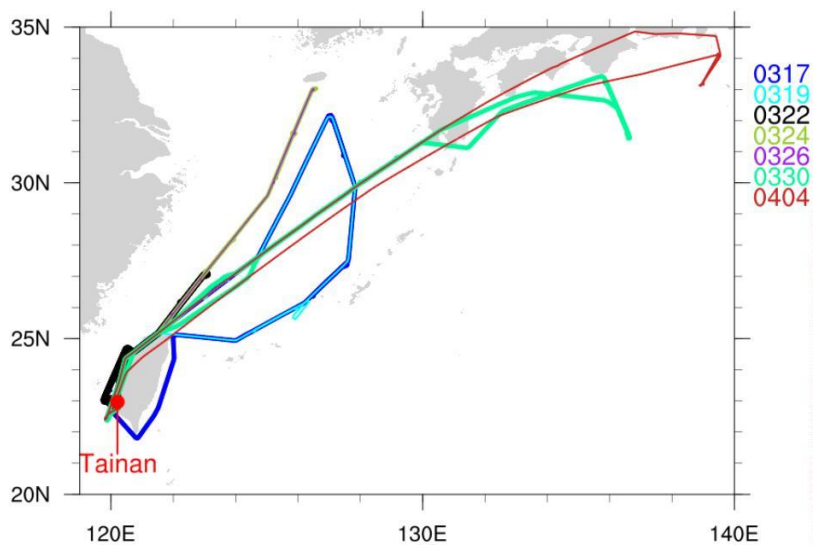


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892 (b)



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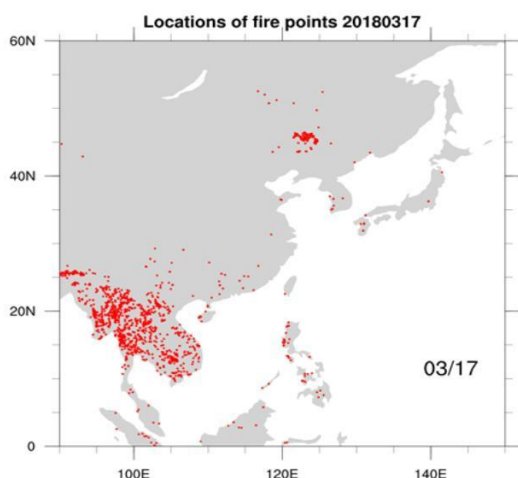
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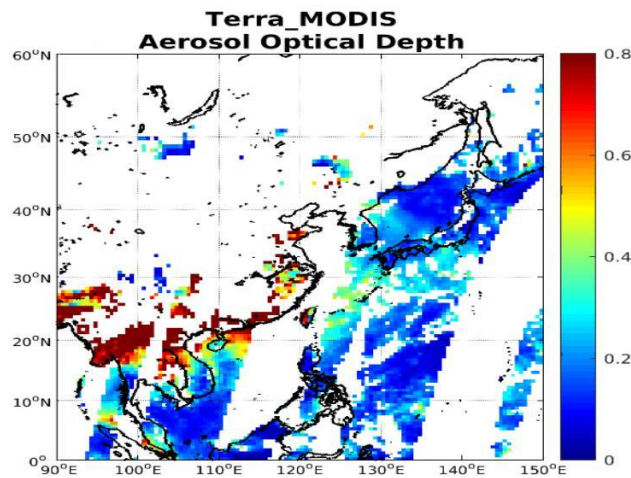
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Figure 1 (a) Configuration of Weather Research and Forecasting model domain, topography, and location of proposed study areas in East Asia, namely IDCA(Indochina area), SCA (southern China area), TWA (Taiwan area) and ECSA (East China Sea area), respectively. (b) The HALO flights on 17, 19, 22, 24, 26, 30 March, and 04 April during EMERGE Asia campaign. Different colors indicated different flights over East Asia. Maps and plots were produced using NCAR Command Language (NCL) version 6.6.2.

912 (a)

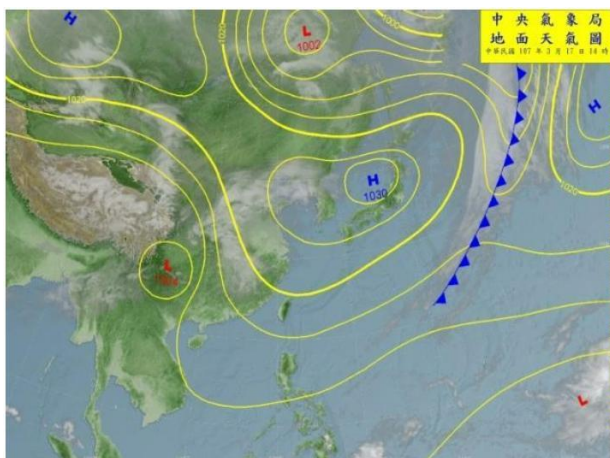


(b)

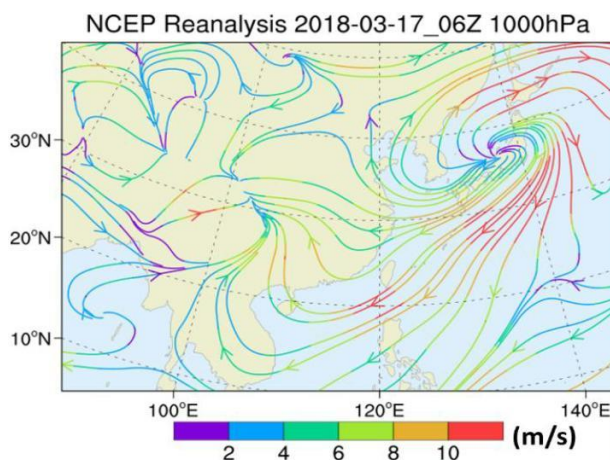


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914 (c)



(d)

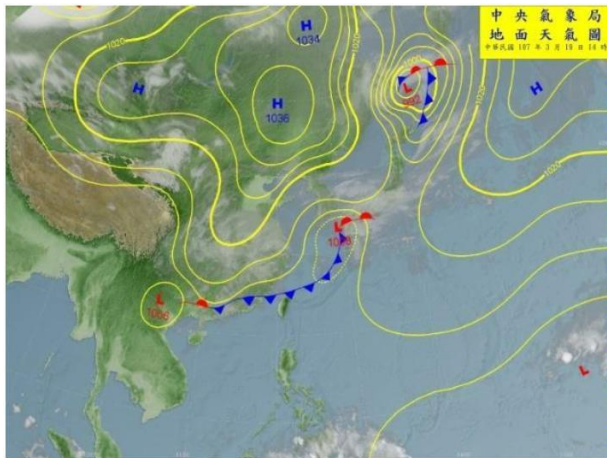


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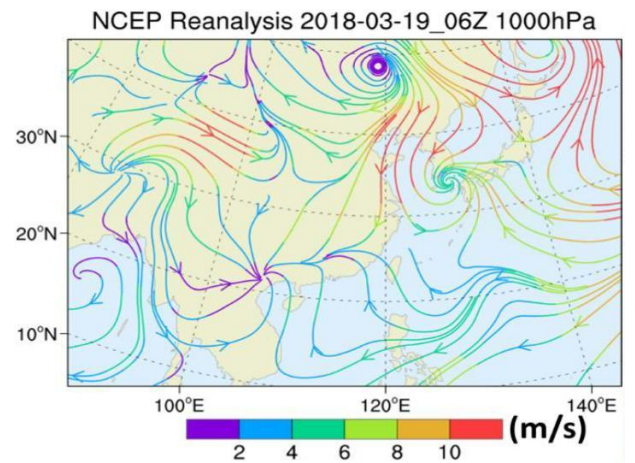
916 Fig.2 (a) MODIS fire hot spots on 17 March 2018 (source: [https://modis-](https://modis-fire.umd.edu/guides.html)
 917 [fire.umd.edu/guides.html](https://modis-fire.umd.edu/guides.html)) and (b) Composited Aerosol Optical Depth (AOD) from
 918 MODIS onboard NASA Terra satellite. The Collection 6.1 AOD is downloaded from
 919 NASA Earth Data website (<https://www.earthdata.nasa.gov/learn/find-data>), and
 920 composited for 0110, 0115, 0120, 0125, 0130, 0250, 0255, 0300, 0305, 0310, 0430,
 921 0435, 0440, 0445, 0610, 0615, 0620, 0745 and 0750UTC data granules on 17 March
 922 2018. (c) weather Chart at 06:00 UTC on 17 March 2018 (d) 1000 hPa streamlines at
 923 06:00 UTC, 17 March 2018 (e) and (f) same as (c) and (d) but on 19 March 2018 ;(g)
 924 700 hPa streamlines at 06:00 UTC, on 17 March 2018 (h) 700 hPa geopotential height
 925 at 06:00 UTC, on 17 March 2018; (i) and (j) same as (g) and (h) but on 19 March
 926 2018.

