- 1 Observation of SOA tracers at a mountainous site in Hong Kong: chemical characteristics,
- 2 origins and implication on particle growth
- X.P. Lyu<sup>1</sup>, H. Guo<sup>1\*</sup>, H.R. Cheng<sup>2\*\*</sup>, X.M. Wang<sup>3</sup>, X. Ding<sup>3</sup>, H.X. Lu<sup>1</sup>, D.W. Yao<sup>1</sup>, and C.
  Xu<sup>1</sup>
- <sup>5</sup> <sup>1</sup> Department of Civil and Environmental Engineering, The Hong Kong Polytechnic University,
- 6 Hong Kong
- <sup>7</sup> <sup>2</sup> Department of Environmental Engineering, School of Resource and Environmental Sciences,
- 8 Wuhan University, Wuhan, China
- <sup>3</sup> State Key Laboratory of Organic Geochemistry, Guangzhou Institute of Geochemistry, Chinese
- 10 Academy of Sciences, Guangzhou, China
- 11 Corresponding author: H. Guo (ceguohai@polyu.edu.hk); H.R. Cheng (chenghr@whu.edu.cn)
- 12

Abstract: Secondary organic aerosol (SOA) is an important constituent of airborne fine particles. 13  $PM_{2.5}$  (particles with aerodynamic diameters  $\leq 2.5 \ \mu m$ ) samples were collected at a mountainous 14 site in Hong Kong in autumn of 2010, and analyzed for SOA tracers. Results indicated that the 15 concentrations of isoprene SOA tracers (54.7 $\pm$ 22.7 ng/m<sup>3</sup>) and aromatics SOA tracers (2.1 $\pm$ 1.6 16 ng/m<sup>3</sup>) were on relatively high levels in Hong Kong. Secondary organic carbon (SOC) derived 17 18 from isoprene, monoterpenes, sesquiterpenes and aromatics was estimated with the SOA tracer based approach, which constituted  $0.35\pm0.15 \ \mu\text{g/m}^3$  (40.6±5.7%),  $0.20\pm0.03 \ \mu\text{g/m}^3$  (30.4±5.5%), 19  $0.05\pm0.02 \ \mu g/m^3$  (5.6±1.7%) and  $0.26\pm0.20 \ \mu g/m^3$  (21.3±8.2%) of the total estimated SOC. 20 Biogenic SOC ( $0.60\pm0.18 \ \mu g/m^3$ ) dominated over anthropogenic SOC ( $0.26\pm0.20 \ \mu g/m^3$ ) at this 21 site. In addition to the total estimated SOC ( $17.8\pm4.6\%$  of organic carbon (OC) in PM<sub>2.5</sub>), 22 primary organic carbon (POC) emitted from biomass burning also accounted for a considerable 23 24 proportion of OC (11.6±3.2%). Insight into the OC origins found that regional transport significantly (p < 0.05) elevated SOC from 0.37±0.17 to 1.04±0.39 µg/m<sup>3</sup>. Besides, SOC load 25 could also increase significantly if there was influence from local ship emission. Biomass 26 burning related POC in regional air masses ( $0.81\pm0.24 \text{ }\mu\text{g/m}^3$ ) was also higher (p<0.05) than that 27 in samples affected by local air  $(0.29\pm0.35 \ \mu g/m^3)$ . Evidences indicated that SOA formation was 28 closely related to new particle formation and the growth of nucleation mode particles, while 29 biomass burning was responsible for some particle burst events in Hong Kong. This is the first 30 SOA study in afforested areas of Hong Kong. 31

Keywords: Secondary organic aerosol, SOA tracer, biogenic SOA, regional transport, particlegrowth

## 34 1. Introduction

35 Atmospheric aerosol has been well recognized to affect global climate change (Stocker et al,

2013), human health (Dockery et al., 1993; Pope III and Dockery, 2006), visibility (Appel et al.,

1985) and sustainability of economy (Chameides et al., 1999). Secondary organic aerosol (SOA)

has been identified to play critical roles in these effects (Maria et al., 2004; Volkamer et al., 2006;

39 Baltensperger et al., 2008), thus receiving sufficient attentions in recent years.

So far, the scientific community has reached a consensus that volatile organic compounds (VOCs) 40 from biogenic emissions and anthropogenic aromatics are key precursors of SOA (Forstner et al., 41 1997; Claeys, et al., 2004). In global scale, biogenic SOA is thought to be the greatest constituent 42 43 of SOA, due to the worldwide largest emission of biogenic VOCs (e.g., isoprene, monoterpenes and sesquiterpenes) and formation of biogenic SOA spanning a wide range of conditions (the 44 level of NO<sub>x</sub>, humidity and aerosol acidity) (Kroll et al., 2006). However, anthropogenic SOA 45 has also been found to be significant in urban areas (Volkamer et al., 2006). Furthermore, upon 46 47 the findings that biogenic SOA correlates well with the indicators of anthropogenic emissions (Goldstein et al., 2009; Hoyle et al., 2011), it is believed that man-made air pollutants promote 48 49 the formation of biogenic SOA, in addition to serving as SOA precursors themselves. These promoting effects at least include forming aerosol seeds, catalyzing photooxidation and 50 51 transformation of biogenic VOCs and their oxidation products and changing the reaction pathways (e.g., atmospheric fate of isoprene in low- and high-NO<sub>x</sub> environments) (Hoyle et al., 52 53 2011). It is believed that SOA is a collection of hundreds to thousands of organic chemicals

54 featuring relatively low volatilities. To understand SOA formation and explore the potential 55 sources, SOA speciation is of great necessity. However, due to the difficulty in chemical analysis

of SOA tracers, the chemical compositions of SOA are far from being well understood. The traditional analysis by gas chromatography-mass spectrometer detector (GC-MSD) generally

58 quantifies a total of less than 20 organic compounds in particles (Offenberg et al., 2007;

59 Kleindienst et al., 2007). Although some advanced instruments have been developed nowadays,

such as aerosol mass spectrometry and thermal desorption aerosol gas chromatography, they are

either fragment-based or highly dependent upon the skills and knowledge of users (Williams et

al., 2014). Instead, SOA tracer based approach is a simplified method widely used to estimate the
amount, precursors and sources of SOA (Kleindienst et al., 2007; Ding et al., 2012).

The SOA tracer based approach applies the laboratory obtained ratios between the sum of 64 specific SOA tracers and total mass of SOA (or secondary organic carbon (SOC)) produced from 65 individual (group of) species to the field measured SOA tracers (Kleindienst et al., 2007), 66 roughly estimating SOA (SOC) derived from an individual VOC or VOC group. Table S1 67 summarizes the VOC precursors, corresponding SOA tracers and the ratios between SOA tracers 68 and SOA (or SOC), as reported by Kleindienst et al. (2007). The drawbacks of this method are 69 obvious. For instance, it is controversial whether the laboratory obtained ratios can be directly 70 applied to the field measured SOA tracers. However, it provides a feasible approach to estimate 71 the SOA concentration, which is especially helpful in the cases of not well knowing the SOA 72 compositions. More importantly, Offenberg et al. (2007) confirmed that the SOA tracer based 73 approach was reliable through comparing with the results obtained from  ${}^{14}C$  contents. 74

Hong Kong, one of the most developed regions in East Asia, has a total territory area of 75 ~ $1.1 \times 10^3$  km<sup>2</sup> and a total population of ~7 million. Despite high population density, it keeps 24 76 77 country parks and a vegetation coverage rate of 70%. Evergreen broadleaf trees are common in Hong Kong. Guenther et al. (2006) suggested that shrubs are large emitters of isoprene, a typical 78 and most abundant biogenic VOC. The total emission amount of biogenic VOCs in Hong Kong 79 is estimated as  $8.6 \times 10^3$  ton C/a (Tsui et al., 2009). Field measurements also revealed that 80 81 isoprene in Hong Kong is relatively high (300-400 pptv) (Guo et al., 2007, 2012a). In contrast, aromatics are largely emitted from vehicular exhaust and solvent usage in Hong Kong. For 82 83 example, toluene is one of the most abundant aromatics, with the level of 3-6 ppby (Guo et al., 2004; Ho et al., 2004). Therefore, local emission of VOCs has a great potential forming SOA in 84 85 Hong Kong. In addition to local SOA formation, regional transport is inevitable due to severe air pollution in upwind direction of Hong Kong (i.e., inland Pearl River Delta (PRD) region). 86 Studies confirmed that Hong Kong received SOA produced by isoprene and toluene from inland 87 PRD region (Hu et al., 2008). However, the previous SOA studies in Hong Kong all focused on 88 urban areas, which are not enough to understand the abundance, compositions and sources of 89 90 SOA in low-altitude mountainous area where both anthropogenic and biogenic emissions are important. 91

92 In this study, SOC derived from isoprene, monoterpenes, sesquiterpenes and aromatics at a 93 mountainous site in Hong Kong were estimated using a SOA tracer based approach. Local and 94 regional contributions to SOC and biomass burning related POC were determined. Furthermore, 95 in combination with particle size distribution simultaneously monitored by a scanning mobility 96 particle sizer (SMPS), the relationships between SOA formation and particle growth were 97 examined.

98 2. Methodology

#### 99 **2.1 Sample collection**

Hong Kong, a coastal city surrounded by the South China Sea (SCS) to the east and south, is 100 located in southern China. Under the dominance of subtropical monsoon climate, Hong Kong 101 receives northerly winds originating from the heavily polluted PRD region in cool seasons 102 (October-March), while prevailing southerly winds bring in clean air from SCS in warm seasons 103 (April-August) (So and Wang, 2003). The sampling site (22.405° N, 114.118° E, 640 m a.s.l) was 104 set up at the mountainside of the highest mountain in Hong Kong (Mt. Tai Mo Shan with the 105 maximum altitude of 957 m, TMS). The vegetation on this mountain mainly includes Acacia 106 107 confusa, Lophotemon confertus, Machilus chekiangensis and Schima superba below 550 m, while it turns to shrubs and grasses above 550 m (Guo et al., 2012b). As demonstrated in 108 109 previous studies (Guo et al., 2013; Ling et al., 2014), regional transport of air pollutants from inland PRD, mesoscale circulation (mountain-valley breezes) and in situ atmospheric chemistry 110 111 are the main processes that significantly influence air quality at this site. Figure 1 shows the geographical locations of the sampling site and air quality monitoring stations (AQMSs) of Hong 112 113 Kong Environmental Protection Department (HKEPD).

