

1 **Formation and sink of glyoxal and methylglyoxal in a polluted subtropical**  
2 **environment: observation-based photochemical analysis and impact evaluation**

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15  
16 **Abstract**

17 The dicarbonyls, glyoxal (Gly) and methylglyoxal (Mgly) have been recognized as  
18 important precursors of secondary organic aerosols (SOAs) through the atmospheric  
19 heterogeneous process. In this study, field measurement was conducted at a receptor  
20 site in the Pearl River Delta (PRD) region in south China, and an observation based  
21 photochemical box model was subsequently applied to investigate the production and  
22 evolution of Gly and Mgly as well as their contributions to SOA formation. The model  
23 was coupled with a detailed gas-phase oxidation mechanism of volatile organic  
24 compounds (VOCs) (*i.e.*, MCM v3.2), heterogeneous processes of Gly and Mgly (*i.e.*,  
25 reversible partitioning in aqueous phase, irreversible volume reactions and irreversible  
26 surface uptake processes), and the gas-particle partitioning of oxidation products. The  
27 results suggested that without considering the heterogeneous processes of Gly and  
28 Mgly on aerosol surfaces, the model would overpredict the mixing ratios of Gly and  
29 Mgly by factors of 3.3 and 3.5 compared to the observed levels. The agreement between  
30 observation and simulation improved significantly when the irreversible uptake and the  
31 reversible partitioning were incorporated into the model, which in total both contributed  
32 ~62% to the destruction of Gly and Mgly during daytime, respectively. Further analysis  
33 on the photochemical budget of Gly and Mgly showed that the oxidation of aromatics  
34 by the OH radical was the major pathway producing Gly and Mgly, followed by

35 degradation of alkynes and alkenes. Furthermore, based on the improved model  
36 mechanism, the contributions of VOCs oxidation to SOA formed from gas-particle  
37 partitioning (SOA<sub>gp</sub>) and from heterogeneous processes of Gly and Mgly (SOA<sub>het</sub>) were  
38 also quantified. It was found that *o*-xylene was the most significant contributor to  
39 SOA<sub>gp</sub> formation (~29%), while *m,p*-xylene and toluene made dominant contributions  
40 to SOA<sub>het</sub> formation. Overall, the heterogeneous processes of Gly and Mgly can explain  
41 ~21% of SOA mass in the PRD region. The results of this study demonstrated the  
42 important roles of heterogeneous processes of Gly and Mgly in SOA formation, and  
43 highlighted the need for a better understanding of the evolution of intermediate  
44 oxidation products.

45

46 **Keywords:** Glyoxal, Methylglyoxal, Secondary organic aerosol, Pearl River Delta,  
47 Volatile organic compound, Photochemical box-model

48

## 49 1. Introduction

50 Organic aerosols (OAs) are important components of atmospheric aerosols, with  
51 important impacts on radiation balance, air quality, atmospheric oxidative capacity, and  
52 climate change (Zhu et al., 2011; Carlton et al., 2009; Hoyle et al., 2009). In addition  
53 to the primary organic components (primary OA, POA) directly emitted from various  
54 sources in the particulate form, a large fraction of OAs are secondarily produced (SOA)  
55 through the aging of POAs, and through complex homogenous/heterogeneous reactions  
56 of volatile or semi-volatile organic compounds (VOCs, SVOCs) (Jimenez et al., 2009;  
57 Steinfeld and Jeffrey, 1998). SOA has frequently been observed to dominate the OA in  
58 many regions, particularly during severe haze pollution events (Guo et al., 2012; Zhang  
59 et al., 2017). However, the characteristics of SOAs are still poorly understood because  
60 of their complicated formation mechanisms, various chemical compositions, and  
61 multitude of precursors from diverse emissions, thus making SOAs an important  
62 research topic in the field of the atmospheric environment.

63 In addition to primary precursors including isoprene, terpene and aromatic  
64 hydrocarbons, glyoxal (Gly) and methylglyoxal (Mgly) have been recognized to be of  
65 critical importance to SOA formation, especially through heterogeneous and multiphase  
66 processes, in many laboratory and model studies (Waxman, et al., 2013, 2015; McNeill

67 et al., 2015; De Haan et al., 2009; Fu et al., 2008). Many efforts have been made to  
68 investigate the sources, evolution of Gly and Mgly and their contributions to SOA  
69 (Benavent et al., 2019; Zhang et al., 2016; Sumner et al., 2014; DiGangi et al., 2012;  
70 Stavarakou et al., 2009). For example, Li et al (2015) constructed a Master Chemical  
71 Mechanism with an equilibrium partitioning module and coupled it in a Community  
72 Air Quality Model (CMAQ) to predict the regional concentrations of SOA from VOCs  
73 in the eastern United States (U.S). It was found that those SOA formed from Gly and  
74 Mgly were accounted for more than 35% of total SOA. Similarly, Ying et al. (2015)  
75 used a modified SAPRC-11 (S11) photochemical mechanism, considering the surface-  
76 controlled reactive uptake of Gly and Mgly, and incorporated the mechanism in the  
77 CMAQ model to simulate ambient SOA concentrations during summer in the eastern  
78 U.S. The results showed that the uptake of Gly and Mgly resulted in the significant  
79 improvement in predicated SOA concentration, and the aerosol surface uptake of  
80 isoprene-generated Gly, Mgly and epoxydiol accounted for more than 45% of total SOA.

81 As two smallest dicarbonyl compounds, the sources of Gly and Mgly are  
82 complicated. It has been well documented that Gly and Mgly have limited primary  
83 sources except biomass burning and biofuel combustion (Grosjean et al., 2001; Zhang  
84 et al., 2016). Furthermore, the primary emissions of Gly and Mgly were much less  
85 significant than those secondarily from photochemical reactions (Lv et al., 2019). Fu et  
86 al. (2008) estimated that primary emissions only accounted for about 4% and 17% to  
87 the total emissions of Mgly and Gly, respectively. On a global scale, isoprene and  
88 ethyne are the most important precursors of Gly and Mgly; on the local scale, however,  
89 degradation of aromatics is the major pathway for the production of Gly and Mgly in  
90 urban and sub-urban areas. For example, the oxidation of aromatics contributed to  
91 approximately 75% of Gly formation in Mexico City (Li et al., 2014; Volkamer et al.,  
92 2007).

93 As for the atmospheric sink for Gly and Mgly, photolysis, reaction with OH, dry  
94 deposition, and heterogenous processes are considered as the main loss pathways,  
95 among which aerosol uptake is most complicated and needs more comprehensive  
96 exploration (De Haan et al., 2018; McNeill, 2015; Knote et al., 2014; Fu et al., 2008).  
97 The uptake of Gly and Mgly onto inorganic or organic particles has been studied in  
98 laboratory experiments under controlled conditions (De Haan et al., 2018; Liggio et al.,  
99 2005), and uptake coefficients ( $\gamma$ ) were measured by the loss of gas phase concentration

100 or the increase of particle organic mass, within the range of  $\sim 10^{-4}$  to  $10^{-2}$  (De Haan et  
101 al., 2018; Pye et al., 2017; Liggio et al., 2005). The lower  $\gamma$  value was probably related  
102 to the kinetic limitations (Ervens and Volkman 2010), while the higher  $\gamma$  value may be  
103 associated with the increased particle acidity (Liggio et al., 2005), relative humidity  
104 (De Haan et al., 2018; Corrigan et al., 2008) and ionic strength (Kroll et al., 2005). In  
105 addition, ammonium-catalyzed and OH reactions were found to have significant  
106 influences on the surface uptake of dicarbonyls (Knote et al., 2014; Kampf et al., 2013;  
107 Noziere et al., 2008), and the rate coefficients were found to increase with the increasing  
108 ammonium ion activity ( $a_{\text{NH}_4^+}$ ) and pH (Noziere et al., 2008). The “salting-in” effects  
109 resulted from the increased ionic strength could cause significant increase ( $\sim 3$  orders  
110 of magnitude) of Henry’s law constant for Gly, affecting the gas-aqueous partitioning of  
111 Gly and enhanced the available Gly for aqueous reactions (Kampf et al., 2013; Knote  
112 et al., 2014; Waxman et al., 2015).

113 The uptake processes of Gly and Mgly derived from the laboratory studies were  
114 incorporated into different models to investigate their formation and destruction (Ge et  
115 al., 2011; Knote et al., 2014; Pye et al., 2017). It was found that solely incorporating  
116 the irreversible uptake pathways of dicarbonyls could lead to high discrepancy between  
117 the observation and simulation results from the global 3D model and other models (Hu  
118 et al., 2017; Li X et al., 2014; Li et al., 2013a), highlighting the needs to consider more  
119 comprehensive processes including both reversible and irreversible pathways for better  
120 simulating the dicarbonyls. Those previous studies showed that the contribution of  
121 heterogeneous processes to the destruction of dicarbonyls varied in the range of 0~80%,  
122 which depended on the relative humidity, the precursors incorporated into the model as  
123 well as the aerosol concentrations for the given region (Knote et al., 2014).