927 Near-surface weather charts and satellite images were provided by Central Weather
 928 Bureau (CWB) Taiwan. The near-surface and 700 hPa streamlines and geopotential
 929 height were deduced from NCEP Reanalysis data. Maps and plots were produced using
 930 NCAR Command Language (NCL) version 6.6.2.

931 (e)



(f)

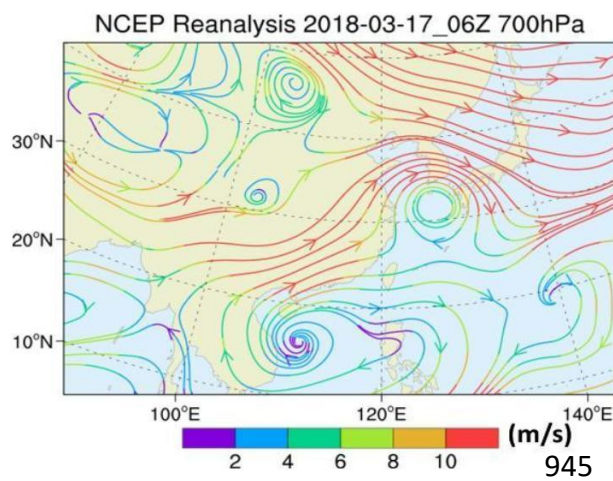


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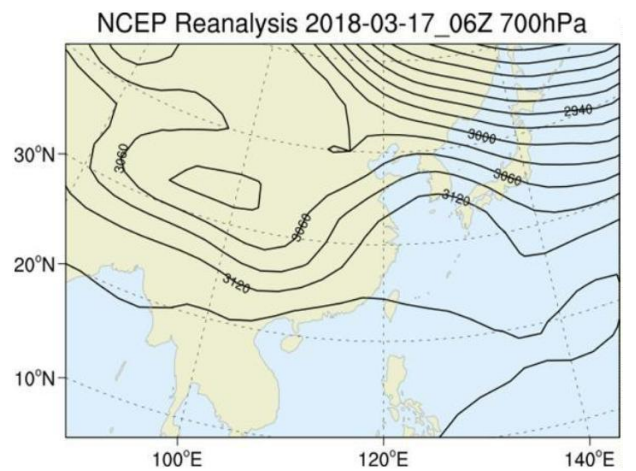
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934 (g)

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(h)



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949 Figure 2 e-h continued

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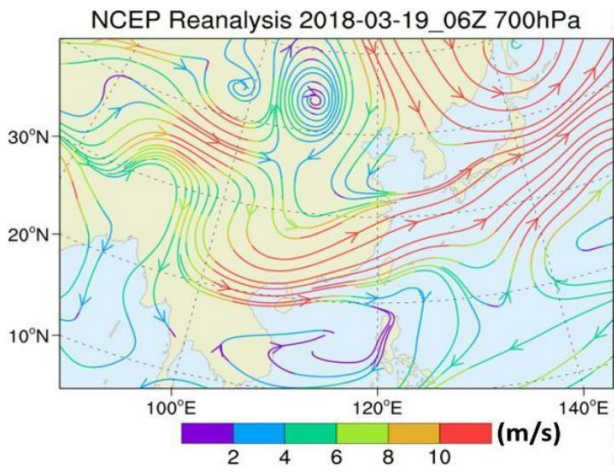
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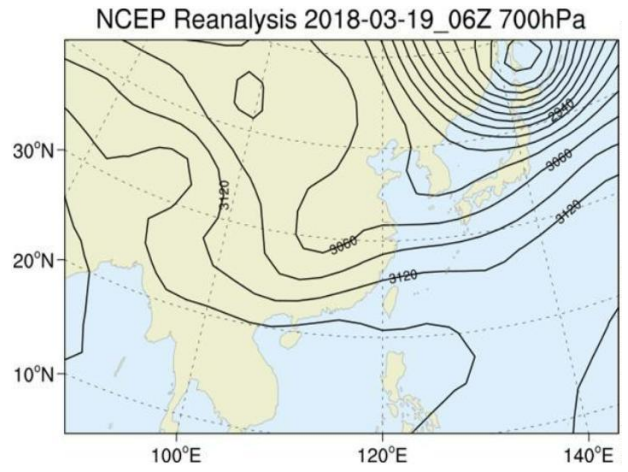
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959 (i)
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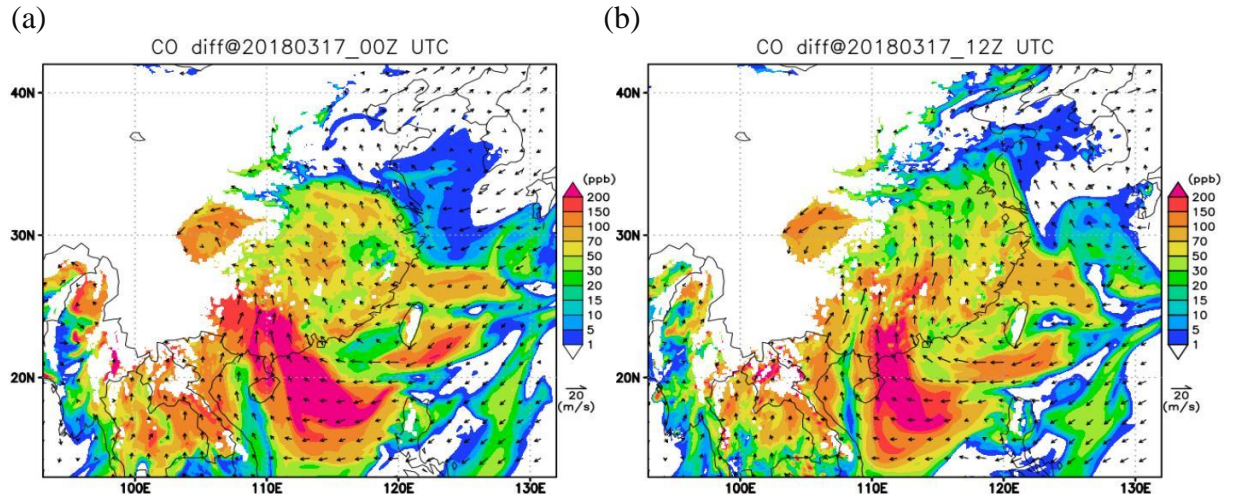
(j)



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Fig. 2 i-j continued

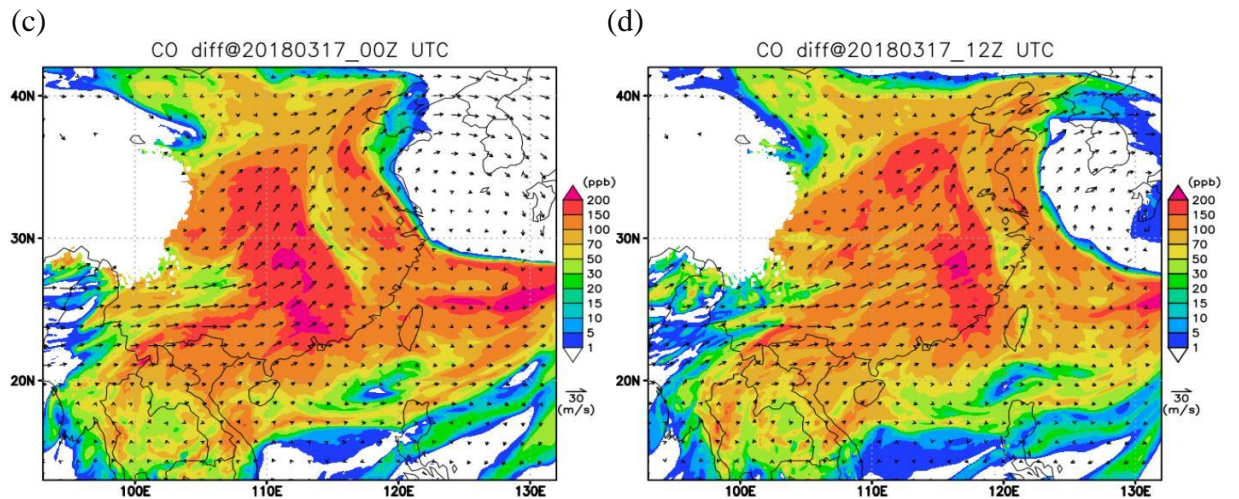
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995 Fig 3 a-d : Simulated wind field ($m s^{-1}$) distribution and concentration (unit: ppb)