From September 7 to November 26, 2010, a total of 19  $PM_{2.5}$  samples were collected. The 114 115 instrument was an Anderson high volume PM2.5 sampler, with a flow rate of 750 L/min. Prebaked A4 size quartz fiber filters were used to collect the samples, which were stored in the 116 refrigerator at -18 °C after sampling. Generally, each filter sampling lasted for 45-55 hours, 117 118 except for the cases that the instrument stopped abnormally on some days due to thunderstormcaused power outages. Table S2 lists the sample IDs, start and end dates & times. The origins of 119 120 air masses, as distinguished by wind field and ratio of  $SO_2/NO_x$  (see section 3.3 for details), are also shown in the table. Additionally, the particle number concentrations and size distributions in 121 the size range of 5.5-350.4 nm in 44 size bins were monitored by a scanning mobility particle 122

sizer (SMPS) from October 27 to November 29. Detailed introductions about the operation
procedures of SMPS and data processing can be found in Guo et al. (2012a). We also collected
ambient VOC samples during September 28-November 21. Inorganic trace gases (SO<sub>2</sub>, CO, NO,
NO<sub>2</sub> and O<sub>3</sub>) and weather conditions were measured simultaneously with the PM<sub>2.5</sub> sampling.
Details about VOC sampling, VOC analysis and monitoring of trace gases are provided in Guo et
al. (2013) and Ling et al. (2014).



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Figure 1 Geographic locations of the sampling site (TMS), AQMSs of HKEPD and the container
terminal area where nine container terminals are located. Capital letters in the brackets are
abbreviations of the site/stations

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### 134 **2.2 Chemical analysis**

OC and element carbon (EC) in PM<sub>2.5</sub> samples were analyzed using the thermo-optical transmittance method recommended by National Institute for Occupational Safety and Health (NIOSH) (Birch, 1998). The concentrations of OC and EC are shown in Figure S1.

The method of SOA analysis was in line with that introduced by Ding et al. (2011). Briefly, the procedures for each sample include solvent extraction, derivation, analysis by GC-MSD, and identification and quantification of SOA tracers. 1/8 of each filter was extracted three times by sonication in the solvent of 40 mL of 1:1 (v/v) dichloride methane (DCM)/methanol mixture. Prior to extraction, the internal standards (hexadecanoic acid-D<sub>31</sub>, phthalic acid-D<sub>4</sub> and

levoglucosan- ${}^{13}C_6$ ) were spiked into the samples. The three-time extracts of each sample were 143 combined, filtered and concentrated to ~2 mL, which was further divided into two parts for 144 methylation and silvlation derivation, respectively. In methylation derivation, the extract 145 experienced a gentle nitrogen blow to dryness, and subsequent addition of 200 µL of DCM, 10 146  $\mu$ L of methanol and 300  $\mu$ L of freshly prepared diazomethane. Then, it was kept in room 147 temperature for one hour to derivatize acids to methyl esters, after which the sample was blown 148 to 200  $\mu$ L and used for analysis of some  $\alpha$ -pinene SOA tracers (Pinonic acid, pinic acid and 3-149 methyl-1,2,3-butanetricarboxylic acid). The silvlation reagent was 100 µL of pyridine and 200 150 uL of N.O-bis-(trimethylsilyl)-trifluoroacetamide (BSTFA) plus 1% trimethylchlorosilane 151 (TMCS). Differently, the derivation was carried out in an oven at 70  $^{\circ}$ C for one hour. One  $\alpha$ -152 pinene SOA tracer (3-hydroxyglutanic acid) and tracers for isoprene SOA, sesquiterpenes SOA, 153 and toluene SOA were analyzed from the silvlated sample. An Agilent 5973N GC/MSD was 154 employed to do the analysis. The identification of SOA tracers was based on the comparison of 155 mass spectra with previous studies, and their retention time in GC chromatogram with other 156 known compounds as the references. Pinonic acid and pinic acid were quantified by authentic 157 158 standards. However, due to lack of standards, other  $\alpha$ -pinene SOA tracers, isoprene SOA tracers,  $\beta$ -Caryophyllenic acid and 2,3-dihydroxy-4-oxopentanoic acid were quantified using pinic acid 159 (PA), erythritol, octadecanoic acid and azelaic acid, respectively. The detection limits (DLs) for 160 pinonic acid, pinic acid, erythritol, octadecanoic acid and azelaic acid were 0.05, 0.07, 0.06, 0.09, 161 and 0.11 ng/m<sup>3</sup>, respectively. Levoglucosan was also quantified with the DL of 0.15 ng/m<sup>3</sup>. The 162 SOA tracers analyzed in this study are highlighted in Table S1. 163

# 164 **2.3 Quality assurance and quality control**

In this study, internal standards were not spiked on the filters before sampling, to avoid their 165 166 influences on OC analysis. According to the saturation concentrations of SOA tracers calculated by Ding et al. (2016), SOA tracers analyzed in this study were of low volatilities, except for 167 pinonic acid. Therefore, pinonic acid in airborne PM<sub>2.5</sub> might be underestimated due to the blow-168 off effect during sampling, which however should not be significant to other SOA tracers. 169 170 Recovery target compounds were spiked in the samples before analysis of SOA tracers. The recovery rates were 104±2%, 68±13%, 62±14%, 78±10%, 81±9% and 87±4% for pinonic acid, 171 pinic acid, erythritol, octadecanoic acid, azelaic acid and levoglucosan, respectively. Since 172

internal standards were added into the samples prior to analysis, we did not use the recoveryrates to correct the concentrations of SOA tracers.

## 175 2.4 Processing of SMPS data

- 176 In this study, particles measured by SMPS were divided into nucleation (5.5-24.7 nm), Aitken
- 177 (24.7-101.4 nm) and accumulation modes (101.4-350.4 nm). Geometric mean diameter (GMD)
- 178 for nucleation mode particles (5.5-24.7 nm) was calculated using the following equations (Guo et
- 179 al., 2012a).

180 GRs = 
$$\frac{a_{GMD}}{d_t}$$
 (Equation (1))

- 181 GMD =  $e^{(\sum n_i ln^{d_i})/N}$  (Equation (2))
- 182  $\frac{d_N}{d_{\log} D_p} = \frac{\Delta N}{\log_{D_p^2} \log_{D_p^1}}$  (Equation (3))
- 183 where  $n_i$  is the particle number concentration in the *i*<sup>th</sup> bin with upper diameter of  $d_i$ , *N* represents 184 the total number concentration (cm<sup>-3</sup>).  $\Delta N$  is the particle number concentration in the size bin 185 with upper and lower limit diameter of  $D_p^2$  and  $D_p^1$ , respectively.
- 186 Condensation sink (CS), which describes the loss rate of vapor molecules and newly formed
- 187 particles on the pre-existing aerosol particles, was calculated as follows (Kulmala et al., 2005):

188 CS =  $2\pi D \int D_p \beta_m(D_p) n(D_p) dD_p = 2\pi D \sum_i \beta_{mi} D_{pi} N_i$  (Equation (4))

- where D is the diffusion coefficient of the condensing vapor,  $\beta_m$  is the transitional regime correction factor,  $D_p$  denotes the particle diameter, *n* and *N* represent the particle numbers.  $\beta_{mi}$ , D<sub>pi</sub> and N<sub>i</sub> are the specific values for a given size bin (*i*). Growth factor calculated according to Laakso et al. (2004) was used to calibrate the dry particle size measured by SMPS.
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# 194 **3. Results and discussion**

# **3.1 Concentrations of SOA tracers**

Table 1 shows the average concentrations of SOA tracers derived from different precursors at TMS. Isoprene SOA tracers were the most abundant  $(54.7\pm22.7 \text{ ng/m}^3)$ , followed by the tracers generated from monoterpenes  $(26.3\pm4.5 \text{ ng/m}^3)$ , aromatics  $(2.1\pm1.6 \text{ ng/m}^3)$  and sesquiterpenes  $(1.1\pm0.4 \text{ ng/m}^3)$ . Figure 2 compares the concentrations of SOA tracers between previous studies and the present study. The number and species of SOA tracers for the same precursor were the same. For the cases that total concentration of monoterpenes SOA tracers was given (*e.g.*, Offenberg et al., 2011), a factor was applied to the total concentration to roughly estimate the 203 sum of monoterpenes SOA tracers with the number and species identical to this study (see section S1 and Figure S2 in the Supplement). It was found that isoprene SOA tracers at TMS 204 were on relatively high level (54.7 $\pm$ 22.7 ng/m<sup>3</sup>), comparable to or even higher than those 205 detected in the forest (61.4±30.4 ng/m<sup>3</sup> in Kleindienst et al. (2007) and 39.0±17.2 ng/m<sup>3</sup> in 206 207 Offenberg et al. (2011)). In fact, isoprene at TMS was relatively low (2-517 pptv), the high isoprene SOA tracers indicated that biogenic SOA formation at this site might be enhanced by 208 209 anthropogenic emissions, e.g., sufficient aerosol seeds from SO<sub>2</sub>-related new particle formation (Guo et al., 2012a). Certainly, regional transport could also contribute to high isoprene SOA 210 tracers, as reported by Hu et al. (2008). For the anthropogenic SOA, the tracer produced by 211 aromatics (2,3-dihydroxy-4-oxopentanoic acid) was noticeable. Although the average 2,3-212 dihydroxy-4-oxopentanoic acid in this study  $(2.1\pm1.6 \text{ ng/m}^3)$  was not significantly higher than 213 that in other studies (p>0.05), its maximum value reached 13.5 ng/m<sup>3</sup>. Given abundant aromatics 214 in the atmosphere of Hong Kong and mesoscale circulations between TMS and the surrounding 215 urban areas, local formation could be an important factor contributing to high aromatics SOA at 216 Moreover, it was reported that 2,3-dihydroxy-4-oxopentanoic acid at an urban 217 TMS. background site in inland PRD was very high (13.1 ng/m<sup>3</sup>) (Ding et al., 2012). Hence, regional 218 transport might be partially responsible for the relatively high aromatics SOA tracer in Hong 219 Kong. SOA tracers derived from monoterpenes and sesquiterpenes were on moderate levels, 220 compared to other studies. 221

222 3-Methyl-1,2,3-butanetricarboxylic acid and 3-hydroxyglutanic acid, formed from monoterpenes oxidation in the presence of NO<sub>x</sub> (Claevs et al., 2007; Eddingsaas et al., 2012), dominated the 223 224 measured monoterpenes SOA tracers, consistent with those at an urban background site in inland PRD (Ding et al., 2012). Pinonic acid and pinic acid, formed from OH oxidation of  $\alpha$ -pinene in 225 226 NO<sub>x</sub> free environment or ozonolysis of  $\alpha$ -pinene (Eddingsaas et al., 2012), were also higher than those in urban areas of Hong Kong (almost below DLs) (Hu et al., 2008). Since NO<sub>x</sub> was at the 227 magnitude of several ppby, the higher pinonic acid and pinic acid were likely due to the higher 228  $O_3$  (69.2±2.4 ppbv at TMS and 30.8±2.6 ppbv at the mountain foot) at this mountainous site. In 229 230 addition, other tracers including 3-hydroxy-4,4-dimethylglutaric acid, 3-isopropylpentanedioic 231 acid, 3-acetylpentanedioic acid and 3-acetylhexanedioic acid, which were not measured in this study, accounted for 46.9±4.0% of the total amount of tracers according to Kleindienst et al. 232 (2007). Of the isoprene SOA tracers, 2-methylthreitol and 2-methylerythritol are representative 233

products formed through the photooxidation of isoprene hydroxyhydroperoxides and acid catalysis of epoxydiols of isoprene in low NO<sub>x</sub> environment when RO<sub>2</sub> reacting with HO<sub>2</sub> dominated the loss of RO<sub>2</sub> (Claeys et al., 2004; Surratt et al., 2010). In this study, they accounted for  $83.5\pm8.1\%$  of the total isoprene tracers. However, NO<sub>x</sub> was not low enough and RO<sub>2</sub> reacting with NO was the main sink of RO<sub>2</sub> (Ling et al., 2014), implying that other mechanisms might enhance the formation of 2-methyltetrols (sum of 2-methylthreitol and 2-methylerythritol) at this site, such as acid catalysis (Surratt et al., 2007).