124 The Pearl River Delta (PRD) region has been experiencing rapid industrialization  
125 and urbanization in the last three decades, making it one of the most developed regions  
126 in China. The field measurement results suggested that OA contributed 30~40% to  
127  $\text{PM}_{2.5}$  mass, and SOA dominated the OA with fractions up to  $\sim 80\%$  in PRD (Huang et  
128 al., 2014; He et al., 2011). Furthermore, the contribution of SOA in  $\text{PM}_{2.5}$  has been  
129 increasing in recent years, highlighting the necessity for better understanding the  
130 formation of SOA in this region (Wu et al., 2019; Wang et al., 2019). However, model  
131 simulation which provides robust information of the influence of physical processes

132 and chemical degradation in SOA formation still underpredict the SOA abundance with  
133 only traditional VOC precursors incorporated, hindering the better understanding the  
134 sources and formation mechanism of SOA in PRD (Wu et al., 2019; Fu et al., 2012;  
135 Wang et al., 2009). It was found that incorporating emissions of Gly and Mgly, and their  
136 degradation mechanisms could effectively narrow the gap between the measured and  
137 modelled SOA (Fu et al., 2012; Li et al., 2013a). However, only the simple  
138 parameterization of surface uptake of Gly and Mgly without detailed physical and  
139 chemical processes (e.g., reversible partitioning of Gly and Mgly into deliquesced  
140 droplets) in the model could bias the evolution of Gly and Mgly, leading to the poor  
141 understanding on the budgets of Gly and Mgly, their relationship with precursors, and  
142 the contributions of precursors to SOA formation in PRD (De Haan et al., 2018;  
143 Waxman et al., 2015; Knote et al., 2014; Li et al., 2013a, 2014; Lu et al., 2013).  
144 Therefore, to improve the model performance for the simulation of Gly and Mgly and  
145 to investigate their evolution and contribution to SOA formation, the observation data  
146 from a receptor site in the PRD region was analyzed by a photochemical box model  
147 with near-explicit chemical mechanisms (*i.e.*, the master chemical mechanism, MCM),  
148 and improvements with reversible and irreversible heterogeneous processes of Gly and  
149 Mgly, and the gas-particle partitioning of oxidation products in the present study. The  
150 production and evolution of Gly, Mgly, and other intermediate products were  
151 investigated. The observed and simulated levels of Gly and Mgly were compared to  
152 evaluate the performance of the model, which was further used to quantify the  
153 contributions of individual VOCs to SOA formation at the receptor site of PRD.

154

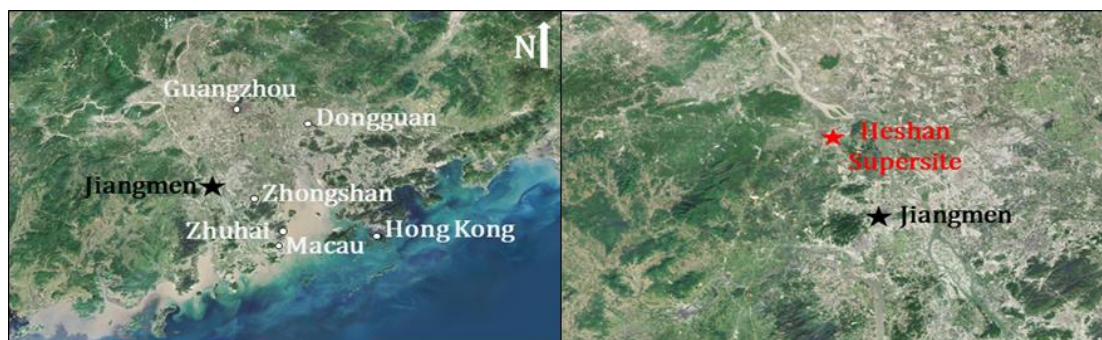
## 155 **2. Methodology**

### 156 **2.1. Field measurement**

157 Field measurements were carried out at Guangdong provincial atmospheric  
158 supersite located at Heshan (22.728°N, 112.929°E, 60 m above sea level) in Jiangmen  
159 City of the PRD region. The sampling site is located about 50 km and 80 km southwest  
160 from Foshan and Guangzhou City, respectively. The Heshan site is surrounded by  
161 mountain areas with trees and subtropical plants, and the location of the site is showed

162 in Figure 1. Ambient measurement of VOCs, carbonyls and other trace gases was  
163 conducted during January 02- 08, 2017, when the dominant wind was mainly from the  
164 southeast where the center of PRD (*i.e.*, Zhuhai and Zhongshan) was located. A detailed  
165 description of the Heshan site and the measurement methodology was provided in our  
166 previous studies (Chang et al., 2019; Yun et al., 2018).

167



168

169 Figure. 1 The location of sampling site and its surrounding environment in the Pearl River Delta  
170 region. The base map was from © Google Maps.

171

172 Briefly, mono-carbonyls, Gly and Mgly were collected with 2,4-  
173 dinitrophenylhydrazine cartridges every 3 h and detected using a high-performance  
174 liquid chromatography (HPLC) system (PerkinElmer 200 Series, US). The hourly  
175 VOCs were measured using a cryogen-free automatic gas chromatograph system  
176 equipped with a mass spectrometer and a flame ionization detector (GC-MS/FID)  
177 (Wang et al., 2014). CO, SO<sub>2</sub>, and O<sub>3</sub> was measured using a gas filter correlation  
178 analyzer, a pulsed fluorescence analyzer, and a UV photometric analyzer, respectively  
179 (Thermo Scientific 48i, 43i, 49i). NO and NO<sub>2</sub> were detected using a  
180 chemiluminescence instrument (Thermo Scientific 42i) with a photolytic converter (Xu  
181 et al., 2013). The method detection limits for non-methane hydrocarbons (NMHCs),  
182 carbonyls, CO, SO<sub>2</sub>, O<sub>3</sub>, NO, and NO<sub>2</sub> were 20-300, 20-450, 4000, 100, 500, 60, and  
183 300 pptv, respectively (Yun et al., 2018; Chang et al., 2019; Li et al., 2020). The  
184 measurement method uncertainty of the retrieved Gly and Mgly mixing ratios was  
185 estimated around 15% (Chang et al., 2019; Li et al., 2020). Furthermore, hourly  
186 meteorological parameters including temperature, wind speed, wind direction, pressure,

187 and relative humidity were recorded using a pyranometer (CMP22, Kipp & Zonen B.V.,  
188 Holland) and a portable weather station (Model WXT520, Vaisala, Finland).

189

## 190 ***2.2. Photochemical box model with master chemical mechanism (PBM-MCM) and*** 191 ***gas-particle partitioning SOA scheme***

192 Photochemical box model (PBM) was employed in this study to simulate the  
193 oxidation of Gly, Mgly and different VOCs, based on a master chemical mechanism  
194 (MCM) coupled with a gas-particle partitioning module to represent the SOA formation  
195 scheme. The MCM (version 3.2) is a near-explicit mechanism including ~16,500  
196 reactions involving ~6,000 chemical species with the latest IUPAC inorganic  
197 nomenclature, which described the chemical degradation of ~143 primary VOCs and  
198 their oxidation products. The MCM scheme has been applied to different photochemical  
199 box models to investigate the oxidation and reactivities of various VOCs, the formation  
200 of photochemical O<sub>3</sub> and secondary organic products, atmospheric radical budget and  
201 propagation, as well as the policy evaluation on mitigating the photochemical smog  
202 (*e.g.*, Ling et al., 2014, 2019; Wang et al., 2017; Lyu et al., 2015; Xue et al., 2014a, b).  
203 The physical processes including dry deposition and atmospheric dilution due to the  
204 variations of planetary boundary layer heights (configured according to the local  
205 observation in the PRD region from previous studies (Li et al., 2014; Wang et al., 2013;  
206 Fan et al., 2011) were considered in the model. Similar to other box models in  
207 simulating the degradation of VOCs and formation of SOA (Aumont, et al., 2012; Lee-  
208 Taylor, et al., 2011; Zhang and Seinfeld, 2013), the PBM-MCM model was developed  
209 by assuming a well-mixed box without consideration of vertical and horizontal  
210 transport, and air pollutants were assumed to be homogeneous (Lam et al., 2013; Ling  
211 et al., 2014). Thus, the influence of horizontal and vertical transport on air pollutants  
212 was not considered in this study.