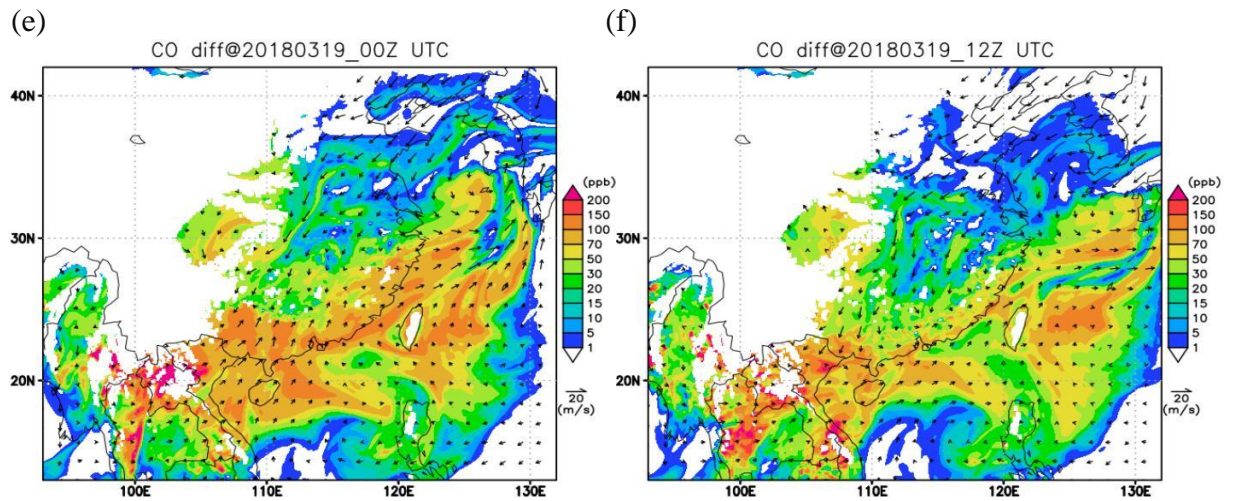
996 difference with and without BB emission for CO on 17 March, 2018 at 00:00 UTC (a,

997 c) and 12:00 UTC (b, d) for 1km altitude (a, b) and 3km altitude (c, d). (unit:ppb)

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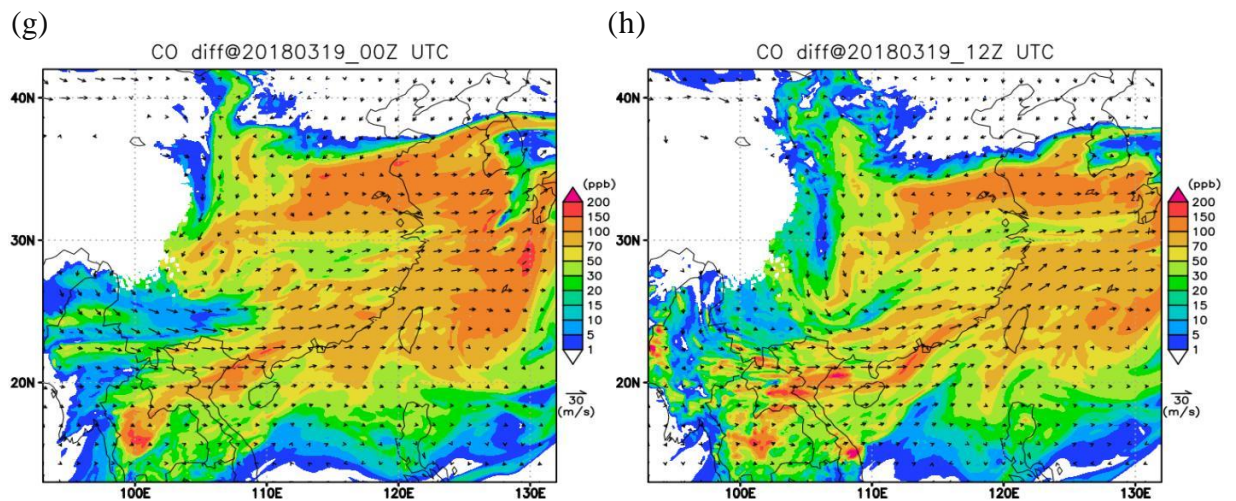
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Fig 3 e-h: Simulated wind field (m s^{-1}) and concentration (unit: ppb) difference with and without BB emission for CO on 19 March, 2018 at 00:00 UTC (e, g) and 12:00 UTC (f, h) for 1km altitude (e, f) and 3km altitude (g, h).

1019 (a)

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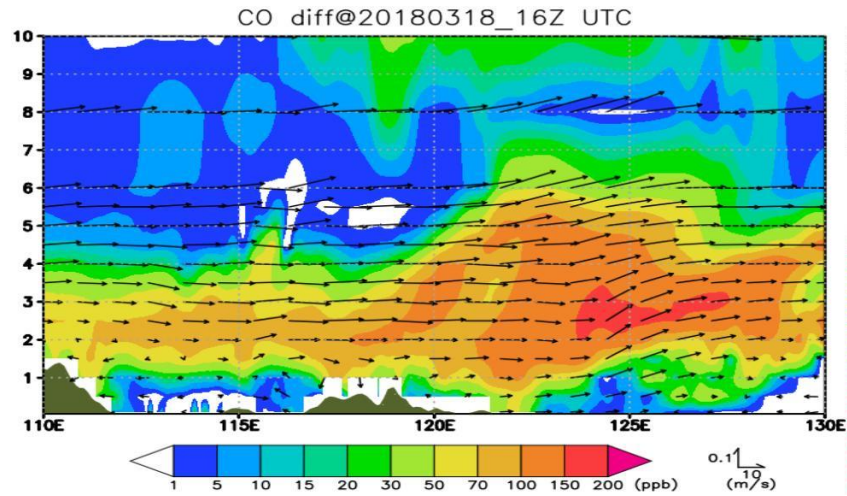
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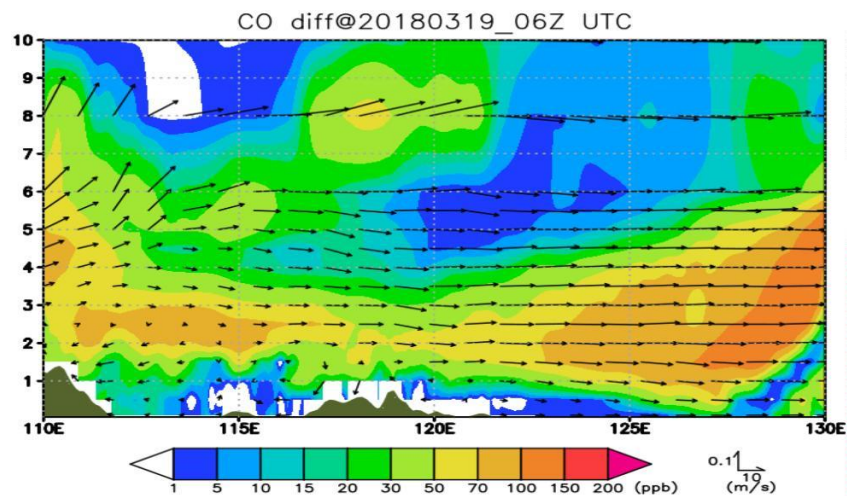
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1027 (b)



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1032 Fig. 4 Simulated wind field (m s^{-1}) distribution and the concentration (ppb) difference

1033 between with and without BB emission for CO at cross-section 30°N (a) 16:00 UTC

1034 18 March 2018 (b) 06:00 UTC, 19 March 2018. Wind vectors represent along section

1035 winds, with scales shown at the down-right corner of plot (unit: m s^{-1})