Table 1 Average concentrations of SOA tracers derived from monoterpenes, isoprene,
sesquiterpenes and aromatics (mean±95% confidence interval (C.I.)).

VOC precursor	SOA tracer	Concentration (ng/m <sup>3</sup> )
Monoterpenes	Pinonic acid	1.2±0.4
	Pinic acid	$2.0\pm0.6$
	3-methyl-1,2,3-butanetricarboxylic acid	16.0±3.5
	3-hydroxyglutanic acid	7.1±2.7
	Sum 1	26.3±4.5
Isoprene	2-Methylthreitol	14.7±6.6
	2-Methylerythritol	31.2±13.1
	2-methylglyceric acid	5.2±3.1
	cis-2-Methyl-1,3,4-trihydroxy-1-butene	$0.8 \pm 0.5$
	trans-2-Methyl-1,3,4-trihydroxy-1-	2.0±1.3
	butene	
	3-Methyl-2,3,4-trihydroxy-1-butene	0.7±0.3
	Sum 2	54.7±22.7
Sesquiterpenes	$\beta$ -Caryophyllenic acid	1.1±0.4
Aromatics	2,3-Dihydroxy-4-oxopentanoic acid	2.1±1.6

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Figure 2 Comparison of SOA tracers between previous studies and this study. <sup>1</sup>Kleindienst et al. (2007); <sup>2</sup>Offenberg et al. (2011); <sup>3</sup>Ding et al. (2012); <sup>4</sup>Hu et al. (2008); <sup>5</sup>Lewandowski et al. (2008); <sup>6</sup>Haddad et al. (2011); <sup>7</sup>this study.

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### 249 **3.2 Estimate of SOC**

Both the EC tracer method and SOA tracer method were used to estimate SOC (details about 250 251 both methods are provided in section S2 of the Supplement). To apply the SOA tracer method, 252 the sum of monoterpenes SOA tracers was scaled by a factor of 1.79 (see section S1 and Figure S2 for details). Figure S3 compares the SOC estimated by these two methods. SOC estimated by 253 EC tracer method (SOC<sub>EC tracer</sub>) were generally higher (p < 0.01) than those estimated by SOA 254 tracer method (SOC<sub>SOA tracer</sub>), except for samples TMS1, TMS3 and TMS8. Ding et al. (2012) 255 indicated that the EC tracer method might overestimate SOC, because it blended some primary 256 OC (POC) from biomass burning with SOC. This inference was supported by the good 257 correlation between the difference of SOC estimated by these two methods (SOC<sub>EC tracer</sub>-SOC<sub>SOA</sub> 258 tracer) and levoglucosan, a tracer of biomass burning. 259

In addition to levoglucosan, methyl chloride (CH<sub>3</sub>Cl) is an indicator of biomass burning 260 261 (Rudolph et al., 1995). Figure 3 plots the correlation between levoglucosan and CH<sub>3</sub>Cl at TMS, as well as that between SOC<sub>EC tracer</sub>-SOC<sub>SOA tracer</sub> and levoglucosan. As expected, levoglucosan 262 fairly correlated with CH<sub>3</sub>Cl ( $R^2$ =0.54), further confirming the reliability of levoglucosan as the 263 tracer of biomass burning. Consistent with Ding et al. (2012), good correlation was found 264 between  $SOC_{EC \text{ tracer}}$ -SOC<sub>SOA tracer</sub> and levoglucosan (R<sup>2</sup>=0.52). This verified that the EC tracer 265 method overestimated SOC due to the interference of biomass burning. Exceptionally, SOC<sub>EC</sub> 266 267 tracer was remarkably lower than  $SOC_{SOA tracer}$  for the samples TMS1, TMS3 and TMS8 (p < 0.05). 268 Further investigation found that on the sampling days of TMS1 and TMS3, the daily maximum hourly  $O_3$  mixing ratio, referred to as peak  $O_3$ , was extremely high, with the values being the 269

270 second (154.4 ppbv) and first (163.4 ppbv) highest among the 19 filter samples, respectively, while during the sampling period of TMS8, the peak O<sub>3</sub> also reached 94.8 ppby. In contrast, the 271 272 average of peak O<sub>3</sub> values on the sampling days of other PM<sub>2.5</sub> filter samples was only 77.6 ppbv. Since high O<sub>3</sub> levels generally imply strong photochemical reactivity, high productions of SOA 273 are also expected on these days. Hence, the higher SOC<sub>SOA tracer</sub> values observed in TMS1, TMS3 274 and TMS8 were understandable. We also found that there was no correlation between SOC<sub>EC</sub> 275  $_{tracer}$  and peak O<sub>3</sub> (not shown), whereas SOC<sub>SOA tracer</sub> correlated well with peak O<sub>3</sub> (R<sup>2</sup>=0.68), as 276 shown in Figure 4, implying that the formation of secondary products, *i.e.*, SOA and O<sub>3</sub>, depends 277 upon the oxidative capacity of the atmosphere and they may also influence each other. Therefore, 278 the SOA tracer method was believed to be more reliable and thus adopted in this study. Hereafter, 279 SOC refers to SOC<sub>SOA tracer</sub>, unless otherwise specified. 280





283 <sub>tracer</sub> and levoglucosan.



Figure 4 Linear correlation between SOC<sub>SOA tracer</sub> and peak O<sub>3</sub>.

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287 Figure 5 shows SOC produced by different groups of VOCs estimated using the SOA tracer based approach, and biomass burning related POC (POC biomass burning). POC biomass burning was 288 estimated by 0.01 ( $\mu$ g/ng) × levoglucosan (ng/m<sup>3</sup>) (Lee et al., 2008). On average, SOC and POC 289 biomass burning constituted 0.86±0.31  $\mu$ g/m<sup>3</sup> (17.8±4.6%) and 0.67±0.22  $\mu$ g/m<sup>3</sup> (11.6±3.2%) of OC 290 291 (provided in Figure S1), respectively. The rest of OC were undetermined due to unknown sources and precursors of OC. The total SOC comprised anthropogenic (*i.e.*, aromatics) and 292 293 biogenic (*i.e.*, isoprene, monoterpenes and sesquiterpenes) SOC, with the fractions of 21.3±8.2%  $(0.26\pm0.20 \ \mu\text{g/m}^3)$  and  $78.7\pm8.2\%$   $(0.60\pm0.18 \ \mu\text{g/m}^3)$ , respectively. Although anthropogenic 294 SOC (ASOC) was significantly lower than biogenic SOC (BSOC) (p<0.05), ASOC had its 295 highest value of 1.71  $\mu$ g/m<sup>3</sup> in the sample TMS1, and BSOC reached its maximum in sample 296 297 TMS3 (2.03  $\mu$ g/m<sup>3</sup>). As mentioned earlier, the two samples were collected on the days with very high O<sub>3</sub>, indicating that aromatics and biogenic VOCs might be responsible for the high SOC in 298 TMS1 and TMS3, respectively. However, VOC samples were not simultaneously collected 299 during the collection of these two samples. Instead, the sample TMS12 was a good example 300 because ASOC was the second highest  $(1.2 \,\mu\text{g/m}^3)$ , and the daily average mixing ratio of toluene 301 coincidently reached 4.0 ppbv during the TMS12 sampling period, the highest value among all 302 VOC samples, which further confirmed the reliability of SOA tracer method in estimating SOC. 303 Among SOC derived from biogenic VOCs, isoprene made the highest contribution (54.2±5.3% 304 of BSOC and 42.7±5.9% of total SOC), followed by monoterpenes (37.9±4.6% and 30.4±5.5%, 305 306 respectively) and sesquiterpenes  $(7.9\pm2.7\% \text{ and } 5.6\pm1.7\%, \text{ respectively})$ .



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Figure 5 Concentrations of estimated SOC derived from monoterpenes, isoprene, sesquiterpenes
 and aromatics and POC <sub>biomass burning</sub>.

# 310 **3.3 Local and regional contributions to SOC**





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Figure 6 Hourly ratios of  $SO_2/NO_x$  for the air masses in different wind directions/speeds at TMS. Table 2 summarizes the concentrations of SOC and POC <sub>biomass burning</sub> in different categories of air masses. Note that since the samples TMS1 and TMS16 were influenced by both regional air and local ship emission, they were separately discussed later. It was found that SOC derived from

334 individual group/species and total SOC in regional air were all significantly (p < 0.05) higher than that in local air, suggesting that SOC at TMS was elevated by regional transport. Despite 335 336 possible influences of local ship emissions in TMS1 and TMS16, the concentrations of SOC and POC in these two samples were well within the ranges of those in regional air, except for 337 aromatics SOC (1.71  $\mu$ g/m<sup>3</sup>) in sample TMS1. The extremely high aromatics SOC in TMS1 338 might be caused by ship emissions which could be laden with high concentrations of aromatics. 339 Contradictorily, SOC derived from aromatics was remarkably lower in TMS16 (0.27  $\mu$ g/m<sup>3</sup>). 340 This discrepancy might be explained by the differences of fuel types and operating conditions of 341 the ship engines, which were repeatedly proved to affect the emission characteristics of ship 342 engines (Reda et al., 2014; Sippula et al., 2014). Similar to SOC, POC also showed a 343 significantly higher level in regional air  $(0.81\pm0.24 \text{ }\mu\text{g/m}^3)$  than that in local air  $(0.29\pm0.35 \text{ }\mu\text{g/m}^3)$ 344  $\mu g/m^3$ ) (p<0.05). Since the high concentration of POC <sub>biomass burning</sub> in regional air was partially 345 contributed by TMS16 (POC <sub>biomass burning</sub> =1.54  $\mu$ g/m<sup>3</sup>), a sample jointly influenced by regional 346 air and local ship emission, specific insight was given to this sample. Firstly, ship emission was 347 not likely to be the culprit of high POC biomass burning, as no study reported ship emission of 348 levoglucosan, the biomass burning tracer used to calculate POC biomass burning in this study. Instead, 349 we found that another sample under the influence of regional air (TMS15) had comparable POC 350 biomass burning (1.64 µg/m<sup>3</sup>). Furthermore, CH<sub>3</sub>Cl, another tracer of biomass burning, increased 351 noticeably under northerly winds in sample TMS16, indicating the regional transport of biomass 352 353 burning plumes into Hong Kong. In fact, nearly no fire spot in local Hong Kong was observed by the satellite during the sampling, compared to some open fires detected in upwind directions (see 354 Figure S5). Therefore, POC biomass burning in TMS16 was reasonably speculated to be elevated by 355 regional biomass burning plumes. Upon this inference, we concluded that regional transport 356 357 significantly contributed to PM<sub>2.5</sub>-bounded POC in Hong Kong.