213 In addition to the gas-phase degradation of VOCs, a gas-particle partitioning  
214 module for the oxidation products of VOCs and those compounds with an estimated  
215 normal boiling temperature greater than 450 K, as developed by Johnson et al (2006),  
216 were incorporated into the model to represent the SOA formation scheme (Johnson et  
217 al., 2005, 2006; Kamens et al., 1999; Stein et al., 1994). In brief, the gas-to-particle  
218 equilibrium partitioning of the species was described by the partitioning coefficient ( $K_p$ ,  
219 unit:  $\text{m}^3\mu\text{g}^{-1}$ ) using Eq. 1 (Johnson et al., 2006).

$$K_p = \frac{7.501 \times 10^{-9} RT}{MW_{om} \zeta P_L^0} \quad (\text{Eq.1})$$

where  $R$ ,  $T$ ,  $MW_{om}$ , and  $\zeta$  are the ideal gas constant ( $8.314 \text{ J K}^{-1} \text{ mol}^{-1}$ ), temperature (K), the mean molecular weight of the absorbing particle organic matter ( $\text{g mol}^{-1}$ ), and the activity coefficient of species in the condensed organic-phase, respectively.  $P_L^0$  is the liquid vapor pressure and was estimated using a semi-empirical expression of the Clausius-Clapeyron equation (Eq. 2):

$$\ln\left(\frac{P_L^0}{760}\right) = -\frac{\Delta S_{vap}(T_b)}{R} \left[1.8\left(\frac{T_b}{T} - 1\right) - 0.8\left(\ln\left(\frac{T_b}{T}\right)\right)\right] \quad (\text{Eq.2})$$

where  $T_b$  was the boiling temperature of different species, which was estimated using a previously described fragmentation method (Stein et al., 1994; Johnson et al., 2006).  $\Delta S_{vap}(T_b)$  was the vaporization entropy change at  $T_b$ , which was estimated using the Trouton-Hildebrand-Everett rule with corrections for polar compounds and compounds with hydrogen-bonding capacity (Baum 1997). The concentration of species  $j$  in the condensed organic-phase ( $F_{j,om}$ ) can be calculated as the following equation (Eq. 3):

$$F_{j,om} = M_{om} \times (K_{p,j} \times A_j) \quad (\text{Eq.3})$$

where  $M_{om}$  is the total mass concentration of each condensed organic material from gas-particle partitioning,  $A_j$  is its gas-phase concentration, and  $K_{p,j}$  is its partitioning coefficient of species  $j$  (Johnson et al., 2006).

The partitioning process was dynamically represented as an equilibrium between absorption and desorption, as described by Kamens et al. (1999). Briefly, the species-dependent  $K_p$  values were defined in terms of absorption ( $k_{in}$ ) and desorption ( $k_{out}$ ) rate coefficients, with  $K_p = k_{in}/k_{out}$ . The value for  $k_{in}$  ( $k_{in} = 6.2 \times 10^{-3} \text{ m}^3 \cdot \mu\text{g}^{-1} \cdot \text{s}^{-1}$ ) was configured as suggested by Johnson et al. (2006). Thus, the  $K_p$  could be expressed in terms of  $k_{out}$ . More detailed description of the equations and parameters are given by Johnson et al. (2005, 2006) and Kamens et al. (1999).

The above gas-particle partitioning of low volatility compounds formed by the gas-phase oxidation of VOCs and other precursors (Aumont, et al., 2012; Lee-Taylor et al., 2011) was configured in the model to estimate the SOA formation. However, the recent experimental results suggested that the formation of SOA in laboratory chambers may be suppressed due to losses of SOA to chamber walls, which leads to



249 underestimates of SOA in air-quality and climate models (Matsunaga and Ziemann  
 250 2010; Zhang et al., 2014). Therefore, to consider the wall loss of SOA, the average wall  
 251 loss rate coefficient of  $6 \times 10^{-5} \text{ s}^{-1}$  was adopted in the model configuration according to  
 252 previous studies on the basis of calculated organic material using an assumed density  
 253 of  $1 \text{ g} \cdot \text{cm}^{-3}$  (Johnson et al., 2004, 2005). In addition, the wall loss of other gaseous  
 254 compounds ( $\text{O}_3$ ,  $\text{NO}_2$  and  $\text{HNO}_3$ ) were implemented in the box model with the average  
 255 parameters of  $3 \times 10^{-6} \text{ s}^{-1}$ ,  $1.15 \times 10^{-5} \text{ s}^{-1}$  and  $8.2 \times 10^{-5} \text{ s}^{-1}$ , respectively. The detailed  
 256 information for the calculation of above parameters was provided in Bloss et al. (2015).

257

### 258 **2.3. Partitioning and reactions of gas-phase dicarbonyls on particles**

259 The partitioning and reactions of dicarbonyls in the aerosol aqueous phase may  
 260 involve both irreversible and reversible processes (Ervens and Volkamer, 2010). In the  
 261 present study, we follow the mechanism proposed by Knote et al. (2014) and consider  
 262 the reversible partitioning in aqueous phase, the irreversible volume reactions and  
 263 irreversible surface uptake processes in our model.

264 The reversible partitioning of Gly and Mgly on aerosols aqueous phase is usually  
 265 described by the Henry's law equilibrium (Kampf et al., 2013) (Eq.4):

$$266 \quad [Gly(Mgly)]_{liquid} = K_H \times [Gly(Mgly)]_{gas} \quad (\text{Eq.4})$$

267 However, hydration of carbonyls function groups and salt-Gly interactions could have  
 268 significant influences on the  $K_H$  value of Gly (Kampf et al., 2013; Waxman et al., 2015),  
 269 and an effective Henry's law coefficient expressed by Eq.5 was often used.

$$270 \quad K_{H, effective} = \frac{K_{H, water}}{10^{(-0.24 \min(12.0, (C_{as} + C_{an})))}} \quad (\text{Eq.5})$$

271 where the  $C_{as}$  and  $C_{an}$  represent the concentrations of ammonium sulfate and nitrate.  
 272 The detailed information on each parameter in these equations have been provided in  
 273 Kampf et al. (2013), Waxman et al. (2015) and the supplementary of the present study.  
 274 As variations were found for the value of  $K_{H, effective}$  under different concentrations of  
 275 ammonium sulfate and nitrate in previous studies (Knote et al., 2014; Kampf et al.,  
 276 2013; Ervens and Volkamer, 2010), the  $C_{as}$  and  $C_{an}$  were calculated every hour in the

277 present study from the measured ammonium sulfate (and ammonium nitrate)  
 278 concentrations ( $\text{mol m}^{-3}$ ) divided by aerosol liquid water content (ALWC,  $\text{kg m}^{-3}$ ),  
 279 which were determined by the aerosol inorganics model (AIM-IV,  
 280 <http://www.aim.env.uea.ac.uk/aim/model4/model4a.php>) with inputs of the observed  
 281 parameters (e.g., ambient relative humidity, temperature, and the moles of each ion) at  
 282 the Heshan site (Chang et al., 2019).

283 The reversible formation of monomer (i.e., glyoxal, glyoxal monohydrate, and  
 284 glyoxal dihydrate) and oligomers are considered with the two important reservoirs (i.e.,  
 285 monomer and oligomer pools, represented as pool1 and pool2) (Knote et al. 2014). The  
 286 variations of the glyoxal monomer ( $[\text{Gly}_{p1}]$ ) and oligomer concentrations ( $[\text{Gly}_{p2}]$ ) with  
 287 time can be represented by the following equations (Erverns and Volkamer, 2010;  
 288 Kampf et al., 2013; Knote et al., 2014):

$$289 \quad \frac{d([\text{Gly}_{p1}])}{dt} = \frac{1}{\tau_1} \times (\text{Gly}_{p1,eq} - \text{Gly}_{p1}) \quad (\text{Eq.6})$$

$$290 \quad \frac{d([\text{Gly}_{p2}])}{dt} = \frac{1}{\tau_2} \times (\text{Gly}_{p2,eq} - \text{Gly}_{p2}) \quad (\text{Eq.7})$$

$$291 \quad \frac{\text{Gly}_{p2,eq}}{\text{Gly}_{p1,eq}} = K_{olig} \quad (\text{Eq.8})$$

292 The equilibrium partitioning between monomers and oligomers was presented as  $K_{olig}$   
 293 (Eq.8). The definition and configuration of each parameters above were provided in the  
 294 supplementary (Section S2) according to Knote et al. (2014) and Kampf et al. (2013).

295 In addition, three irreversible pathways of Gly, including 1) the ammonium-  
 296 catalyzed volume pathway, 2) the OH-reaction volume pathway, and 3) the irreversible  
 297 surface uptake, were parameterized in the model (Knote et al., 2014; Ervens and  
 298 Volkman 2010). The ammonium-catalyzed reactions, with rate constant depending on  
 299 both particle acidity (pH) and the activity of the ammonium ion ( $a_{\text{NH}_4^+}$ ), were  
 300 parameterized as follows when the monomer and oligomer concentrations were in  
 301 equilibrium (Eq. 9):

$$302 \quad K = 2 \times 10^{-10} \times \exp(1.5 \times a_{\text{NH}_4}) \times \exp(2.5 \times \text{pH}) \times \text{Gly}_{p1} \quad (\text{Eq.9})$$

303 This parameterization was configured based on the assumption that only total  
 304 concentration in the monomer pool was the only particulate glyoxal available to the  
 305 ammonium-catalyzed reaction as the reversibly formed oligomers do not evaporate  
 306 easily (Knote et al., 2014; De Haan et al., 2009; Noziere et al., 2008).