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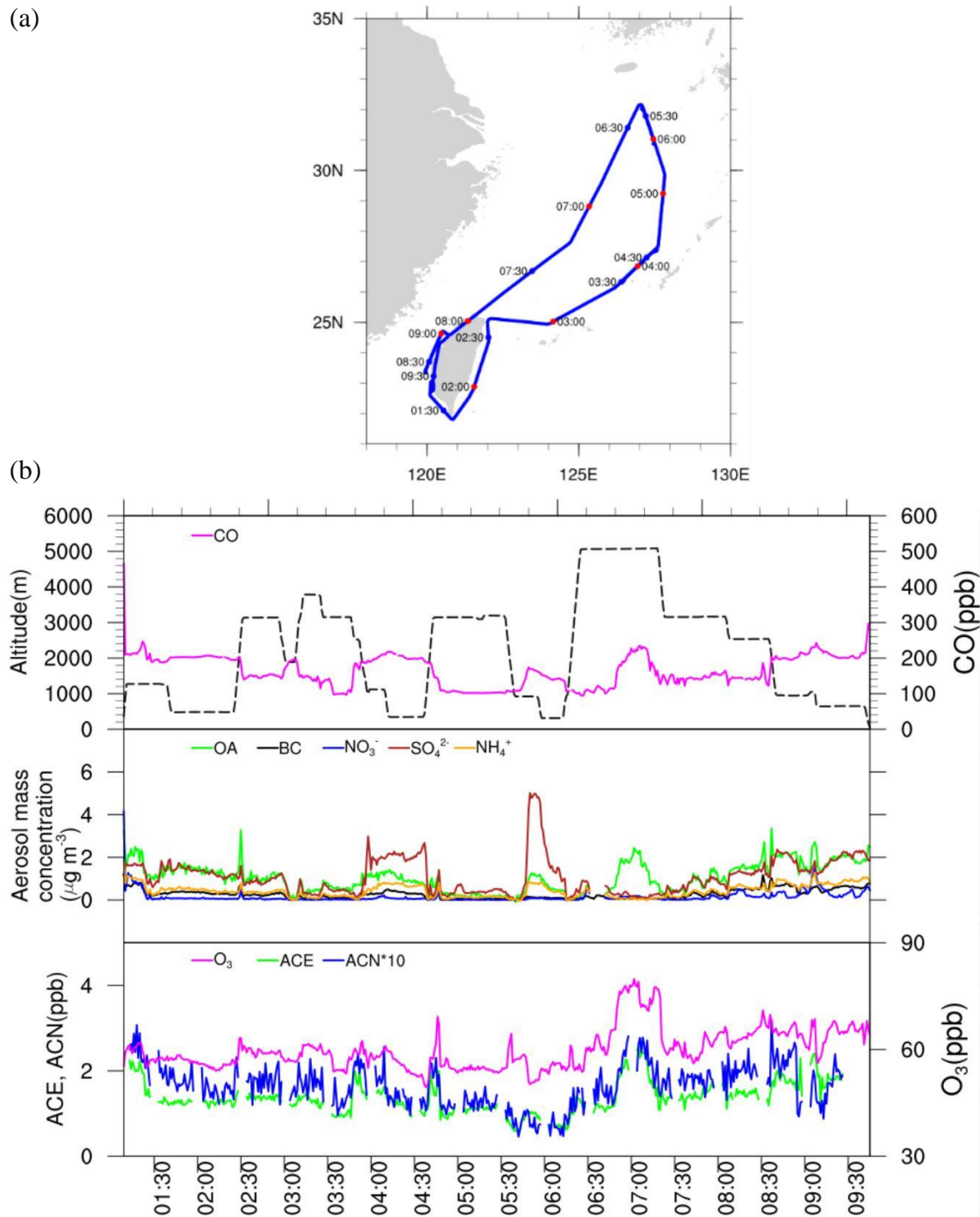
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1041 (a)
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 1052 (b)



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Fig. 5 (a) The HALO flight and detailed locations on 17 March 2018. (b) Flight altitude and 1-min mean of observed concentrations for CO (upper), Organic aerosol (OA), BC aerosol (BC), SO_4^{2-} , NO_3^- , NH_4^+ (middle), O_3 , acetone (ACE) and acetonitrile (ACN) (bottom) on 17 March. (c) The observed SO_4^{2-} mass concentration by HALO along with height-latitude variations on 17 March 2018 (d) The observed OA mass concentration by HALO along with height-latitude variations on 17 March 2018 (e) Result of the HYSPLIT model backward trajectory analysis started at the location of the HALO flight path at 02:00, 04:00, 06:00, 09:00 UTC on 17 March 2018.

1062 (c)

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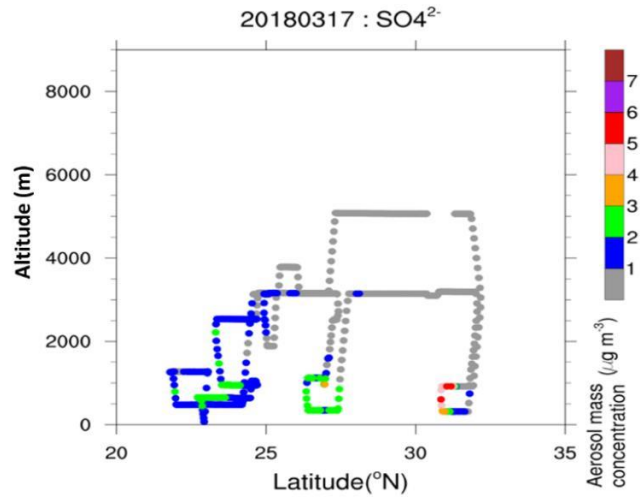
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1074 (d)

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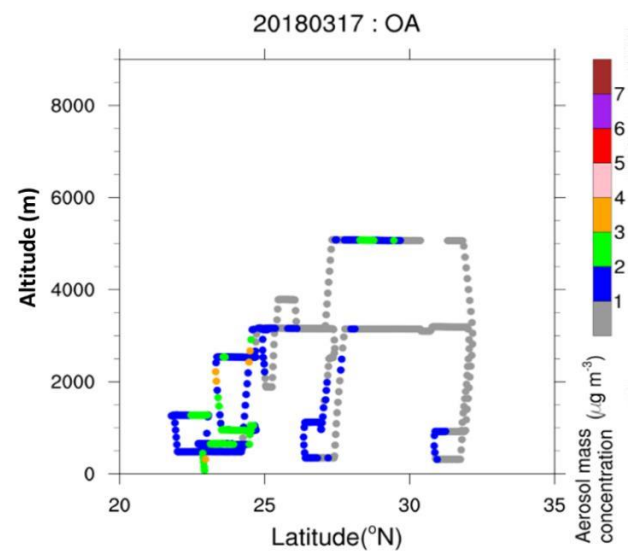
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1086 (e)