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Table 2 Mean $\pm$ 95% C.I. of SOC and POC in different categories of air masses (Unit:  $\mu$ g/m<sup>3</sup>).

	Local air	Regional air	TMS1 <sup>*</sup>	TMS16 <sup>*</sup>
SOC (monoterpenes)	0.13±0.06	0.23±0.03	0.23	0.28
SOC (isoprene)	$0.16 \pm 0.07$	$0.42 \pm 0.18$	0.51	0.41
SOC (sesquiterpenes)	$0.01 \pm 0.01$	$0.06 \pm 0.02$	0.16	0.07
SOC (aromatics)	$0.07 \pm 0.04$	0.33±0.26	1.71	0.27

POC biomass burning	$0.29 \pm 0.35$	0.81±0.24	0.17	1.54
Total SOC	0.37±0.17	1.04±0.39	2.61	1.04

<sup>\*</sup> 95% C.I. is not available for a single sample.

## **361 3.4 Implications to particle formation and growth**

As an important constituent of airborne particles, SOA plays critical role in particle formation 362 and growth (Jang et al., 2003). The relationships between SOA formation and particle 363 364 formation/growth were investigated on three selected days (October 31, November 09 and November 19) when SOC were among the highest of all the samples and SMPS data were 365 available. Figures 7-9 show the evolutions of particle numbers, GMD of nucleation mode 366 particles, CS, the simulated sulfuric acid (SA) vapor, oxidized intermediates of aromatics (AOI), 367 oxidized intermediates of biogenic VOCs (BOI), and the measured inorganic trace gases. Details 368 about the modelling of SA vapor, AOI and BOI are provided in Section S3 and Table S3 in the 369 Supplement. For convenience of analysis, the hourly mixing ratios of biogenic VOCs (BVOCs), 370 aromatics and CH<sub>3</sub>Cl are presented in Figure S6. 371

In Figures 7-8, the number concentration of nucleation mode particles  $(N_{nuc})$  increased 372 substantially in the morning of October 31 (11:00-12:00) and November 09 (10:00-11:00), 373 followed by the increases of number of Aitken mode particles (NAit). Prior to the increases of 374 N<sub>nuc</sub>, the simulated SA vapor began to increase about 3 hours earlier. These were consistent with 375 the findings in Guo et al. (2012a) who reported that the increase of  $N_{nuc}$  was caused by new 376 particle formation (NPF) events occurred on both days, and SA vapor played important roles in 377 NPF. However, we further found that the oxidation intermediates of BVOCs (*i.e.*, BOI) also 378 increased slightly ahead of the rise of N<sub>nuc</sub>, which might suggest that the oxidation of BVOCs 379 also made some contributions to NPF. In fact, the involvement of BVOCs in NPF at this 380 381 afforested site has been speculated by Guo et al. (2012a), which is confirmed with the aid of model simulations in this study. 382

Moreover, we noticed that the nucleation mode particles experienced obvious growth with the rate of 1.9 nm/h (15:00-16:00) and 1.4 nm/h (14:00-16:00) in the afternoon of October 31 and November 09, respectively. Both growths occurred under the conditions of high  $N_{Ait}$  and high CS, differing from NPF events. Meanwhile,  $O_3$  was on high level, which meant strong oxidative capacity of the atmosphere. Correspondingly, the simulated SA vapor, AOI and BOI showed great increments simultaneously with or 1-2 hours earlier than the increase of GMD. It is 389 noteworthy that the significant increases of AOI and BOI were also attributable to the rapid 390 increases of VOCs (see Figure S6), in addition to strong atmospheric oxidative capacity. This 391 suggested that the photo-oxidation of VOCs also facilitated the growth of nucleation mode particles. The prompt responses of particle growth to the increments of oxidation products on 392 October 31 (rather than 1.5 hours' delay on November 09) were likely caused by the much more 393 abundant BOI (~22.7 pptv) than that on November 09 (~13.5 pptv). Besides, the lower initial 394 GMD before its increase on October 31 (~14 nm compared to ~16 nm on November 09) implied 395 higher surface area and subsequently quicker growth. Since the aforementioned days featured 396 high SOA, the roles of photo-oxidation of VOCs in the formation and growth of nucleation mode 397 particles might reflect the very initial stages of SOA formation. However, to better understand 398 399 the relationships between SOA formation and the formation/growth of particles, data with higher resolution and more comprehensive chemical information of SOA are crucially needed. 400



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402 Figure 7 Evolutions of particle numbers, GMD of nucleation mode particles, CS, simulated SA

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Figure 8 Evolutions of particle numbers, GMD of nucleation mode particles, CS, simulated SA
vapor, AOI, BOI and inorganic trace gases on November 09 (second sampling day of TMS14).

Similarly, the growth of nucleation mode particles was also observed on November 19 (15:00-408 409 16:00) (Figure 9). However, a distinct phenomenon was that the numbers of particles in nucleation, Aitken and accumulation modes all showed rapid increases simultaneously from 410 411 around 14:30 and reached the highest values at ~15:00, which indicated that the particles in 412 different modes shared a common source. Although SO<sub>2</sub> and the simulated SA vapor began to 413 increase 1.5 hours earlier, this could not be a NPF event, as the increase of NAit had no delay (no banana shape) and the CS was high. In addition, the levels of primary air pollutants (e.g., NO<sub>x</sub> 414 415 and CO) were high at the peak hours of all three-mode particle numbers. More importantly, we found that CH<sub>3</sub>Cl largely increased from 13:00 to 15:00 and reached the maximum at 15:00 (see 416 417 Figure S6). Therefore, we suspected that biomass burning might be responsible for the increase of particle numbers. This coincided with the high POC biomass burning  $(1.54 \ \mu g/m^3)$  in PM<sub>2.5</sub> sample 418 collected on this day (TMS16). Generally, particles emitted from biomass burning are in Aitken 419 and accumulation modes (Reid et al., 2005). However, in vicinity of fire, nucleation mode can 420 also exist (Janhall et al., 2010). Here, although nucleation mode particles increased in the particle 421

burst event,  $N_{nuc}$  was very low (maximum= $2.1 \times 10^3$  cm<sup>-3</sup>), indicating that this was not a local biomass burning and nucleation mode particles converted to larger size particles from the source region to this site. This was also consistent with the regional influence on this day identified by the wind fields (see section 3.3).



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Figure 9 Evolutions of particle numbers, GMD of nucleation mode particles, CS, simulated SA
vapor, AOI, BOI and inorganic trace gases on November 19 (sampling day of TMS16).

429

# 430 4. Conclusions

PM<sub>2.5</sub> samples were collected at a mountainous site in Hong Kong in autumn of 2010. Nine SOA 431 tracers in PM<sub>2.5</sub> were analyzed, which helped to understand the compositions and sources of 432 SOC in this study. Results indicated that isoprene made the highest contribution to SOC 433 formation at this site, followed by monoterpenes, aromatics and sesquiterpenes. Averagely, 434 biogenic SOC dominated over anthropogenic SOC. However, anthropogenic SOC cannot be 435 neglected, particularly under the influences of regional transport (e.g., 1.2  $\mu$ g/m<sup>3</sup> in sample 436 TMS12) and/or local ship emission (e.g.,  $1.7 \text{ }\mu\text{g/m}^3$  in sample TMS1). The simultaneous 437 observation of VOCs confirmed the role of aromatics in contributing to high concentrations of 438 439 anthropogenic SOC. In terms of SOC origins, regional transport caused nearly two-fold increase

440 of SOC, relative to local air. However, SOC load could also be significantly elevated by local ship emissions possibly containing abundant VOC precursors and SO<sub>2</sub>, which promoted SOC 441 442 formation. In addition, the regional air was generally characterized with high biomass burning 443 related POC, aggravating the PM<sub>2.5</sub>-bounded POC in Hong Kong. In combination with the SMPS data, we found that the formation of SOA (particularly the biogenic SOA) might be partially 444 responsible for the new particle formation and growth of nucleation mode particles. Primary 445 emissions, such as biomass burning, could cause particle burst events and lead to POC increases. 446 To our knowledge, this is the first SOA study carried out in low-altitude (640 m) mountainous 447 area of Hong Kong, where the air quality is under the combined influence of anthropogenic and 448 biogenic emissions. This study also demonstrates the urgency of data acquisition with more 449 comprehensive chemical information and higher time resolution in future SOA studies over this 450 region. 451

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- **1** Observation of SOA tracers at a mountainous site in Hong Kong: chemical characteristics,
- 2 origins and implication on particle growth
- X.P. Lyu<sup>1</sup>, H. Guo<sup>1\*</sup>, H.R. Cheng<sup>2\*\*</sup>, X.M. Wang<sup>3</sup>, X. Ding<sup>3</sup>, H.X. Lu<sup>1</sup>, D.W. Yao<sup>1</sup>, and C.
  Xu<sup>1</sup>
- <sup>5</sup> <sup>1</sup> Department of Civil and Environmental Engineering, The Hong Kong Polytechnic University,
- 6 Hong Kong
- <sup>7</sup> <sup>2</sup> Department of Environmental Engineering, School of Resource and Environmental Sciences,
- 8 Wuhan University, Wuhan, China
- <sup>3</sup> State Key Laboratory of Organic Geochemistry, Guangzhou Institute of Geochemistry, Chinese
- 10 Academy of Sciences, Guangzhou, China