307 For OH pathway, the gas-phase OH was in equilibrium with liquid-phase OH by  
 308 a Henry's law constant ( $K_{L,OH} = 25 \text{ M atm}^{-1}$ ) with the consideration of the "salting-in"  
 309 impact (Ervens and Volkamer 2010), and constant of reactions between OH and Gly  
 310 was  $1.1 \times 10^{-9} \text{ M}^{-1} \text{ s}^{-1}$  (Buxton et al., 1997). As suggested by Knote et al. (2014), the Gly  
 311 concentration available to the OH-reaction pathway was the total glyoxal concentration  
 312 in the monomer pool.

313 Surface-controlled irreversible uptake of Gly has been widely employed in  
 314 different modeling studies (Ervens et al., 2011; Li et al., 2014; Liu et al., 2007), was  
 315 parameterized as follows (Eq.10):

$$316 \quad K_r = -\frac{\gamma_{gly(mgly)} \times S_{aw} \times v_{gly(mgly)} \times C_{gly(mgly)}^*}{4} \quad (\text{Eq.10})$$

317 where  $C^*$  and  $v$  are the gas-phase concentration and mean molecular velocity,  
 318 respectively.  $\gamma$  represents the uptake coefficient for Gly and Mgly. Here we use the  
 319 surface uptake coefficients ( $\gamma_{gly} = 1.0 \times 10^{-3}$  and  $\gamma_{Mgly} = 2.6 \times 10^{-4}$ ) to account for the  
 320 irreversible surface uptake of Gly and Mgly, respectively. It is noted that the surface  
 321 uptake coefficient of Gly was configured according to the results of uptake kinetics  
 322 experiments from Schweitzer et al. (1998), which has been used in the model simulation  
 323 of Gly in the previous PRD study (Li et al., 2014). On the other hand, the surface uptake  
 324 coefficient of Mgly was obtained via scaling to glyoxal uptake coefficient by the  
 325 relative Henry's law coefficient suggested by Pye et al. (2017).  $S_{aw}$   
 326 ( $S_{aw} = S_a \times f(\text{RH}) = S_a \times (1 + a \times (\text{RH})^b)$ ) is the RH corrected aerosol surface area density (Li  
 327 et al., 2014). The value for  $a$  (2.06) was configured as those suggested previously (Liu  
 328 et al., 2007), while the dry aerosol surface concentration ( $S_a$ ) was obtained from the  
 329 measurement at the Heshan site (Yun et al., 2018). In this study, the mean molecular  
 330 velocities of Gly were calculated by the HyperPhysics model (<http://hyperphysics.phy->

331 astr.gsu.edu/hbase, last access date: 06 June 2019). The carbonaceous and insoluble  
332 components were considered as an aqueous shell for aerosols, whereas the aerosol  
333 surface was fully covered with an aqueous layer (Li et al., 2015).

334 On the other hand, though heterogeneous processes of Mgly are similar to those  
335 of Gly, some difference between these two species were found. The Henry's law  
336 constant for Mgly is not as effective as that for Gly. Hence, a Henry's law constant ( $3.7$   
337  $\times 10^3 \text{ M atm}^{-1}$ ) for Mgly we used (Zhou and Mopper 1990). In fact, Kroll et al. (2005)  
338 suggested that no obviously aerosol growth was observed from gas-phase Mgly  
339 presumably because of its more stable (less electron deficient) ketone moiety, and a  
340 recent study indicated that less Mgly would partition into the aerosols than expected  
341 according to Henry's law (Waxman et al., 2015). In addition, the surface uptake  
342 coefficient ( $\gamma_{\text{Mgly}} = 2.6 \times 10^{-4}$ ) suggested by Pye et al. (2017) is lower than that extracted  
343 from the chamber study (De Haan et al., 2018), which reported the value of  $\gamma_{\text{Mgly}}$  could  
344 increase to  $3.7 \times 10^{-3}$  at 95% RH and even larger than Gly in a high relative-humidity  
345 environment ( $\geq 95\%$ ). However, they also figured out that treating the surface uptake of  
346 Mgly on aerosols as an irreversible pathway could probably overestimate its positive  
347 effect for SOA formation via heterogeneous processes, because  $\sim 20\%$  of SOA which  
348 were formed from Mgly via aqueous processes would further hydrolyze.

349

#### 350 **2.4. Model scenarios**

351 According to the discussion above, it could be seen that the heterogeneous  
352 processes we described for Gly was more complicated than that for Mgly, as the  
353 parameterization for the sink of Gly from laboratory and model studies were more  
354 robust. Therefore, the present study put more emphasis on the evolution of Gly for  
355 better understanding and evaluating the effects of the different sink pathways on  
356 dicarbonyls and its influence on SOA formation. Table 1 provides detailed information  
357 regarding all the model scenarios for the simulation of Gly, while the model scenarios  
358 for Mgly are also given in Table S1 in the supplementary.

359

Table 1. Model scenarios used for gas-phase Gly

Scenarios	Description	Purpose
INITIAL	Default MCMv3.2, without considering the reversible and irreversible uptake of Gly and the gas-particle partitioning of other oxidation products	Base run
scenario 1	As INITIAL, also considers ammonium-catalyzed reactions of Gly through monomers pool 1 without the reversible formation of oligomers pool 2.	Investigating the influence of Ammonium reactions on the destruction of Gly
scenario 2	As scenario 1, also considers OH reactions of Gly through monomers pool 1 without the reversible formation of oligomers pool 2.	Investigating the influence of OH reactions for the destruction of Gly
scenario 3	As scenario 2, and considers the aqueous oligomers formation (pool 2) and revisable process with monomers (pool 1).	Investigating the “salting in” impact
scenario 4	As scenario 3, and considers surface uptake by aerosols of Gly with the uptake coefficient of $1 \times 10^{-3}$ suggested by Li et al. (2014).	Investigating the influence of surface uptake

360

361 In this study, hourly observation data of CO, SO<sub>2</sub>, NO, NO<sub>2</sub>, O<sub>3</sub>, NMHC and  
362 meteorological parameters were used as input and constraints in the model. By taking  
363 the NMHC species incorporating in the MCM mechanism into account (MCM website,  
364 <http://mcm.leeds.ac.uk/MCM/roots.htm>, access date: 22 June 2020), observations of  
365 total 44 NMHC species, including 18 alkanes, 11 alkenes, ethyne and 14 aromatics were  
366 used as input for the model simulation (Table S2 in the supplementary). The selected  
367 NMHCs contributed about 98% and 99% to the total mixing ratios and photochemical  
368 reactivities of all measured NMHCs at the Heshan site. Furthermore, the selected VOCs  
369 are the major precursors for Gly, Mgly, photochemical O<sub>3</sub> and SOA (Ding et al., 2016,  
370 2017; Li et al., 2014; Lou et al., 2010; Yuan et al., 2013), and have been frequently used  
371 to drive box model for studies on SOA, photochemical O<sub>3</sub> and photochemical reactivity  
372 (Hofzumahaus et al., 2009; Lee-Taylor, et al., 2011).

373 The photolysis rates, which were not measured, were modified in the model using  
374 the photon fluxes from the Tropospheric Ultraviolet and Visible Radiation (TUV-v5)  
375 model (Madronich and Flocke 1997) according to the sampling location and modeling  
376 period. Model simulation on Gly and Mgly was performed on January 07-08, 2017,  
377 when both daily Gly and Mgly data were available, with 00:00 LT (local time) as the

378 initial time. Before the simulation, the model was pre-run for 5 days using the observed  
379 variability of the input species during the whole sampling period to achieve a steady  
380 state for the unmeasured species with a short lifetime, *i.e.*, OH and HO<sub>2</sub> radicals (Xue  
381 et al., 2014a, b).