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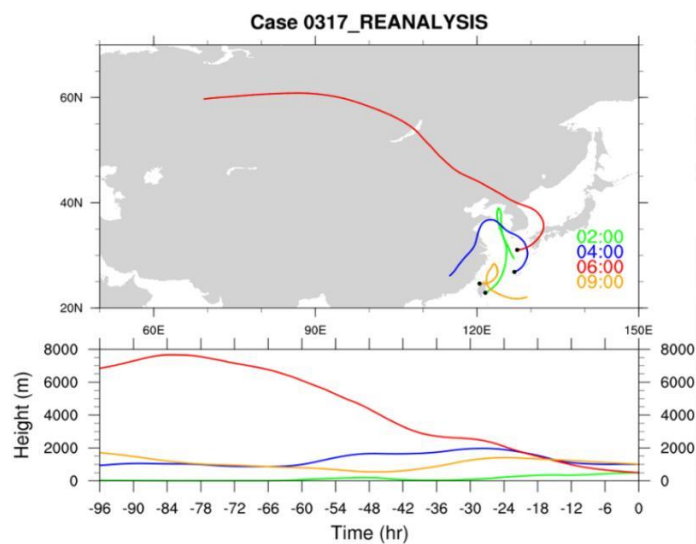
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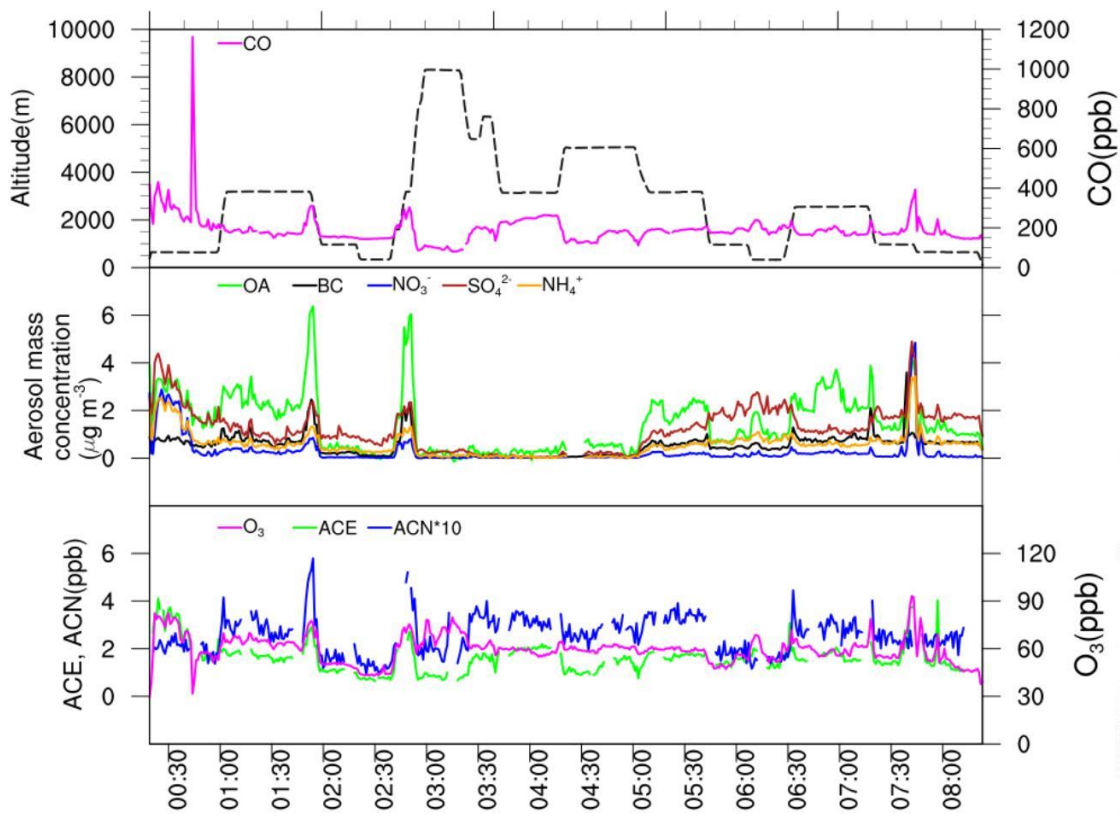
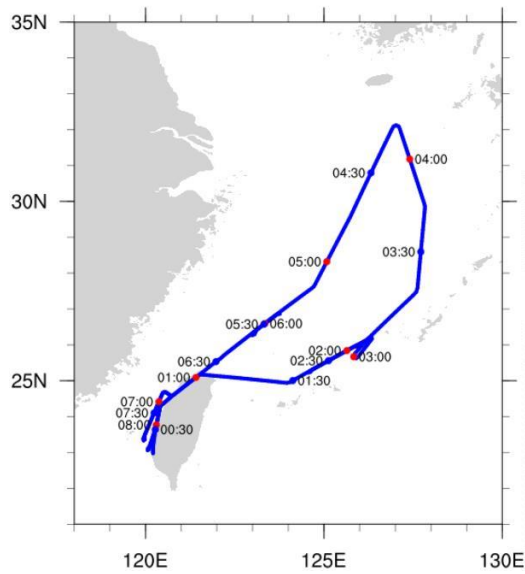
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1098 Figure 5 c-e

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11100 (a)
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 11112 (b)



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Figure 6 (a) The HALO flight and detailed locations on 19 March. (b) Flight altitude and 1-min mean of observed concentrations for CO (upper), Organic aerosol (OA), BC aerosol (BC), SO_4^{2-} , NO_3^- , NH_4^+ (middle), O_3 , acetone (ACE) and Acetonitrile (ACN) (bottom) on 19 March 2018. (c) The observed SO_4^{2-} mass concentration by HALO along with height-latitude variations on 19 March 2018 (d) The observed OA mass concentration by HALO along with height-latitude variations on 19 March 2018 (e) Result of the HYSPLIT model backward trajectory analysis started at the location of the HALO flight path at 02:00, 04:00, 05:00, 07:00 UTC on 19 March 2018.

1122 (c)

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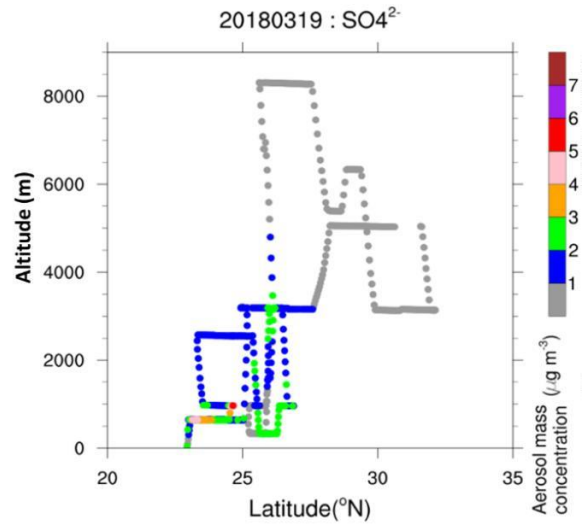
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1134 (d)

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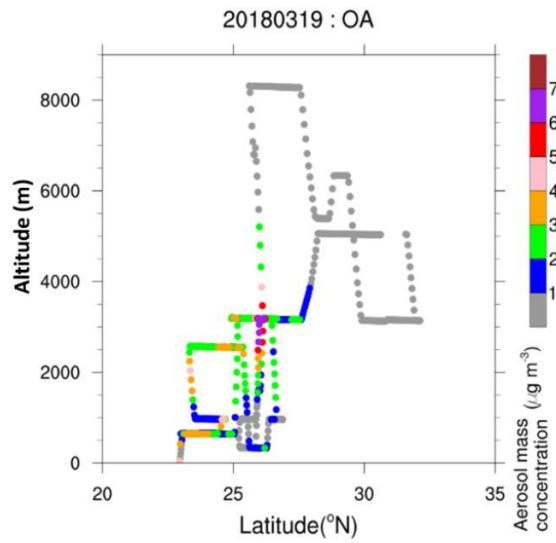
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1147 (e)

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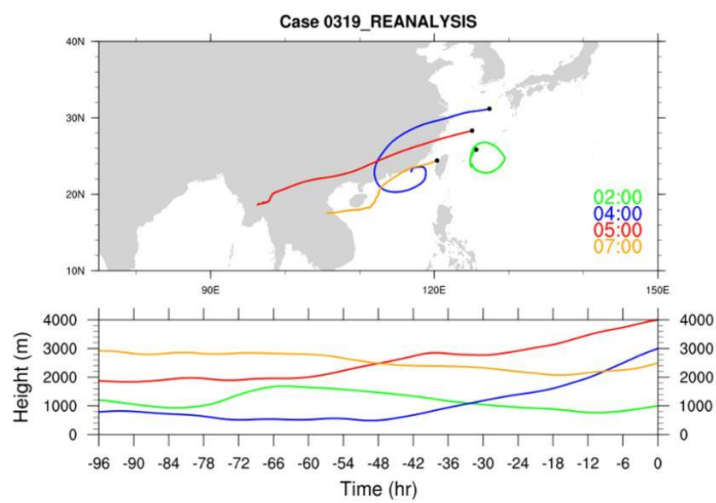
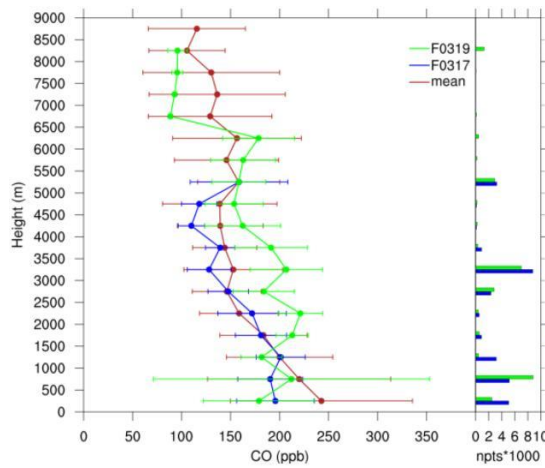
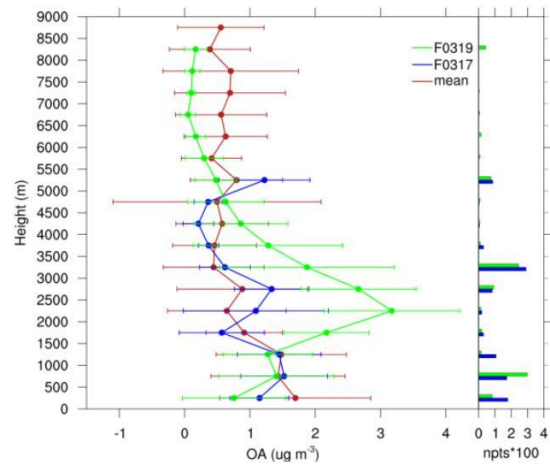