11 Corresponding author: H. Guo (<u>ceguohai@polyu.edu.hk</u>); H.R. Cheng (<u>chenghr@whu.edu.cn</u>)

12

Abstract: Secondary organic aerosol (SOA) is an important constituent of airborne fine particles. 13  $PM_{2.5}$  (particles with aerodynamic diameters  $\leq 2.5 \ \mu m$ ) samples were collected at a mountainous 14 site in Hong Kong in autumn of 2010, and analyzed for SOA tracers. Results indicated that the 15 concentrations of isoprene SOA tracers (54.7 $\pm$ 22.7 ng/m<sup>3</sup>) and aromatics SOA tracers (2.1 $\pm$ 1.6 16  $ng/m^3$ ) were on relatively high levels in Hong Kong. Secondary organic carbon (SOC) derived 17 from isoprene, monoterpenes, sesquiterpenes and aromatics was estimated with the SOA tracer 18 based approach, which constituted  $0.35\pm0.15 \ \mu\text{g/m}^3$  (40.6±5.7%),  $0.20\pm0.03 \ \mu\text{g/m}^3$  (30.4±5.5%), 19  $0.05\pm0.02 \ \mu g/m^3$  (5.6±1.7%) and  $0.26\pm0.20 \ \mu g/m^3$  (21.3±8.2%) of the total estimated SOC. 20 Biogenic SOC ( $0.60\pm0.18 \text{ }\mu\text{g/m}^3$ ) dominated over anthropogenic SOC ( $0.26\pm0.20 \text{ }\mu\text{g/m}^3$ ) at this 21 site. In addition to the total estimated SOC ( $17.8\pm4.6\%$  of organic carbon (OC) in PM<sub>2.5</sub>), 22 primary organic carbon (POC) emitted from biomass burning also accounted for a considerable 23 24 proportion of OC (11.6±3.2%). Insight into the OC origins found that regional transport significantly (p < 0.05) elevated SOC from 0.37±0.17 to 1.04±0.39 µg/m<sup>3</sup>. Besides, SOC load 25 26 could also increase significantly if there was influence from local ship emission. Biomass burning related POC in regional air masses ( $0.81\pm0.24 \text{ }\mu\text{g/m}^3$ ) was also higher (p<0.05) than that 27 in samples affected by local air  $(0.29\pm0.35 \ \mu g/m^3)$ . Evidences indicated that SOA formation was 28 closely related to new particle formation and the growth of nucleation mode particles, while 29 30 biomass burning was responsible for some particle burst events in Hong Kong. This is the first SOA study in afforested areas of Hong Kong. 31

Keywords: Secondary organic aerosol, SOA tracer, biogenic SOA, regional transport, particlegrowth

## 34 1. Introduction

Atmospheric aerosol has been well recognized to affect global climate change (Stocker et al, 2013), human health (Dockery et al., 1993; Pope III and Dockery, 2006), visibility (Appel et al., 1985) and sustainability of economy (Chameides et al., 1999). Secondary organic aerosol (SOA) has been identified to play critical roles in these effects (Maria et al., 2004; Volkamer et al., 2006; Baltensperger et al., 2008), thus receiving sufficient attentions in recent years.

So far, the scientific community has reached a consensus that volatile organic compounds (VOCs) 40 from biogenic emissions and anthropogenic aromatics are key precursors of SOA (Forstner et al., 41 1997; Claeys, et al., 2004). In global scale, biogenic SOA is thought to be the greatest constituent 42 43 of SOA, due to the worldwide largest emission of biogenic VOCs (e.g., isoprene, monoterpenes and sesquiterpenes) and formation of biogenic SOA spanning a wide range of conditions (the 44 45 level of NO<sub>x</sub>, humidity and aerosol acidity) (Kroll et al., 2006). However, anthropogenic SOA has also been found to be significant in urban areas (Volkamer et al., 2006). Furthermore, upon 46 47 the findings that biogenic SOA correlates well with the indicators of anthropogenic emissions (Goldstein et al., 2009; Hoyle et al., 2011), it is believed that man-made air pollutants promote 48 49 the formation of biogenic SOA, in addition to serving as SOA precursors themselves. These promoting effects at least include forming aerosol seeds, catalyzing photooxidation and 50 51 transformation of biogenic VOCs and their oxidation products and changing the reaction pathways (e.g., atmospheric fate of isoprene in low- and high-NO<sub>x</sub> environments) (Hoyle et al., 52 53 2011). It is believed that SOA is a collection of hundreds to thousands of organic chemicals featuring relatively low volatilities. To understand SOA formation and explore the potential 54 55 sources, SOA speciation is of great necessity. However, due to the difficulty in chemical analysis of SOA tracers, the chemical compositions of SOA are far from being well understood. The 56 traditional analysis by gas chromatography-mass spectrometer detector (GC-MSD) generally 57 quantifies a total of less than 20 organic compounds in particles (Offenberg et al., 2007; 58 Kleindienst et al., 2007). Although some advanced instruments have been developed nowadays, 59 60 such as aerosol mass spectrometry and thermal desorption aerosol gas chromatography, they are either fragment-based or highly dependent upon the skills and knowledge of users (Williams et 61

al., 2014). Instead, SOA tracer based approach is a simplified method widely used to estimate the
amount, precursors and sources of SOA (Kleindienst et al., 2007; Ding et al., 2012).

The SOA tracer based approach applies the laboratory obtained ratios between the sum of 64 specific SOA tracers and total mass of SOA (or secondary organic carbon (SOC)) produced from 65 individual (group of) species to the field measured SOA tracers (Kleindienst et al., 2007), 66 roughly estimating SOA (SOC) derived from an individual VOC or VOC group. Table S1 67 summarizes the VOC precursors, corresponding SOA tracers and the ratios between SOA tracers 68 and SOA (or SOC), as reported by Kleindienst et al. (2007). The drawbacks of this method are 69 obvious. For instance, it is controversial whether the laboratory obtained ratios can be directly 70 applied to the field measured SOA tracers. However, it provides a feasible approach to estimate 71 the SOA concentration, which is especially helpful in the cases of not well knowing the SOA 72 compositions. More importantly, Offenberg et al. (2007) confirmed that the SOA tracer based 73 approach was reliable through comparing with the results obtained from <sup>14</sup>C contents. 74

Hong Kong, one of the most developed regions in East Asia, has a total territory area of 75 ~ $1.1 \times 10^3$  km<sup>2</sup> and a total population of ~7 million. Despite high population density, it keeps 24 76 77 country parks and a vegetation coverage rate of 70%. Evergreen broadleaf trees are common in Hong Kong. Guenther et al. (2006) suggested that shrubs are large emitters of isoprene, a typical 78 and most abundant biogenic VOC. The total emission amount of biogenic VOCs in Hong Kong 79 is estimated as  $8.6 \times 10^3$  ton C/a (Tsui et al., 2009). Field measurements also revealed that 80 81 isoprene in Hong Kong is relatively high (300-400 pptv) (Guo et al., 2007, 2012a). In contrast, aromatics are largely emitted from vehicular exhaust and solvent usage in Hong Kong. For 82 83 example, toluene is one of the most abundant aromatics, with the level of 3-6 ppby (Guo et al., 2004; Ho et al., 2004). Therefore, local emission of VOCs has a great potential forming SOA in 84 85 Hong Kong. In addition to local SOA formation, regional transport is inevitable due to severe air pollution in upwind direction of Hong Kong (i.e., inland Pearl River Delta (PRD) region). 86 Studies confirmed that Hong Kong received SOA produced by isoprene and toluene from inland 87 PRD region (Hu et al., 2008). However, the previous SOA studies in Hong Kong all focused on 88 urban areas, which are not enough to understand the abundance, compositions and sources of 89 90 SOA in low-altitude mountainous area where both anthropogenic and biogenic emissions are important. 91

92 In this study, SOC derived from isoprene, monoterpenes, sesquiterpenes and aromatics at a 93 mountainous site in Hong Kong were estimated using a SOA tracer based approach. Local and 94 regional contributions to SOC and biomass burning related POC were determined. Furthermore, 95 in combination with particle size distribution simultaneously monitored by a scanning mobility 96 particle sizer (SMPS), the relationships between SOA formation and particle growth were 97 examined.

98 **2. Methodology** 

#### 99 **2.1 Sample collection**

Hong Kong, a coastal city surrounded by the South China Sea (SCS) to the east and south, is 100 located in southern China. Under the dominance of subtropical monsoon climate, Hong Kong 101 receives northerly winds originating from the heavily polluted PRD region in cool seasons 102 (October-March), while prevailing southerly winds bring in clean air from SCS in warm seasons 103 (April-August) (So and Wang, 2003). The sampling site (22.405° N, 114.118° E, 640 m a.s.l) was 104 set up at the mountainside of the highest mountain in Hong Kong (Mt. Tai Mo Shan with the 105 maximum altitude of 957 m, TMS). The vegetation on this mountain mainly includes Acacia 106 107 confusa, Lophotemon confertus, Machilus chekiangensis and Schima superba below 550 m, while it turns to shrubs and grasses above 550 m (Guo et al., 2012b). As demonstrated in 108 109 previous studies (Guo et al., 2013; Ling et al., 2014), regional transport of air pollutants from inland PRD, mesoscale circulation (mountain-valley breezes) and in situ atmospheric chemistry 110 111 are the main processes that significantly influence air quality at this site. Figure 1 shows the geographical locations of the sampling site and air quality monitoring stations (AQMSs) of Hong 112 113 Kong Environmental Protection Department (HKEPD).

From September 7 to November 26, 2010, a total of 19  $PM_{2.5}$  samples were collected. The 114 115 instrument was an Anderson high volume PM2.5 sampler, with a flow rate of 750 L/min. Prebaked A4 size quartz fiber filters were used to collect the samples, which were stored in the 116 refrigerator at -18 °C after sampling. Generally, each filter sampling lasted for 45-55 hours, 117 118 except for the cases that the instrument stopped abnormally on some days due to thunderstormcaused power outages. Table S2 lists the sample IDs, start and end dates & times. The origins of 119 120 air masses, as distinguished by wind field and ratio of  $SO_2/NO_x$  (see section 3.3 for details), are also shown in the table. Additionally, the particle number concentrations and size distributions in 121 the size range of 5.5-350.4 nm in 44 size bins were monitored by a scanning mobility particle 122

sizer (SMPS) from October 27 to November 29. Detailed introductions about the operation
procedures of SMPS and data processing can be found in Guo et al. (2012a). We also collected
ambient VOC samples during September 28-November 21. Inorganic trace gases (SO<sub>2</sub>, CO, NO,
NO<sub>2</sub> and O<sub>3</sub>) and weather conditions were measured simultaneously with the PM<sub>2.5</sub> sampling.
Details about VOC sampling, VOC analysis and monitoring of trace gases are provided in Guo et
al. (2013) and Ling et al. (2014).