382 In this study, the simulation on the diurnal variations of OH and HO<sub>2</sub> was  
383 performed well, with peak values at noon, consistent with those measured and  
384 simulated in PRD (Hofzumahaus et al., 2009 and related papers; Tan et al., 2019). The  
385 simulated mean mixing ratios of OH and HO<sub>2</sub> radicals from the model in the present  
386 study were  $\sim 1.6 \times 10^6$  molecule·cm<sup>-3</sup> and  $\sim 3 \times 10^7$  molecule·cm<sup>-3</sup>, which are comparable  
387 to the winter observations at Beijing, Tokyo, and New York (Kanaya et al., 2007; Ren  
388 et al., 2006; Ma et al., 2019), and lower than the measurement and simulation values in  
389 summer (e.g., July) or autumn (e.g., October to November) in the PRD region (Table  
390 S3 in the supplementary) (Hofzumahaus et al., 2009; Tan et al., 2019). Note that the  
391 variations of simulation results in the present study and those observation results in  
392 previous studies in PRD may be associated with differences in the levels of O<sub>3</sub> and its  
393 precursors, different photolysis rates, and to a lesser extent, meteorological conditions  
394 (Hofzumahaus et al., 2009). The higher OH and HO<sub>2</sub> mixing ratios were expected in  
395 summer and autumn than winter due to the stronger solar radiation and higher  
396 temperature, as well as the variations of O<sub>3</sub> and its precursors in different sites, though  
397 the measurement of OH/HO<sub>2</sub> radicals has been very challenging, and significant  
398 uncertainties still exist in the measurement values of the radicals (Hofzumahaus et al.,  
399 2009; Tan et al., 2019). Furthermore, the comparison between the simulation of a box  
400 model and observation results suggested that the higher observed mixing ratios of OH  
401 and HO<sub>2</sub> radicals were related to an unidentified source of OH at the backgarden site of  
402 PRD in summer of 2006, while the comparison between the observed OH/HO<sub>2</sub>  
403 variations and those calculated from the parameterization of HO<sub>x</sub> (HO<sub>x</sub> = OH + HO<sub>2</sub>)  
404 production and destruction indicated a missing OH source of 4-6 ppbv·h<sup>-1</sup> and an  
405 unknown RO<sub>2</sub> loss at the Heshan site in autumn of 2014.

406 In addition to the simulation of OH and HO<sub>2</sub> radicals, as there were no direct  
407 measured SOA data in this study (Chang et al., 2019), the model performance was

408 evaluated by the comparison between the model simulated SOA with those calculated  
409 using the EC (elemental carbon)-tracer method, and by the comparison between the  
410 simulated and observed concentrations of other secondary products, which have been  
411 provided in detail in the supplementary (Section S3). For example, the simulated  
412 concentration of SOA was about 85% of those calculated by the EC-tracer method  
413 based on the observed hourly data (Chang et al., 2019). Furthermore, the simulated  
414 concentrations of acetic acid, formic acid and pyruvic acid were close to those observed  
415 at the Heshan site, accounting for ~80%, 70% and 88% of observed values for acetic  
416 acid, formic acid and pyruvic acid, respectively. The results confirmed that secondary  
417 formation was the dominant source of above species at the Heshan site, and suggested  
418 that the PBM-MCM model could provide robust performance on simulating the  
419 abundance of above secondary species and SOA.

420

## 421 **2.5. Model uncertainty**

422 Uncertainties in the simulation of Gly and Mgly by the model were noted. The total  
423 model errors could be calculated conservatively from 1) the uncertainties in the  
424 measurement of trace gases and NMHCs; 2) the measured data of meteorological  
425 parameters, *i.e.*, temperature  $T$ , pressure  $P$ , and the calculated photolysis frequencies  $J$   
426 based on meteorological conditions; 3) reaction rate constants  $k$ ; and 4) the dry  
427 deposition. In this study, following Li et al. (2014) and Lu et al. (2013), the uncertainty  
428 factors for the above parameters were adopted as suggested previously (Table S4 in the  
429 supplementary), and all parameters were divided into three groups (*i.e.*, physical  
430 parameters, radical and trace gas concentrations, and reaction rate constants of non-  
431 photolytic reactions). Each parameter was multiplied by its uncertainty factor first, and  
432 the gaussian error propagation was then applied within each group. We run the model  
433  $n$  times ( $n$  is the number of parameters considered). The mean diurnal variation of the  
434 uncertainty of modeled Gly and Mgly is shown in Figure S1. The total uncertainties of  
435 the modelled Gly and Mgly were both estimated to be around 39% with the  
436 contributions from radical and trace gas concentrations (~19%), physical parameters  
437 (~13%) (included photolysis frequencies, deposition lifetime,  $T$ , etc.) and reaction rate  
438 constants of non-photolytic reactions (~7%), respectively.

439

### 440 **3. Results and Discussion**

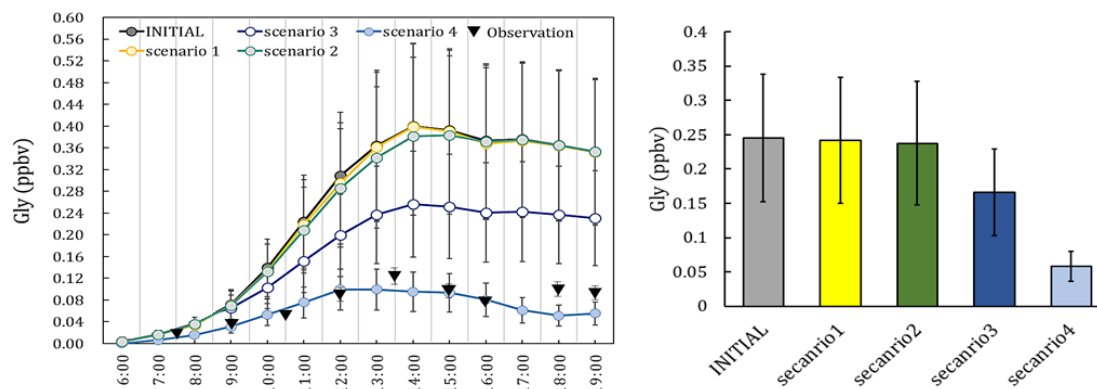
#### 441 **3.1. Comparison between the simulation and observation**

442 In this study, the simulated Gly and Mgly were secondarily formed from the  
443 oxidation of their VOC precursors. Therefore, before the comparison between the  
444 simulation and observation results, the contributions of primary and secondary sources  
445 to the measured Gly and Mgly were preliminarily estimated by a correlation-based  
446 source apportionment method suggested by previous studies (Friedfeld et al., 2002;  
447 Yuan et al., 2013). Table S5 in the supplementary shows linear regression coefficients  
448 and relative source contributions of Gly and Mgly. It was found that the contributions  
449 from primary sources (3.46% and 3.51% for Gly and Mgly, respectively) were  
450 significantly lower than those from secondary sources (96.14% and 96.44%,  
451 respectively), confirming that observed Gly and Mgly in the present study were mostly  
452 related to secondary formation.

453 The simulated Gly and Mgly from the photochemical box model under different  
454 scenarios were examined and compared with the observation. The predicted Gly and  
455 Mgly from in-situ formation in the INITIAL scenario was found to generally  
456 overpredict the mixing ratios of Gly and Mgly, and were about 3.3 and 3.5 times of the  
457 observed concentrations at the Heshan site, respectively. Similar overestimation has  
458 been reported in previous modeling studies, for example, the simulations with only the  
459 MCM gas-phases schemes overpredicted the Gly concentration by factors of 2-6 in both  
460 urban Mexico City (Volkamer et al., 2007) and a semi-rural site of the PRD region (Li  
461 et al., 2014). The significant overestimation in simulation results indicated that there  
462 were important loss pathways for Gly and Mgly other than the oxidation reactions (*e.g.*,  
463 by OH and NO<sub>3</sub> radicals). A sensitivity analysis was firstly conducted with twice the  
464 dilution rate (and deposition velocities) in INITIAL scenario, which resulted in the  
465 reduction of modeled concentrations of Gly and Mgly by 9.2% (3.2%) and 7.9% (2.8%),  
466 respectively (Table S6 in the supplementary). Though these enlarged rates were higher  
467 than the upper limits of the dilution rate and deposition velocities in previous studies  
468 (Fan et al., 2011; Wang et al., 2013; Li et al., 2014), the predicted mixing ratios of Gly  
469 and Mgly were still three times higher than the measured levels, suggesting that the



470 dilution and dry deposition configuration could not be the main causes for the  
471 overestimation of modeled Gly and Mgly mixing ratios (Volkamer et al., 2006, 2007).



472  
473 Figure 2. The observation data, the concentrations and the daily average concentration of Gly  
474 predicted from the different scenarios.

475  
476 To identify the causes of discrepancy and improve model performance, sensitivity  
477 analysis with different heterogeneous mechanisms incorporated in the model scheme (as  
478 scenarios listed in Table 1) was conducted. The average diurnal patterns of Gly  
479 simulated by different model scenarios are shown in Figure 2. It can be seen that in  
480 the early morning (*i.e.*, 0600 to 0800 LT, local time), the predicted mixing ratios of Gly  
481 in different scenarios were comparable to the observation. In contrast, the difference  
482 became larger from 0900 LT onwards, though the predicted peaks of Gly by different  
483 model scenarios were all presented at the early afternoon, following by a slow decrease  
484 in the late afternoon. The model results from Scenario 4 were more consistent with the  
485 observational data. The relative changes of modeled Gly concentrations by adding  
486 additional heterogeneous processes to the model scenario INITIAL (*i.e.*, model  
487 scenarios 1-4) were summarized in Table S7 in the supplementary.