Figure 6 c-e

1160 (a)
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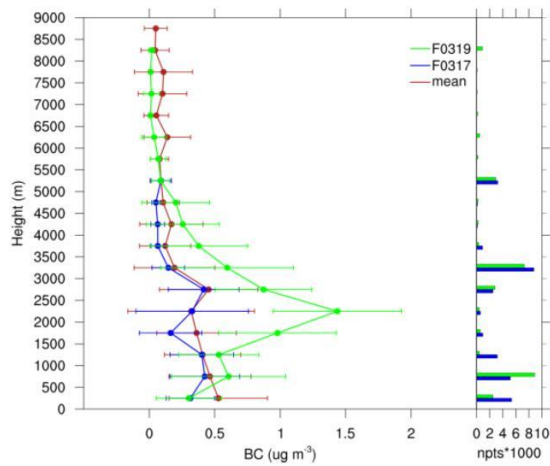


(b)

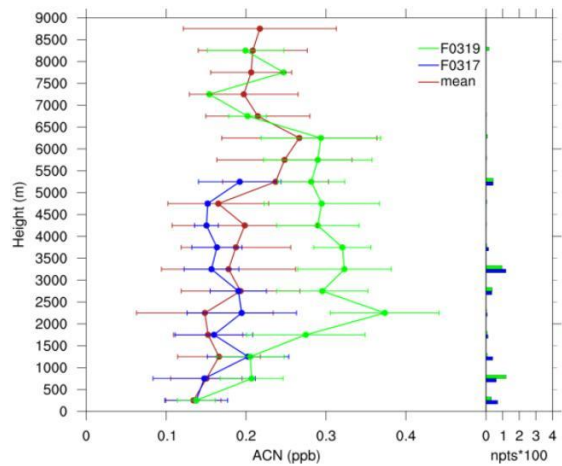


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1163 (c)



(d)



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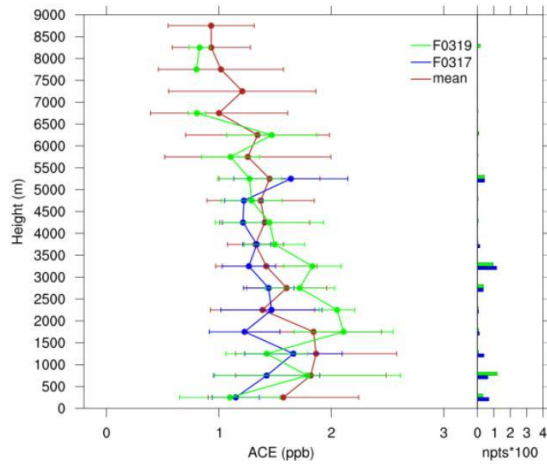
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1168 Fig.7 Observed vertical distribution calculated as 1-min mean and 500 m interval with
1169 one standard deviation of the concentrations for the mean profiles (red) (including 17,
1170 19, 22, 24, 26, 30 March, and 04 April 2018) and flights on 17 (blue) and 19 (green)
1171 March 2018. (a) CO (b) OA (c) BC (d) Acetonitrile (ACN) (e) Acetone (ACE) (f) O₃
1172 (g) J (O¹D) (h) NO_y. The number of data points is shown in the right panel.

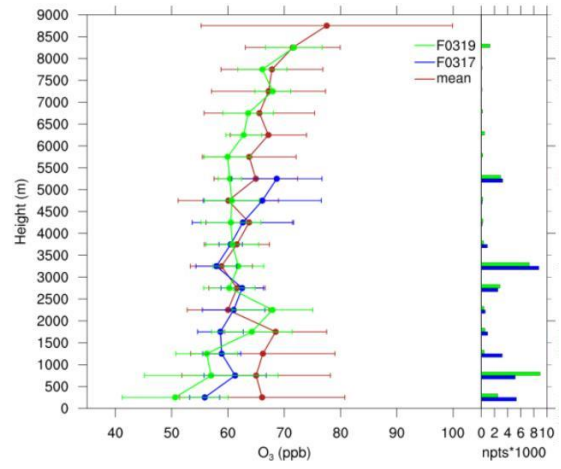
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1175 (e)

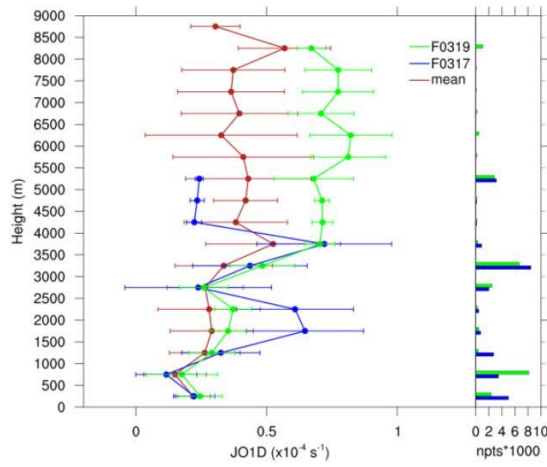


(f)

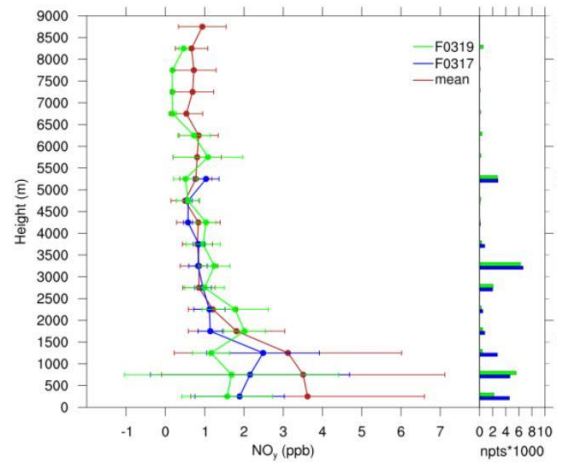


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1177 (g)



(h)



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1184 Figure 7 continued

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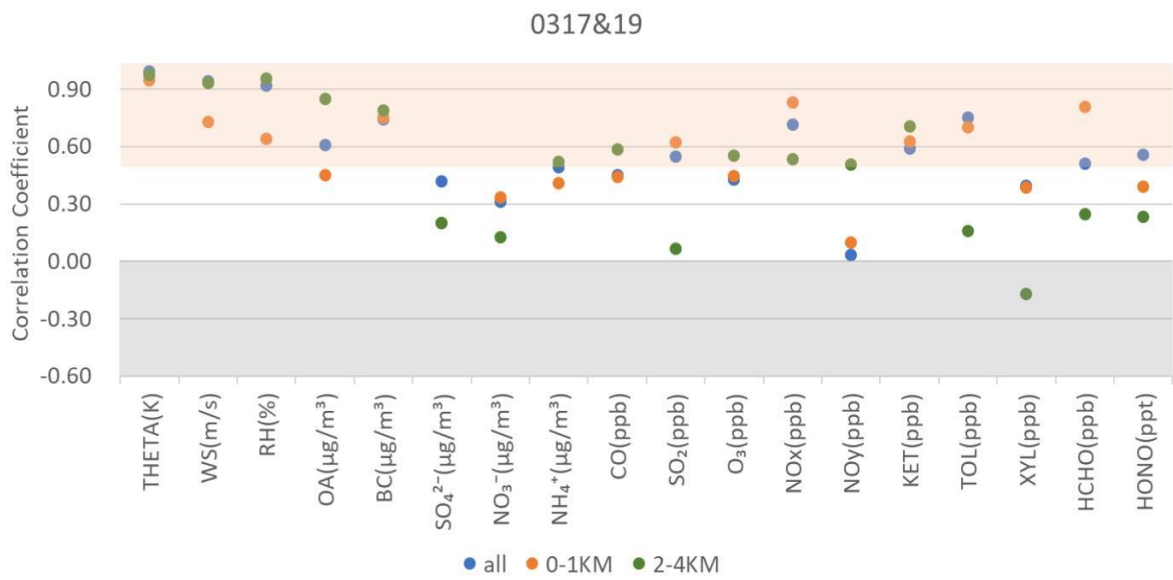
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1196 Fig. 8 Correlation Coefficient (R) between observation and simulation along with the
1197 HALO flights at the elevations 0-1 km, 2-4 km, and the whole track (all) on 17 and 19
1198 March 2018.