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Figure 1 Geographic locations of the sampling site (TMS), AQMSs of HKEPD and the container
terminal area where nine container terminals are located. Capital letters in the brackets are
abbreviations of the site/stations

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### 134 **2.2 Chemical analysis**

OC and element carbon (EC) in PM<sub>2.5</sub> samples were analyzed using the thermo-optical transmittance method recommended by National Institute for Occupational Safety and Health (NIOSH) (Birch, 1998). The concentrations of OC and EC are shown in Figure S1.

The method of SOA analysis was in line with that introduced by Ding et al. (2011). Briefly, the procedures for each sample include solvent extraction, derivation, analysis by GC-MSD, and identification and quantification of SOA tracers. 1/8 of each filter was extracted three times by sonication in the solvent of 40 mL of 1:1 (v/v) dichloride methane (DCM)/methanol mixture. Prior to extraction, the internal standards (hexadecanoic acid-D<sub>31</sub>, phthalic acid-D<sub>4</sub> and

levoglucosan- ${}^{13}C_6$ ) were spiked into the samples. The three-time extracts of each sample were 143 combined, filtered and concentrated to ~2 mL, which was further divided into two parts for 144 methylation and silvlation derivation, respectively. In methylation derivation, the extract 145 experienced a gentle nitrogen blow to dryness, and subsequent addition of 200 µL of DCM, 10 146  $\mu$ L of methanol and 300  $\mu$ L of freshly prepared diazomethane. Then, it was kept in room 147 temperature for one hour to derivatize acids to methyl esters, after which the sample was blown 148 to 200  $\mu$ L and used for analysis of some  $\alpha$ -pinene SOA tracers (Pinonic acid, pinic acid and 3-149 methyl-1,2,3-butanetricarboxylic acid). The silvlation reagent was 100 µL of pyridine and 200 150 uL of N.O-bis-(trimethylsilyl)-trifluoroacetamide (BSTFA) plus 1% trimethylchlorosilane 151 (TMCS). Differently, the derivation was carried out in an oven at 70  $^{\circ}$ C for one hour. One  $\alpha$ -152 pinene SOA tracer (3-hydroxyglutanic acid) and tracers for isoprene SOA, sesquiterpenes SOA, 153 and toluene SOA were analyzed from the silvlated sample. An Agilent 5973N GC/MSD was 154 employed to do the analysis. The identification of SOA tracers was based on the comparison of 155 mass spectra with previous studies, and their retention time in GC chromatogram with other 156 known compounds as the references. Pinonic acid and pinic acid were quantified by authentic 157 158 standards. However, due to lack of standards, other  $\alpha$ -pinene SOA tracers, isoprene SOA tracers,  $\beta$ -Caryophyllenic acid and 2,3-dihydroxy-4-oxopentanoic acid were quantified using pinic acid 159 (PA), erythritol, octadecanoic acid and azelaic acid, respectively. The detection limits (DLs) for 160 pinonic acid, pinic acid, erythritol, octadecanoic acid and azelaic acid were 0.05, 0.07, 0.06, 0.09, 161 and 0.11 ng/m<sup>3</sup>, respectively. Levoglucosan was also quantified with the DL of 0.15 ng/m<sup>3</sup>. The 162 SOA tracers analyzed in this study are highlighted in Table S1. 163

# 164 **2.3 Quality assurance and quality control**

In this study, internal standards were not spiked on the filters before sampling, to avoid their 165 166 influences on OC analysis. According to the saturation concentrations of SOA tracers calculated by Ding et al. (2016), SOA tracers analyzed in this study were of low volatilities, except for 167 pinonic acid. Therefore, pinonic acid in airborne PM<sub>2.5</sub> might be underestimated due to the blow-168 off effect during sampling, which however should not be significant to other SOA tracers. 169 170 Recovery target compounds were spiked in the samples before analysis of SOA tracers. The recovery rates were 104±2%, 68±13%, 62±14%, 78±10%, 81±9% and 87±4% for pinonic acid, 171 pinic acid, erythritol, octadecanoic acid, azelaic acid and levoglucosan, respectively. Since 172

internal standards were added into the samples prior to analysis, we did not use the recoveryrates to correct the concentrations of SOA tracers.

## 175 2.4 Processing of SMPS data

- 176 In this study, particles measured by SMPS were divided into nucleation (5.5-24.7 nm), Aitken
- 177 (24.7-101.4 nm) and accumulation modes (101.4-350.4 nm). Geometric mean diameter (GMD)
- 178 for nucleation mode particles (5.5-24.7 nm) was calculated using the following equations (Guo et
- 179 al., 2012a).

180 GRs = 
$$\frac{a_{GMD}}{d_t}$$
 (Equation (1))

- 181 GMD =  $e^{(\sum n_i ln^{d_i})/N}$  (Equation (2))
- 182  $\frac{d_N}{d_{\log} D_p} = \frac{\Delta N}{\log_{D_p^2} \log_{D_p^1}}$  (Equation (3))
- 183 where  $n_i$  is the particle number concentration in the *i*<sup>th</sup> bin with upper diameter of  $d_i$ , *N* represents 184 the total number concentration (cm<sup>-3</sup>).  $\Delta N$  is the particle number concentration in the size bin 185 with upper and lower limit diameter of  $D_p^2$  and  $D_p^1$ , respectively.
- 186 Condensation sink (CS), which describes the loss rate of vapor molecules and newly formed
- 187 particles on the pre-existing aerosol particles, was calculated as follows (Kulmala et al., 2005):

188 CS =  $2\pi D \int D_p \beta_m(D_p) n(D_p) dD_p = 2\pi D \sum_i \beta_{mi} D_{pi} N_i$  (Equation (4))

- where D is the diffusion coefficient of the condensing vapor,  $\beta_m$  is the transitional regime correction factor,  $D_p$  denotes the particle diameter, *n* and *N* represent the particle numbers.  $\beta_{mi}$ ,  $D_{pi}$  and  $N_i$  are the specific values for a given size bin (*i*). Growth factor calculated according to Laakso et al. (2004) was used to calibrate the dry particle size measured by SMPS.
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# 194 **3. Results and discussion**

# **3.1 Concentrations of SOA tracers**

Table 1 shows the average concentrations of SOA tracers derived from different precursors at TMS. Isoprene SOA tracers were the most abundant  $(54.7\pm22.7 \text{ ng/m}^3)$ , followed by the tracers generated from monoterpenes  $(26.3\pm4.5 \text{ ng/m}^3)$ , aromatics  $(2.1\pm1.6 \text{ ng/m}^3)$  and sesquiterpenes  $(1.1\pm0.4 \text{ ng/m}^3)$ . Figure 2 compares the concentrations of SOA tracers between previous studies and the present study. The number and species of SOA tracers for the same precursor were the same. For the cases that total concentration of monoterpenes SOA tracers was given (*e.g.*, Offenberg et al., 2011), a factor was applied to the total concentration to roughly estimate the 203 sum of monoterpenes SOA tracers with the number and species identical to this study (see section S1 and Figure S2 in the Supplement). It was found that isoprene SOA tracers at TMS 204 were on relatively high level (54.7 $\pm$ 22.7 ng/m<sup>3</sup>), comparable to or even higher than those 205 detected in the forest (61.4±30.4 ng/m<sup>3</sup> in Kleindienst et al. (2007) and 39.0±17.2 ng/m<sup>3</sup> in 206 207 Offenberg et al. (2011)). In fact, isoprene at TMS was relatively low (2-517 pptv), the high isoprene SOA tracers indicated that biogenic SOA formation at this site might be enhanced by 208 209 anthropogenic emissions, e.g., sufficient aerosol seeds from SO<sub>2</sub>-related new particle formation (Guo et al., 2012a). Certainly, regional transport could also contribute to high isoprene SOA 210 tracers, as reported by Hu et al. (2008). For the anthropogenic SOA, the tracer produced by 211 aromatics (2,3-dihydroxy-4-oxopentanoic acid) was noticeable. Although the average 2,3-212 dihydroxy-4-oxopentanoic acid in this study  $(2.1\pm1.6 \text{ ng/m}^3)$  was not significantly higher than 213 that in other studies (p>0.05), its maximum value reached 13.5 ng/m<sup>3</sup>. Given abundant aromatics 214 in the atmosphere of Hong Kong and mesoscale circulations between TMS and the surrounding 215 urban areas, local formation could be an important factor contributing to high aromatics SOA at 216 Moreover, it was reported that 2,3-dihydroxy-4-oxopentanoic acid at an urban 217 TMS. background site in inland PRD was very high (13.1 ng/m<sup>3</sup>) (Ding et al., 2012). Hence, regional 218 transport might be partially responsible for the relatively high aromatics SOA tracer in Hong 219 Kong. SOA tracers derived from monoterpenes and sesquiterpenes were on moderate levels, 220 compared to other studies. 221

222 3-Methyl-1,2,3-butanetricarboxylic acid and 3-hydroxyglutanic acid, formed from monoterpenes oxidation in the presence of NO<sub>x</sub> (Claevs et al., 2007; Eddingsaas et al., 2012), dominated the 223 224 measured monoterpenes SOA tracers, consistent with those at an urban background site in inland PRD (Ding et al., 2012). Pinonic acid and pinic acid, formed from OH oxidation of  $\alpha$ -pinene in 225 226 NO<sub>x</sub> free environment or ozonolysis of  $\alpha$ -pinene (Eddingsaas et al., 2012), were also higher than those in urban areas of Hong Kong (almost below DLs) (Hu et al., 2008). Since NO<sub>x</sub> was at the 227 magnitude of several ppby, the higher pinonic acid and pinic acid were likely due to the higher 228  $O_3$  (69.2±2.4 ppbv at TMS and 30.8±2.6 ppbv at the mountain foot) at this mountainous site. In 229 230 addition, other tracers including 3-hydroxy-4,4-dimethylglutaric acid, 3-isopropylpentanedioic 231 acid, 3-acetylpentanedioic acid and 3-acetylhexanedioic acid, which were not measured in this study, accounted for 46.9±4.0% of the total amount of tracers according to Kleindienst et al. 232 (2007). Of the isoprene SOA tracers, 2-methylthreitol and 2-methylerythritol are representative 233

products formed through the photooxidation of isoprene hydroxyhydroperoxides and acid catalysis of epoxydiols of isoprene in low NO<sub>x</sub> environment when RO<sub>2</sub> reacting with HO<sub>2</sub> dominated the loss of RO<sub>2</sub> (Claeys et al., 2004; Surratt et al., 2010). In this study, they accounted for  $83.5\pm8.1\%$  of the total isoprene tracers. However, NO<sub>x</sub> was not low enough and RO<sub>2</sub> reacting with NO was the main sink of RO<sub>2</sub> (Ling et al., 2014), implying that other mechanisms might enhance the formation of 2-methyltetrols (sum of 2-methylthreitol and 2-methylerythritol) at this site, such as acid catalysis (Surratt et al., 2007).