488 On average, by including additional irreversible and reversible pathways, the  
489 modeled Gly concentrations during daytime (06:00-19:00) decrease by 72.3% of the  
490 values predicted by the INITIAL scenario, and a significant decrease of Gly  
491 concentration occurred by adding the effect of surface uptake pathway (*i.e.*, model  
492 scenarios 3-4). Similar results have been obtained in a previous study in summer in the  
493 PRD region (Li et al., 2014), which found that the simulated Gly concentration decrease  
494 significantly (~45 %) in the daytime (*i.e.*, 0600 to 1900 LT) when heterogeneous uptake  
495 process was considered with the incorporation of a single uptake coefficient.

496 The contribution of different heterogeneous sink pathways is calculated based on

497 the scenario 4 (Figure S2). The irreversible pathways of Gly (*i.e.*, surface uptake by  
498 aerosols, OH and ammonium reactions) accounted for 67.3% of the total sink of Gly,  
499 among which the surface uptake was a dominant pathway (62%) comparing to the  
500 ammonium and OH reactions (2.4% and 2.9%, respectively). The reversible pathway  
501 made a relatively lower contribution to the total sink of Gly (32.7%).

502 In addition, the heterogeneous irreversibly and reversibly pathways of Mgly was  
503 also investigated in the sensitivity analysis (*i.e.*, model scenarios M1-2, listed in Table  
504 S8 in the supplementary), and the predicted Mgly concentrations during daytime (0600-  
505 1900 LT) decreased by 73.0% of the values estimated in INITIAL model scenario. The  
506 surface uptake pathway in the scenario M2 was found to be the most important  
507 heterogeneous pathway for the loss of Mgly, and contributed to 64.1% of the total  
508 heterogeneous sink of Mgly. The contributions of the reversible pathway to the  
509 destruction of Mgly was around 35.9% (Figure S3).

510 Overall, by incorporating a more detailed heterogeneous processes of Gly and  
511 Mgly, the results of scenarios 4 and M2 provided better agreement between the modeled  
512 and measured Gly and Mgly. The results demonstrated the significance of  
513 heterogeneous uptake processes on the destruction of Gly and Mgly, and adopting the  
514 irreversible/reversible pathways (*i.e.*, the reversible partitioning, volume reactions, and  
515 the surface uptake) could reasonably reproduce the variations of Gly and Mgly at the  
516 Heshan site in the PRD region.

517

### 518 **3.2. Process analysis on the production and destruction of Gly and Mgly**

519 The scenarios 4 and M2 simulation with the best agreement with measurement  
520 were further analyzed to investigate the photochemical budget of Gly and Mgly at the  
521 Heshan site, respectively (Table 2 and Table S1). It was found that OH oxidation of  
522 aromatics was the most important contributor for the Gly and Mgly production, with  
523 mean contributions of ~80% and ~94%, respectively. Among all the aromatic  
524 precursors, toluene and *m,p*-xylene were the two major precursors for the formation of  
525 Gly and Mgly, with total contributions of ~43% and ~56% of Gly and Mgly formation,  
526 respectively. In contrast, because of the relatively low photochemical reactivity,  
527 benzene and alkanes had lower contributions to the formation of Gly and Mgly,  
528 although they can travel a long distance and contribute to secondary Gly and Mgly in

529 areas far from their emissions (Lv et al., 2019). Different from previous studies that  
530 found the isoprene as the key precursor for Gly and Mgly formation (Li et al., 2014;  
531 Lou et al., 2010), the contributions of isoprene oxidation at the Heshan site in the  
532 present study were much lower than that of aromatics, with only mean contributions of  
533 ~2% and ~3%, respectively. It can be attributed to the lower mixing ratios of isoprene  
534 (*i.e.*,  $70 \pm 10$  pptv) observed at the Heshan site because of the lower temperature in  
535 winter, comparing to the much higher concentration observed during summer (average  
536 of ~1 ppbv and maximum of ~4 ppbv in the afternoon) in the rural and forest areas in  
537 this region (Li et al., 2014; Lou et al., 2010).

538 In addition, the relative contributions of different loss pathways of Gly and Mgly,  
539 including physical processes (vertical dilution and dry deposition), reaction with  
540 radicals (e.g., OH and NO<sub>3</sub>), and the heterogeneous processes of Gly and Mgly on  
541 aerosols at daytime were also estimated from the PBM-MCM model results (Table 2).  
542 Consistent with previous studies (Atkinson and Arey, 2003; Ervens et al., 2011),  
543 heterogeneous processes were the most important pathway for the destruction of Gly  
544 and Mgly (with contributions of ~62% during daytime), followed by photolysis (with  
545 contributions of ~26% and ~25%, respectively). It should be noted that the oxidation  
546 of Gly and Mgly by O<sub>3</sub> was not considered in this study as the reaction rate constants  
547 of Gly and Mgly with O<sub>3</sub> are  $< 3$  and  $< 6 \times 10^{-21}$  cm<sup>3</sup>·molecule<sup>-1</sup>·s<sup>-1</sup>, respectively,  
548 which are 6 order of magnitude lower than the reaction rate constants with NO<sub>3</sub>  
549 (which  $> 1$  and  $> 2 \times 10^{-15}$  cm<sup>3</sup>·molecule<sup>-1</sup>·s<sup>-1</sup>, respectively), and are 9 order of  
550 magnitude lower than the reaction rate constants with OH ( $9$  and  $13 \times 10^{-12}$   
551 cm<sup>3</sup>·molecule<sup>-1</sup>·s<sup>-1</sup> for the reactions of Gly and Mgly with O<sub>3</sub>, respectively) (Mellouki  
552 et al., 2015). Therefore, we believe that the influence of O<sub>3</sub> on the removal of Gly and  
553 Mgly was negligible (Mellouki et al., 2015). Furthermore, there were few  
554 parameterizations for the reaction mechanism of Gly/Mgly with O<sub>3</sub> due to their low  
555 reaction rates with O<sub>3</sub>.

556 On the other hand, at nighttime, only the heterogenous processes made the main

557 contribution to Gly and Mgly destruction, with contributions higher than 90% to the  
 558 total destruction of Gly and Mgly at night (Table S9 in the supplementary), consistent  
 559 with previous studies (Washenfelter et al., 2011; Gomez et al., 2015). The lower  
 560 contributions of Gly and Mgly with radicals were mainly because of the low OH  
 561 concentration at night and their relatively lower reactivities with NO<sub>3</sub> radical (e.g., the  
 562 reaction rate constants of Gly/Mgly with NO<sub>3</sub> are ~1000 times lower than those with  
 563 OH radical) (Calvert et al., 2011; Mellouki et al., 2015).

564

565 Table 2 Production and destruction of Gly and Mgly from model simulation at daytime

Precursor	Oxidant	Gly		Mgly	
		Molar yield (%) <sup>a</sup>	Contribution (%)	Molar yield (%) <sup>a</sup>	Contribution(%)
<b>Aromatics</b>					
benzene, %	OH	32	4.94	-	-
toluene, %	OH	30.6	23.41	21.5	23.80
<i>m, p</i> -xylene, %	OH	25.2	19.22	35.1	32.08
<i>o</i> -xylene, %	OH	12.7	15.04	33.1	14.49
1,2,4-trimethylbenzene, %	OH	7.2	1.40	27.2	5.98
1,2,3-trimethylbenzene, %	OH	7.8	1.43	15.1	4.54
1,3,5-trimethylbenzene, %	OH	- <sup>c</sup>	-	58.1	13.21
ethylbenzene, %	OH	55	6.62	-	-
<i>p</i> -ethyltoluene, %	OH	31.9	5.45	-	-
<i>m</i> -ethyltoluene, %	OH	7.9	1.52	-	-
<i>o</i> -ethyltoluene, %	OH	8	0.51	-	-
<b>Sum</b>			79.54		94.10
<b>Alkanes</b>					
propane, %	OH	-	-	11	0.73
> C3 alkanes <sup>b</sup> , %	OH	1	0.19	3.2	0.71
<b>Sum</b>			0.19		1.44
<b>Alkenes</b>					
	OH	6.2	0.43	25	0.57
isoprene, %	NO <sub>3</sub>	43.7	1.34	37.8	2.83
	O <sub>3</sub>	4	0.20	-	-
ethene, %	OH	5.7	1.08	-	-
	O <sub>3</sub>	0.44	1.15	-	-

> C2 alkenes <sup>b</sup> , %	OH	-	-	7.7	1.06
propene, %	O <sub>3</sub>	8.3	1.01	-	-
1-pentene, %	O <sub>3</sub>	2	0.73	-	-
Sum			5.94		4.46
Acetylene	OH	63.5	14.33	-	-

### Loss pathways

photolysis, %	26.2	25.1
NO <sub>3</sub> ,OH-reaction, %	4.06	7.87
dry deposition, %	2.23	1.73
dilution, %	5.71	3.30
heterogeneous <sup>d</sup> , %		
Irreversible processes, %	41.0	39.8
Reversible processes, %	20.8	22.2

566 <sup>a</sup> Molar yields were taken from previous studies (Fu et al., 2008) (Fick et al., 2003) (Nishino  
567 et al., 2010) (Calvert 2000; Volkamer et al., 2006).