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1218 (a)

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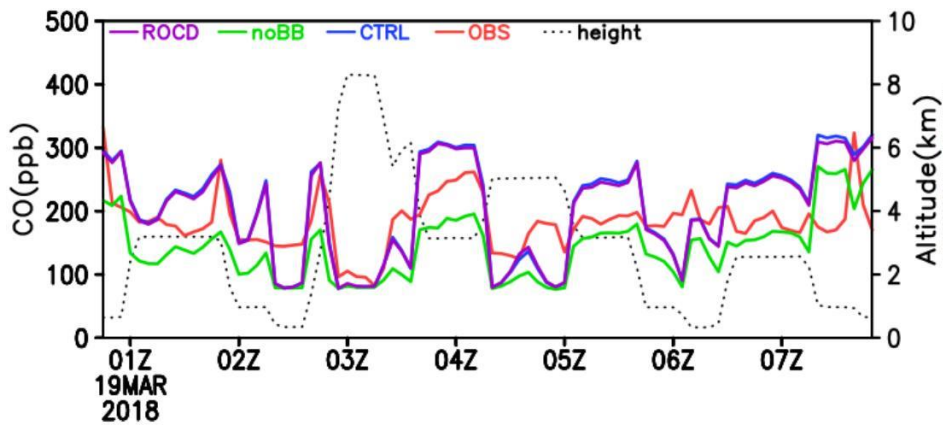
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1229 (b)

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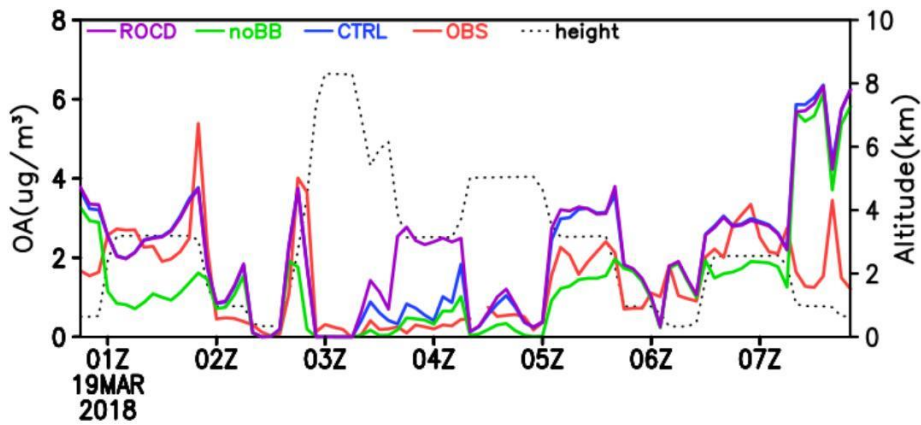
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1240 (c)

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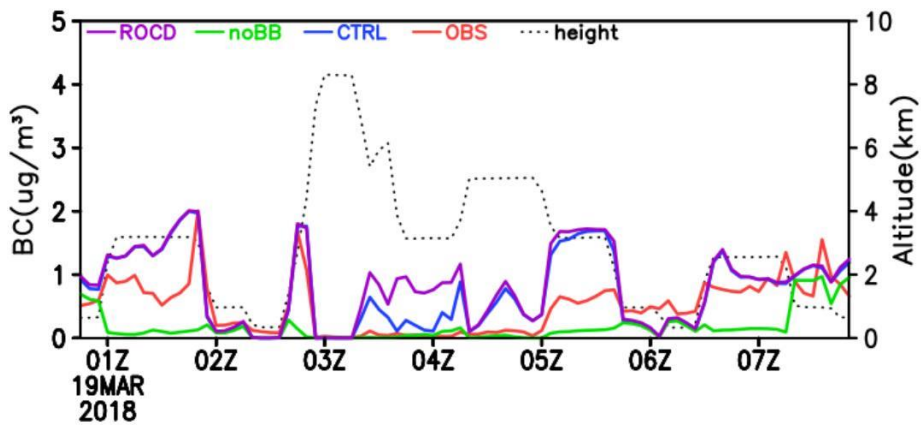
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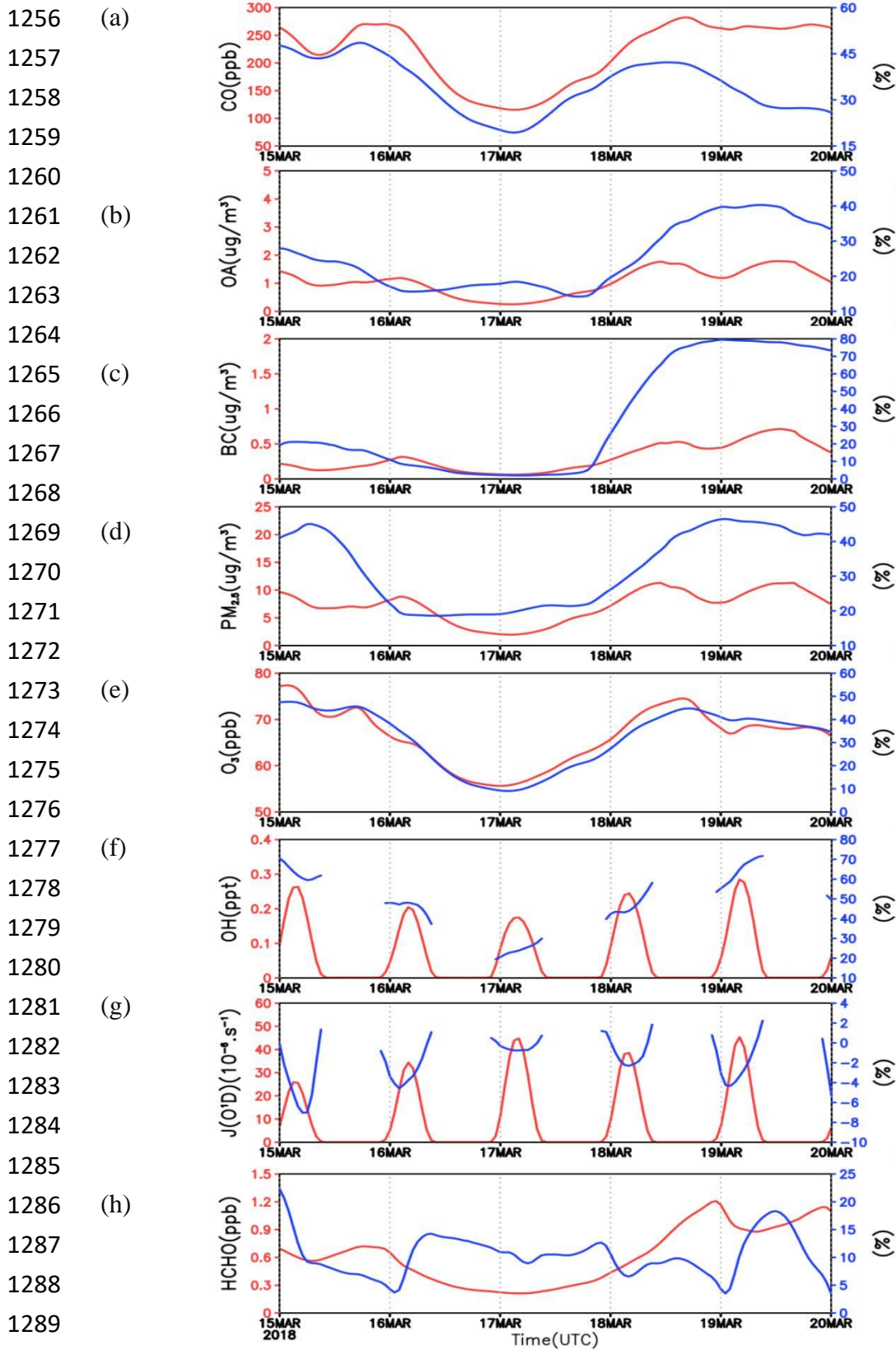
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1252 Fig.9 Observed (OBS, red) and simulated concentration (CTRL, blue), and the simulation

1253 without indirect effect (ROCD, purple), without BB emission (noBB, green) along with the

1254 flight altitude for (a) CO(ppb) (b) OA($\mu\text{g m}^{-3}$) (c) BC ($\mu\text{g m}^{-3}$) on 19 March 2018.