Table 1 Average concentrations of SOA tracers derived from monoterpenes, isoprene,
sesquiterpenes and aromatics (mean±95% confidence interval (C.I.)).

VOC precursor	SOA tracer	Concentration (ng/m <sup>3</sup> )
Monoterpenes	Pinonic acid	1.2±0.4
	Pinic acid	2.0±0.6
	3-methyl-1,2,3-butanetricarboxylic acid	16.0±3.5
	3-hydroxyglutanic acid	7.1±2.7
	Sum 1	26.3±4.5
Isoprene	2-Methylthreitol	14.7±6.6
	2-Methylerythritol	31.2±13.1
	2-methylglyceric acid	5.2±3.1
	cis-2-Methyl-1,3,4-trihydroxy-1-butene	$0.8 \pm 0.5$
	trans-2-Methyl-1,3,4-trihydroxy-1-	2.0±1.3
	butene	
	3-Methyl-2,3,4-trihydroxy-1-butene	0.7±0.3
	Sum 2	54.7±22.7
Sesquiterpenes	$\beta$ -Caryophyllenic acid	1.1±0.4
Aromatics	2,3-Dihydroxy-4-oxopentanoic acid	2.1±1.6

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Figure 2 Comparison of SOA tracers between previous studies and this study. <sup>1</sup>Kleindienst et al. (2007); <sup>2</sup>Offenberg et al. (2011); <sup>3</sup>Ding et al. (2012); <sup>4</sup>Hu et al. (2008); <sup>5</sup>Lewandowski et al. (2008); <sup>6</sup>Haddad et al. (2011); <sup>7</sup>this study.

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### 249 **3.2 Estimate of SOC**

Both the EC tracer method and SOA tracer method were used to estimate SOC (details about 250 251 both methods are provided in section S2 of the Supplement). To apply the SOA tracer method, 252 the sum of monoterpenes SOA tracers was scaled by a factor of 1.79 (see section S1 and Figure S2 for details). Figure S3 compares the SOC estimated by these two methods. SOC estimated by 253 EC tracer method (SOC<sub>EC tracer</sub>) were generally higher (p < 0.01) than those estimated by SOA 254 tracer method (SOC<sub>SOA tracer</sub>), except for samples TMS1, TMS3 and TMS8. Ding et al. (2012) 255 indicated that the EC tracer method might overestimate SOC, because it blended some primary 256 OC (POC) from biomass burning with SOC. This inference was supported by the good 257 correlation between the difference of SOC estimated by these two methods (SOC<sub>EC tracer</sub>-SOC<sub>SOA</sub> 258 tracer) and levoglucosan, a tracer of biomass burning. 259

In addition to levoglucosan, methyl chloride (CH<sub>3</sub>Cl) is an indicator of biomass burning 260 261 (Rudolph et al., 1995). Figure 3 plots the correlation between levoglucosan and CH<sub>3</sub>Cl at TMS, as well as that between SOC<sub>EC tracer</sub>-SOC<sub>SOA tracer</sub> and levoglucosan. As expected, levoglucosan 262 fairly correlated with CH<sub>3</sub>Cl ( $R^2$ =0.54), further confirming the reliability of levoglucosan as the 263 tracer of biomass burning. Consistent with Ding et al. (2012), good correlation was found 264 between  $SOC_{EC \text{ tracer}}$ -SOC<sub>SOA tracer</sub> and levoglucosan (R<sup>2</sup>=0.52). This verified that the EC tracer 265 method overestimated SOC due to the interference of biomass burning. Exceptionally, SOC<sub>EC</sub> 266 267 tracer was remarkably lower than SOC<sub>SOA tracer</sub> for the samples TMS1, TMS3 and TMS8 (p<0.05). 268 Further investigation found that on the sampling days of TMS1 and TMS3, the daily maximum hourly  $O_3$  mixing ratio, referred to as peak  $O_3$ , was extremely high, with the values being the 269

270 second (154.4 ppbv) and first (163.4 ppbv) highest among the 19 filter samples, respectively, while during the sampling period of TMS8, the peak O<sub>3</sub> also reached 94.8 ppby. In contrast, the 271 272 average of peak O<sub>3</sub> values on the sampling days of other PM<sub>2.5</sub> filter samples was only 77.6 ppbv. Since high O<sub>3</sub> levels generally imply strong photochemical reactivity, high productions of SOA 273 are also expected on these days. Hence, the higher SOC<sub>SOA tracer</sub> values observed in TMS1, TMS3 274 and TMS8 were understandable. We also found that there was no correlation between SOC<sub>EC</sub> 275  $_{tracer}$  and peak O<sub>3</sub> (not shown), whereas SOC<sub>SOA tracer</sub> correlated well with peak O<sub>3</sub> (R<sup>2</sup>=0.68), as 276 shown in Figure 4, implying that the formation of secondary products, *i.e.*, SOA and O<sub>3</sub>, depends 277 upon the oxidative capacity of the atmosphere and they may also influence each other. Therefore, 278 the SOA tracer method was believed to be more reliable and thus adopted in this study. Hereafter, 279 SOC refers to SOC<sub>SOA tracer</sub>, unless otherwise specified. 280





283 <sub>tracer</sub> and levoglucosan.



Figure 4 Linear correlation between SOC<sub>SOA tracer</sub> and peak O<sub>3</sub>.

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287 Figure 5 shows SOC produced by different groups of VOCs estimated using the SOA tracer based approach, and biomass burning related POC (POC biomass burning). POC biomass burning was 288 estimated by 0.01 ( $\mu$ g/ng) × levoglucosan (ng/m<sup>3</sup>) (Lee et al., 2008). On average, SOC and POC 289 biomass burning constituted 0.86±0.31  $\mu$ g/m<sup>3</sup> (17.8±4.6%) and 0.67±0.22  $\mu$ g/m<sup>3</sup> (11.6±3.2%) of OC 290 291 (provided in Figure S1), respectively. The rest of OC were undetermined due to unknown sources and precursors of OC. The total SOC comprised anthropogenic (*i.e.*, aromatics) and 292 293 biogenic (*i.e.*, isoprene, monoterpenes and sesquiterpenes) SOC, with the fractions of 21.3±8.2%  $(0.26\pm0.20 \ \mu\text{g/m}^3)$  and  $78.7\pm8.2\%$   $(0.60\pm0.18 \ \mu\text{g/m}^3)$ , respectively. Although anthropogenic 294 SOC (ASOC) was significantly lower than biogenic SOC (BSOC) (p<0.05), ASOC had its 295 highest value of 1.71  $\mu$ g/m<sup>3</sup> in the sample TMS1, and BSOC reached its maximum in sample 296 297 TMS3 (2.03  $\mu$ g/m<sup>3</sup>). As mentioned earlier, the two samples were collected on the days with very high O<sub>3</sub>, indicating that aromatics and biogenic VOCs might be responsible for the high SOC in 298 TMS1 and TMS3, respectively. However, VOC samples were not simultaneously collected 299 during the collection of these two samples. Instead, the sample TMS12 was a good example 300 because ASOC was the second highest  $(1.2 \,\mu\text{g/m}^3)$ , and the daily average mixing ratio of toluene 301 coincidently reached 4.0 ppbv during the TMS12 sampling period, the highest value among all 302 VOC samples, which further confirmed the reliability of SOA tracer method in estimating SOC. 303 Among SOC derived from biogenic VOCs, isoprene made the highest contribution (54.2±5.3% 304 of BSOC and 42.7±5.9% of total SOC), followed by monoterpenes (37.9±4.6% and 30.4±5.5%, 305 306 respectively) and sesquiterpenes  $(7.9\pm2.7\% \text{ and } 5.6\pm1.7\%, \text{ respectively})$ .



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Figure 5 Concentrations of estimated SOC derived from monoterpenes, isoprene, sesquiterpenes
 and aromatics and POC <sub>biomass burning</sub>.

# 310 **3.3 Local and regional contributions to SOC**

311 In line with the method used in Guo et al. (2013), the local and regional air masses were distinguished with the wind direction (WD) and wind speed (WS) monitored at TMS. Briefly, 312 the northerly winds  $(0^{\circ} < WD < 90^{\circ} \text{ or } 270^{\circ} < WD < 360^{\circ})$  with WS higher than 2 m/s were considered 313 to be capable of delivering air pollutants from the heavily polluted inland PRD region to the site. 314 In these cases, the air masses were believed to be subject to regional influences. Otherwise, the 315 site was dominated by local air. To validate this method, Figure 6 shows the hourly ratios of 316  $SO_2/NO_x$  for the air masses in different wind directions/speeds. Overall, the  $SO_2/NO_x$  ratios were 317 higher for regional air masses, particularly when 0°<WD<90° and WS>2 m/s. This coincided 318 with the lower sulfur content in vehicle fuels and higher vehicle emissions of NO<sub>x</sub> in Hong Kong 319 (Wang and So, 2003; Guo et al., 2009). For the 24-48 hr PM<sub>2.5</sub> samples (*i.e.*, duration of 1-2 320 days), the regional influence was not unexpected given that the northerly wind was stronger than 321 2 m/s for at least one hour on each sampling day. It was noteworthy that during the sampling 322 periods of TMS1 and TMS16 (influenced by regional air), SO<sub>2</sub> was exclusively high at the 323 HKEPD AQMSs in vicinity of the ship container terminals, e.g., KC, SSP and TW (see Figure 1). 324 Figure S4 presents SO<sub>2</sub> distributions at 14 AQMSs in Hong Kong during these two periods. In 325 view of the fact that ship is a significant emitter of SO<sub>2</sub>, the influence of local ship emission was 326 also suspected for the two samples. According to these inferences, Table S2 lists the categories 327 328 of the samples affected by regional air, local air, and local ship emissions, respectively.