568 <sup>b</sup> >2 alkenes (include 3 alkenes) and >3 alkanes (include 17 alkanes) are represented in this  
569 study as a single lumped species (Lv et al., 2019).

570 <sup>c</sup> “-” not applicable.

571 <sup>d</sup> Considered both irreversible and reversible parameterizations of the aerosol sinks (*i.e.*,  
572 scenario 4 and M2 in the supplementary).

573

### 574 **3.3 Implications for secondary organic aerosol formation**

575 By incorporating both the traditional gas-particle partitioning (of VOC oxidation  
576 products) and the heterogeneous processes (of Gly and Mgly) into the model, we  
577 investigated the contributions of different mechanism in SOA formation through  
578 sensitivity analysis. The contributions of VOC oxidations to SOA formed from gas-  
579 particle partitioning (SOA<sub>gp</sub>) and SOA formed from heterogeneous processes of Gly  
580 and Mgly (SOA<sub>het</sub>) were quantified.

581 On the other hand, only based on the SOA<sub>gp</sub> formation scheme, the relative  
582 importance of each VOC precursor in SOA<sub>gp</sub> formation was further evaluated to  
583 provide a complete picture for SOA<sub>gp</sub> formation and its relationship with precursors.  
584 As with O<sub>3</sub> formation, the roles of individual VOC precursors in SOA<sub>gp</sub> formation were  
585 calculated using relative increment reactivity ( $RIR_{SOA_{gp}}$ ) method, which have been  
586 widely used to present the percentage change in the production of secondary products  
587 per percent change in precursors. The  $RIR_{SOA_{gp}}$  of a specific precursor  $X$  at site  $Z$  is  
588 given by Eq. 11:

$$RIR_{SOA_{gp}}^Z(X) = \frac{[P_{SOA_{gp}}^Z(X) - P_{SOA_{gp}}^Z(X - \Delta X)] / P_{SOA_{gp}}^Z(X)}{\Delta Z(X) / Z(X)} \quad (\text{Eq. 11})$$

where  $Z(X)$  represents the measured concentration of precursor  $X$ , including the amounts emitted at the site and those transported to the site, and  $\Delta X$  is the change in the concentration of precursor  $X$  caused by a hypothetical change  $\Delta Z(X)$  (10%  $Z(X)$  in this study). Here,  $P_{SOA_{gp}}^Z(X)$  represents the  $SOA_{gp}$  formation potential. A large positive  $RIR_{SOA_{gp}}$  value of a specific precursor suggests that  $SOA_{gp}$  formation could be significantly decreased if the emissions of this precursor were controlled. Figure 3 depicts the top 10 VOC precursors with high  $RIR_{SOA_{gp}}$  values at day time. Both *m,p*-xylene and *o*-xylene had the highest  $RIR_{SOA_{gp}}$  value (~0.35), followed by toluene (~0.2) and ethylbenzene (~0.06). As *m,p*-xylene, *o*-xylene and toluene can also have a significant impact on dicarbonyls production, they are likely to make a noticeable contribution to both  $SOA_{gp}$  and  $SOA_{het}$  formation.

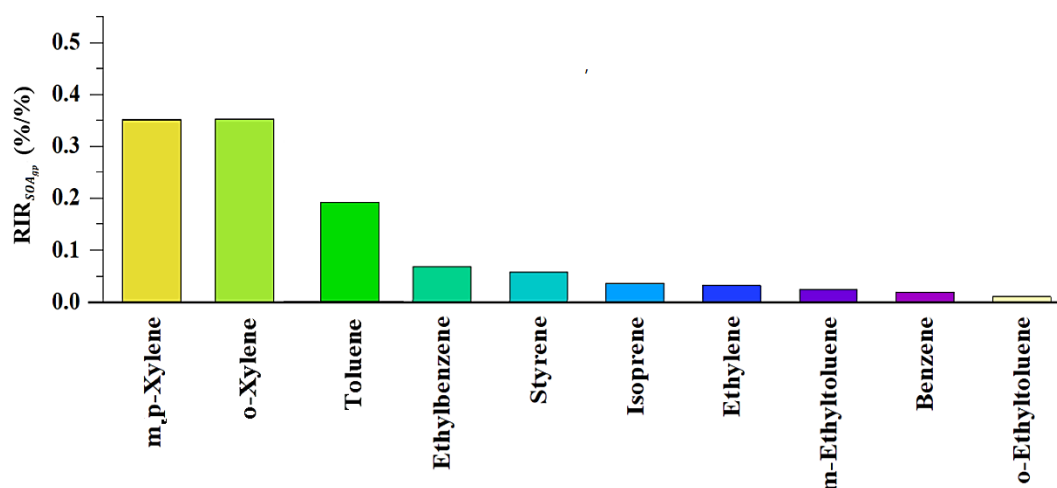
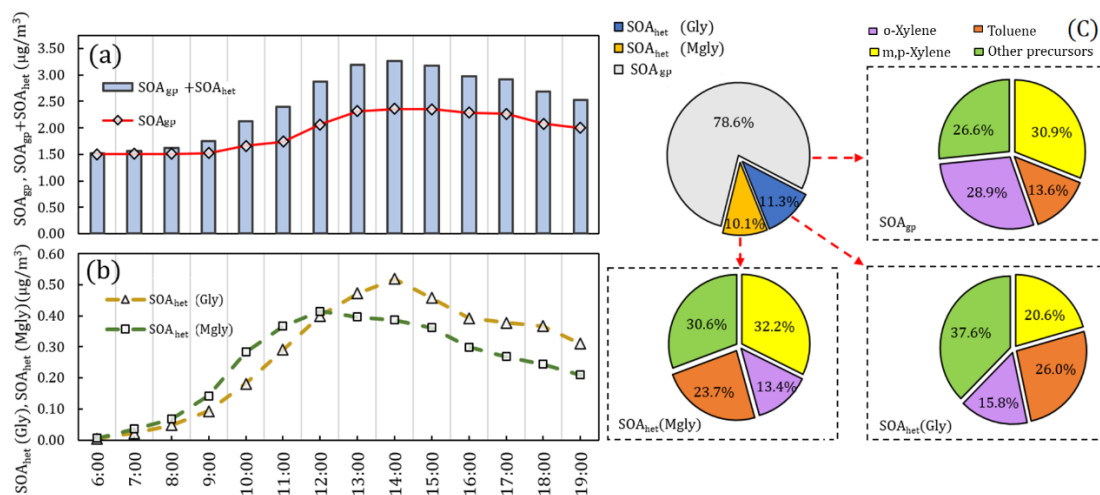


Figure 3. Top 10 VOC precursors with high  $RIR_{SOA_{gp}}$  values at daytime.

The SOA production from Gly and Mgly was further explored by the model simulation with and without the consideration of Gly and Mgly. It was found that by incorporating the evolution of Gly and Mgly, the SOA production has been improved apparently from 1.83 to 2.47  $\mu\text{g}/\text{m}^3$ . The total contribution of the Gly and Mgly contributed ~26% to the simulated SOA concentrations, of which ~21% was from the heterogeneous processes of Gly and Mgly ( $SOA_{het}$ ), further demonstrating that the heterogeneous processes have significant influences on the SOA formation from Gly and Mgly.