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1291 Figure 10 Hourly variation of simulated mean concentration (red) and contributed by
 1292 BB (%) (blue) between 2 km and 4 km over the region ECSA in Fig.1a during 15-19
 1293 March 2018. (a) CO (b) OA (c) BC (d) PM_{2.5} (e) O₃ (f) OH (g) J(O¹D), and (h) HCHO

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1295 (a)

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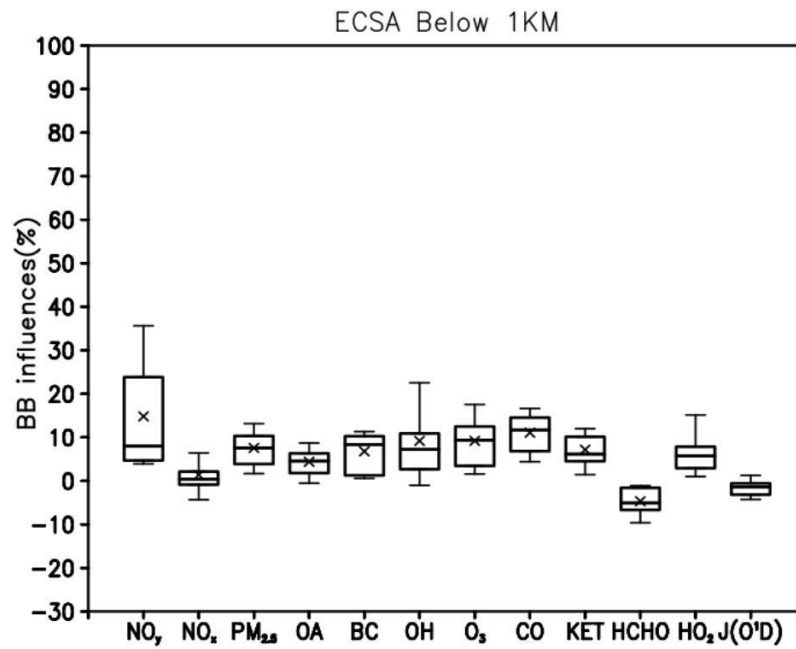
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1310 (b)

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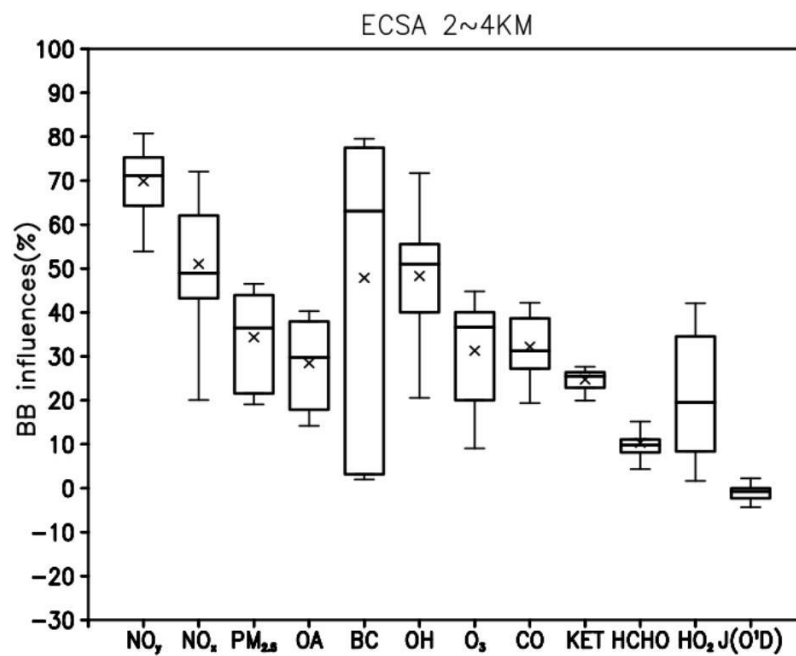
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1327 Figure 11 Box plots of simulated BB influences (%) on NO_y, NO_x, PM_{2.5}, OA, BC, OH,

1328 O₃, CO, KET, HCHO, HO₂, and J(O¹D) over the region ECSA in Fig. 1a on 17 and 19

1329 March 2018. (a) below 1 km, (b) between 2 km and 4 km

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1332 (a)

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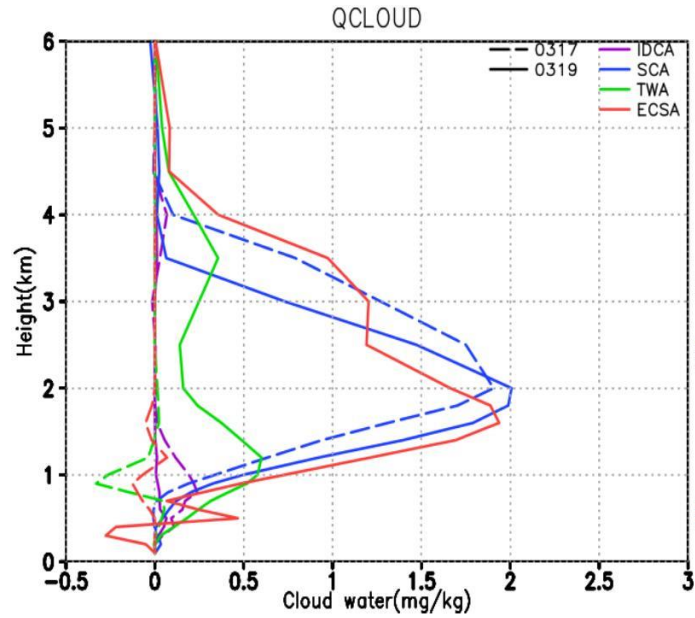
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1347 (b)

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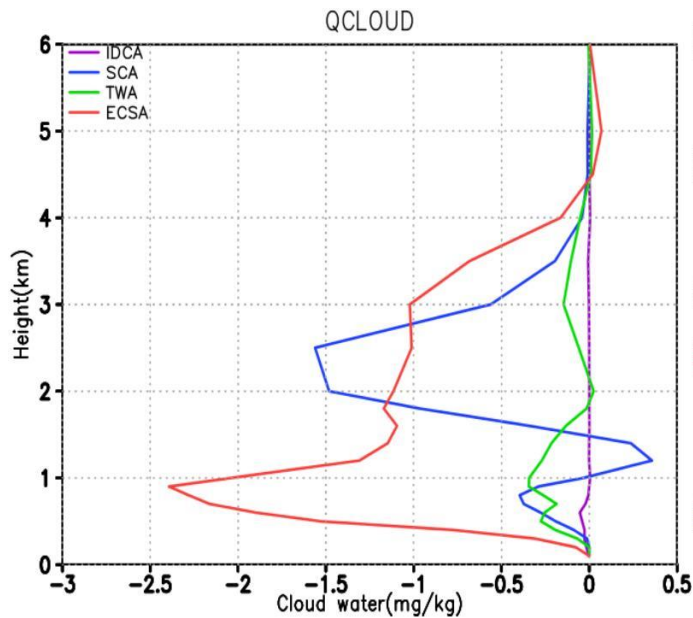
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1364 Figure 12 (a) Simulated vertical distribution of BB influences on cloud water difference

1365 between with and without BB emission on 17 (dash) and 19 (solid) March 2018. (b)

1366 Simulated vertical distribution of cloud water difference between with and without

1367 indirect effect in the model on 19 March 2018.

1368 Regions include IDCA, SCA, TWA, and ECSA as shown in Figure 1a.

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