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Figure 6 Hourly ratios of  $SO_2/NO_x$  for the air masses in different wind directions/speeds at TMS. Table 2 summarizes the concentrations of SOC and POC <sub>biomass burning</sub> in different categories of air masses. Note that since the samples TMS1 and TMS16 were influenced by both regional air and local ship emission, they were separately discussed later. It was found that SOC derived from

334 individual group/species and total SOC in regional air were all significantly (p < 0.05) higher than that in local air, suggesting that SOC at TMS was elevated by regional transport. Despite 335 336 possible influences of local ship emissions in TMS1 and TMS16, the concentrations of SOC and POC in these two samples were well within the ranges of those in regional air, except for 337 aromatics SOC (1.71  $\mu$ g/m<sup>3</sup>) in sample TMS1. The extremely high aromatics SOC in TMS1 338 might be caused by ship emissions which could be laden with high concentrations of aromatics. 339 Contradictorily, SOC derived from aromatics was remarkably lower in TMS16 (0.27  $\mu$ g/m<sup>3</sup>). 340 This discrepancy might be explained by the differences of fuel types and operating conditions of 341 the ship engines, which were repeatedly proved to affect the emission characteristics of ship 342 engines (Reda et al., 2014; Sippula et al., 2014). Similar to SOC, POC also showed a 343 significantly higher level in regional air  $(0.81\pm0.24 \text{ }\mu\text{g/m}^3)$  than that in local air  $(0.29\pm0.35 \text{ }\mu\text{g/m}^3)$ 344  $\mu g/m^3$ ) (p<0.05). Since the high concentration of POC <sub>biomass burning</sub> in regional air was partially 345 contributed by TMS16 (POC <sub>biomass burning</sub> =1.54  $\mu$ g/m<sup>3</sup>), a sample jointly influenced by regional 346 air and local ship emission, specific insight was given to this sample. Firstly, ship emission was 347 not likely to be the culprit of high POC biomass burning, as no study reported ship emission of 348 levoglucosan, the biomass burning tracer used to calculate POC biomass burning in this study. Instead, 349 we found that another sample under the influence of regional air (TMS15) had comparable POC 350 biomass burning (1.64 µg/m<sup>3</sup>). Furthermore, CH<sub>3</sub>Cl, another tracer of biomass burning, increased 351 noticeably under northerly winds in sample TMS16, indicating the regional transport of biomass 352 353 burning plumes into Hong Kong. In fact, nearly no fire spot in local Hong Kong was observed by the satellite during the sampling, compared to some open fires detected in upwind directions (see 354 Figure S5). Therefore, POC biomass burning in TMS16 was reasonably speculated to be elevated by 355 regional biomass burning plumes. Upon this inference, we concluded that regional transport 356 357 significantly contributed to PM<sub>2.5</sub>-bounded POC in Hong Kong.

358

Table 2 Mean $\pm$ 95% C.I. of SOC and POC in different categories of air masses (Unit:  $\mu$ g/m<sup>3</sup>).

	Local air	Regional air	TMS1 <sup>*</sup>	TMS16 <sup>*</sup>
SOC (monoterpenes)	0.13±0.06	0.23±0.03	0.23	0.28
SOC (isoprene)	$0.16 \pm 0.07$	$0.42 \pm 0.18$	0.51	0.41
SOC (sesquiterpenes)	$0.01 \pm 0.01$	$0.06 \pm 0.02$	0.16	0.07
SOC (aromatics)	$0.07 \pm 0.04$	0.33±0.26	1.71	0.27

POC biomass burning	$0.29 \pm 0.35$	0.81±0.24	0.17	1.54
Total SOC	0.37±0.17	1.04±0.39	2.61	1.04

<sup>\*</sup> 95% C.I. is not available for a single sample.

## **361 3.4 Implications to particle formation and growth**

As an important constituent of airborne particles, SOA plays critical role in particle formation 362 and growth (Jang et al., 2003). The relationships between SOA formation and particle 363 364 formation/growth were investigated on three selected days (October 31, November 09 and November 19) when SOC were among the highest of all the samples and SMPS data were 365 available. Figures 7-9 show the evolutions of particle numbers, GMD of nucleation mode 366 particles, CS, the simulated sulfuric acid (SA) vapor, oxidized intermediates of aromatics (AOI), 367 oxidized intermediates of biogenic VOCs (BOI), and the measured inorganic trace gases. Details 368 about the modelling of SA vapor, AOI and BOI are provided in Section S3 and Table S3 in the 369 Supplement. For convenience of analysis, the hourly mixing ratios of biogenic VOCs (BVOCs), 370 aromatics and CH<sub>3</sub>Cl are presented in Figure S6. 371

In Figures 7-8, the number concentration of nucleation mode particles (N<sub>nuc</sub>) increased 372 substantially in the morning of October 31 (11:00-12:00) and November 09 (10:00-11:00), 373 followed by the increases of number of Aitken mode particles (NAit). Prior to the increases of 374 N<sub>nuc</sub>, the simulated SA vapor began to increase about 3 hours earlier. These were consistent with 375 the findings in Guo et al. (2012a) who reported that the increase of  $N_{nuc}$  was caused by new 376 particle formation (NPF) events occurred on both days, and SA vapor played important roles in 377 NPF. However, we further found that the oxidation intermediates of BVOCs (*i.e.*, BOI) also 378 increased slightly ahead of the rise of N<sub>nuc</sub>, which might suggest that the oxidation of BVOCs 379 also made some contributions to NPF. In fact, the involvement of BVOCs in NPF at this 380 381 afforested site has been speculated by Guo et al. (2012a), which is confirmed with the aid of model simulations in this study. 382

Moreover, we noticed that the nucleation mode particles experienced obvious growth with the rate of 1.9 nm/h (15:00-16:00) and 1.4 nm/h (14:00-16:00) in the afternoon of October 31 and November 09, respectively. Both growths occurred under the conditions of high  $N_{Ait}$  and high CS, differing from NPF events. Meanwhile,  $O_3$  was on high level, which meant strong oxidative capacity of the atmosphere. Correspondingly, the simulated SA vapor, AOI and BOI showed great increments simultaneously with or 1-2 hours earlier than the increase of GMD. It is 389 noteworthy that the significant increases of AOI and BOI were also attributable to the rapid 390 increases of VOCs (see Figure S6), in addition to strong atmospheric oxidative capacity. This 391 suggested that the photo-oxidation of VOCs also facilitated the growth of nucleation mode particles. The prompt responses of particle growth to the increments of oxidation products on 392 October 31 (rather than 1.5 hours' delay on November 09) were likely caused by the much more 393 abundant BOI (~22.7 pptv) than that on November 09 (~13.5 pptv). Besides, the lower initial 394 GMD before its increase on October 31 (~14 nm compared to ~16 nm on November 09) implied 395 higher surface area and subsequently quicker growth. Since the aforementioned days featured 396 high SOA, the roles of photo-oxidation of VOCs in the formation and growth of nucleation mode 397 particles might reflect the very initial stages of SOA formation. However, to better understand 398 399 the relationships between SOA formation and the formation/growth of particles, data with higher resolution and more comprehensive chemical information of SOA are crucially needed. 400



401

Figure 7 Evolutions of particle numbers, GMD of nucleation mode particles, CS, simulated SA



vapor, AOI, BOI and inorganic trace gases on October 31 (first sampling day of TMS12).



404

Figure 8 Evolutions of particle numbers, GMD of nucleation mode particles, CS, simulated SA
vapor, AOI, BOI and inorganic trace gases on November 09 (second sampling day of TMS14).

Similarly, the growth of nucleation mode particles was also observed on November 19 (15:00-408 409 16:00) (Figure 9). However, a distinct phenomenon was that the numbers of particles in nucleation, Aitken and accumulation modes all showed rapid increases simultaneously from 410 411 around 14:30 and reached the highest values at ~15:00, which indicated that the particles in 412 different modes shared a common source. Although SO<sub>2</sub> and the simulated SA vapor began to 413 increase 1.5 hours earlier, this could not be a NPF event, as the increase of NAit had no delay (no banana shape) and the CS was high. In addition, the levels of primary air pollutants (e.g., NO<sub>x</sub> 414 415 and CO) were high at the peak hours of all three-mode particle numbers. More importantly, we found that CH<sub>3</sub>Cl largely increased from 13:00 to 15:00 and reached the maximum at 15:00 (see 416 417 Figure S6). Therefore, we suspected that biomass burning might be responsible for the increase of particle numbers. This coincided with the high POC biomass burning  $(1.54 \ \mu g/m^3)$  in PM<sub>2.5</sub> sample 418 collected on this day (TMS16). Generally, particles emitted from biomass burning are in Aitken 419 and accumulation modes (Reid et al., 2005). However, in vicinity of fire, nucleation mode can 420 also exist (Janhall et al., 2010). Here, although nucleation mode particles increased in the particle 421

burst event,  $N_{nuc}$  was very low (maximum= $2.1 \times 10^3$  cm<sup>-3</sup>), indicating that this was not a local biomass burning and nucleation mode particles converted to larger size particles from the source region to this site. This was also consistent with the regional influence on this day identified by the wind fields (see section 3.3).



426

Figure 9 Evolutions of particle numbers, GMD of nucleation mode particles, CS, simulated SA
vapor, AOI, BOI and inorganic trace gases on November 19 (sampling day of TMS16).

429

# 430 4. Conclusions

PM<sub>2.5</sub> samples were collected at a mountainous site in Hong Kong in autumn of 2010. Nine SOA 431 tracers in PM<sub>2.5</sub> were analyzed, which helped to understand the compositions and sources of 432 SOC in this study. Results indicated that isoprene made the highest contribution to SOC 433 formation at this site, followed by monoterpenes, aromatics and sesquiterpenes. Averagely, 434 biogenic SOC dominated over anthropogenic SOC. However, anthropogenic SOC cannot be 435 neglected, particularly under the influences of regional transport (e.g., 1.2  $\mu$ g/m<sup>3</sup> in sample 436 TMS12) and/or local ship emission (e.g.,  $1.7 \text{ }\mu\text{g/m}^3$  in sample TMS1). The simultaneous 437 observation of VOCs confirmed the role of aromatics in contributing to high concentrations of 438 439 anthropogenic SOC. In terms of SOC origins, regional transport caused nearly two-fold increase

440 of SOC, relative to local air. However, SOC load could also be significantly elevated by local ship emissions possibly containing abundant VOC precursors and SO<sub>2</sub>, which promoted SOC 441 442 formation. In addition, the regional air was generally characterized with high biomass burning 443 related POC, aggravating the PM<sub>2.5</sub>-bounded POC in Hong Kong. In combination with the SMPS data, we found that the formation of SOA (particularly the biogenic SOA) might be partially 444 responsible for the new particle formation and growth of nucleation mode particles. Primary 445 emissions, such as biomass burning, could cause particle burst events and lead to POC increases. 446 To our knowledge, this is the first SOA study carried out in low-altitude (640 m) mountainous 447 area of Hong Kong, where the air quality is under the combined influence of anthropogenic and 448 biogenic emissions. This study also demonstrates the urgency of data acquisition with more 449 comprehensive chemical information and higher time resolution in future SOA studies over this 450 region. 451

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