To further highlight the roles of heterogeneous processes of Gly and Mgly on the SOA production ( $SOA_{het}$ ) and to evaluate the contributions of different VOCs, the

613 average diurnal variations of SOA<sub>het</sub> concentration formed from the heterogeneous  
614 processes of Gly and Mgly were showed in Figure 4. Both SOA<sub>het</sub> (Gly) and SOA<sub>het</sub>  
615 (Mgly) concentrations presented photochemistry-driven diurnal patterns, and started to  
616 increase in the morning before reaching the maximum value (0.52 and 0.42  $\mu\text{g}/\text{m}^3$ ) at  
617 1400 and 1200 LT, respectively. It is consistent with the diurnal pattern of SOA<sub>gp</sub>, which  
618 could be formed from the oxidation of VOCs (including NMHCs and the gaseous  
619 oxidation of Gly and Mgly which were formed from the oxidation of NMHCs), due to  
620 the high photochemical reactivity at noon, which further converted to SOA<sub>gp</sub> through  
621 gas-particle partitioning. In general, SOA<sub>gp</sub> made a higher contribution to total SOA  
622 (78.6%) than SOA<sub>het</sub> (21.4%). Previous studies have indicated that the more abundant  
623 anthropogenic precursors than biogenic ones under NO<sub>x</sub> saturated environment could  
624 lead to greater contribution of SOA<sub>gp</sub> to total SOA despite that the oxidation of  
625 anthropogenic species (*i.e.*, aromatics) could lead to relatively higher yields of Gly and  
626 Mgly (Knote et al., 2014; Ervens et al., 2011). Ervens et al. (2011) has found that in  
627 areas with high concentrations of biogenic precursors at high relative humidity, the  
628 SOA<sub>het</sub> and SOA<sub>gp</sub> were equally important for total SOA, while in the anthropogenic  
629 dominated areas, the contribution of SOA<sub>het</sub> to the total predicted SOA mass was around  
630 30%. Similarly, the formation of SOA<sub>gp</sub> and SOA<sub>het</sub> were both dominated by xylenes  
631 and toluene, contributing to ~74%, ~62% and ~69% of SOA<sub>gp</sub>, SOA<sub>het</sub> (Gly) and SOA<sub>het</sub>  
632 (Mgly), respectively. Furthermore, *o*-xylene was the most important precursor to the  
633 SOA<sub>gp</sub> (~29%), but only contributed ~16% and ~13% to SOA<sub>het</sub> (Gly) and SOA<sub>het</sub>  
634 (Mgly) formation, respectively. The toluene and *m,p*-xylene made the most significant  
635 contributions to the SOA<sub>het</sub> (Gly) (~26%) and SOA<sub>het</sub> (Mgly) (~32%) formation,  
636 respectively, consistent with the chamber results from the oxidations of different  
637 precursors (Ervens et al., 2011).



638

639 Figure 4. The average diurnal variations of  $SOA_{gp}$ ,  $SOA_{het}$  (Gly) (*i.e.*,  $SOA_{het}$  formed from  
 640 heterogeneous processes of Gly),  $SOA_{het}$  (Mgly) (*i.e.*,  $SOA_{het}$  formed heterogeneous processes  
 641 from Mgly) and total SOA formation ( $SOA_{gp} + SOA_{het}$ ) were showed in Figure 4a and 4b. The  
 642 proportion of  $SOA_{gp}$ ,  $SOA_{het}$  (Gly) and  $SOA_{het}$  (Mgly) in total SOA as well as the contribution  
 643 of VOCs precursors to  $SOA_{gp}$ ,  $SOA_{het}$  (Gly) and  $SOA_{het}$  (Mgly) formation, were represented  
 644 in the pie charts in Figure 4c.

645

### 646 3.4 Comparison with previous studies in PRD

647 Previous studies have been conducted to investigate the evolution of Gly and/or  
 648 Mgly using observation and model simulation in the PRD region. However, one must  
 649 bear in mind that the mechanisms of the formation and evolution of Gly and/or Mgly  
 650 were more detailed in the present study, as previous studies in PRD only incorporated  
 651 the irreversible surface uptake process with a single coefficient for the heterogeneous  
 652 pathway for dicarbonyls. This may not reflect the current knowledge for the formation  
 653 and evolution of Gly and/or Mgly and their influence on SOA formation (Knote et al.,  
 654 2014; Waxman et al., 2015; Sumner et al., 2014).

655 Li et al. (2013a) used the regional air quality model CMAQ to investigate the  
 656 contributions of the aerosol surface uptake of Gly and Mgly to SOA formation in the  
 657 PRD region, and an uptake coefficient of  $2.9 \times 10^{-3}$  was used for both Gly and Mgly in  
 658 the model. Interestingly, the results from their model were about 30% higher than that  
 659 in our study (Table S10 in the supplementary). For example, Li et al. (2013a) concluded  
 660 that SOA formed from the heterogeneous processes of dicarbonyls may contribute  
 661 higher than 50% to the total SOA mass in the PRD region, while our study showed that  
 662 the contribution of  $SOA_{het}$  to total SOA mass was  $\sim 21\%$  (*i.e.*,  $\sim 11\%$  of  $SOA_{het}$  formed  
 663 from Gly;  $\sim 10\%$  of  $SOA_{het}$  formed from Mgly). In addition, the averaged concentration  
 664 of  $SOA_{het}$  from Gly ( $0.28 \mu\text{g}/\text{m}^3$ ) and Mgly ( $0.25 \mu\text{g}/\text{m}^3$ ) in our study is one order of



665 magnitude lower than that in Li et al. (2013a) (*i.e.*, 2.33 and 2.51  $\mu\text{g}/\text{m}^3$ , respectively).  
666 The discrepancy was mainly due to the different parameterizations of heterogeneous  
667 processes of dicarbonyls. The parameterization in the Li et al. (2013a) and other previous  
668 studies did not consider the reversible processes of dicarbonyls, but used one constant  
669 surface uptake coefficient to represent all the heterogeneous processes, which could  
670 result in bias in SOA formation if there are available aerosol surfaces without  
671 considering the influence of aerosols composition and phase state. Moreover, most of  
672 the previous studies using higher surface uptake coefficients intended to narrow the  
673 discrepancy between observed and simulated SOA mass without direct comparison  
674 between observed and simulated concentrations of dicarbonyls (*i.e.*, Li et al., 2013a;  
675 Waxman et al., 2013; Fu et al., 2008; Vokalmer et al., 2007). For example, Knote et al.  
676 (2014) conducted a total of seven simulations to investigate the SOA formation from  
677 Gly over California. Their results showed that the SOA concentration in SIMPLE  
678 scenario (characterized by a single uptake coefficient of  $3.3 \times 10^{-3}$ ) was an order of  
679 magnitude higher than that in HYBRID scenario (characterized by an uptake coefficient  
680 of  $1.0 \times 10^{-3}$  and also considered more comprehensive parameterization of  
681 heterogeneous processes). In fact, if we only consider the surface uptake by aerosols  
682 for dicarbonyls using the same uptake coefficient for dicarbonyls ( $2.9 \times 10^{-3}$ ) as Li et al.,  
683 (2013a), the contribution of  $\text{SOA}_{\text{het}}$  to total SOA mass would increase to 72% (*i.e.*, 37%  
684 of  $\text{SOA}_{\text{het}}$  formed from Gly; 35% of  $\text{SOA}_{\text{het}}$  formed from Mgly) (Table S10 in the  
685 supplementary). However, this configuration may not reflect the real evolution of  
686 dicarbonyls, resulting in the underestimation on the dicarbonyls concentrations (*i.e.*, the  
687 simulated concentration is at least one order of magnitude lower than the observation)  
688 (Figure S4) and overprediction of the contribution of  $\text{SOA}_{\text{het}}$  to total SOA mass in this  
689 study (~51%).

690 Table 3 compares the surface uptake coefficient derived from laboratory  
691 experiments and those used in different model simulation. It could be found that there  
692 was a large variation range for the surface uptake coefficients of Gly, while the studies  
693 on Mgly were still limited. For example, the laboratory experiment reported the surface  
694 uptake coefficients of Gly in the range of (0.8-6.6) and  $(\leq 1 - 9) \times 10^{-3}$  on aqueous  
695 inorganic aerosols and cloud droplet/ice crystals, respectively (Volkamer et al., 2007;  
696 Loggio et al., 2005), and the coefficients were found to be  $> 2.3 \times 10^{-3}$  for particles  
697 with high acidity (pH values within the range of -0.44 to -1.3) (Loggio et al., 2005). On

698 the other hand, Schweitzer et al. (1998) reported that the uptake coefficient of  $> 0.001$   
 699 was only observed for lower temperature conditions, and the experimentally measured  
 700 coefficient ranged from  $(1.2 \pm 0.06) \times 10^{-2}$  to  $(2.5 \pm 0.01) \times 10^{-3}$  on acidic solution (*i.e.*,  
 701 60-93 wt% H<sub>2</sub>SO<sub>4</sub>) at 253-273 k (Gomez et al., 2015; Zhang et al., 2015). It is suggested  
 702 that more accurate and comprehensive parameterization of heterogeneous processes of  
 703 dicarbonyls still needs deeper exploration for further model development. The  
 704 parameterization used in this study were mostly adopted from previous results, though  
 705 it may still have limitations and uncertainties, the results of simulation at this site show  
 706 better agreement with the observation.

707

708 Table 3 Surface uptake coefficient of Gly from laboratory experiments and used in the  
 709 model simulation in the present and previous studies

Coefficient	References
$(0.8-7.3) \times 10^{-3}$ , on aqueous inorganic aerosols	Volkamer et al., 2007; Loggio et al., 2005;
$(\leq 1 - 9) \times 10^{-3}$ , and on cloud droplet/ice crystals	Volkamer et al., 2007; Loggio et al., 2005;
$(1.2 \pm 0.06) \times 10^{-2} - (2.5 \pm 0.01) \times 10^{-3}$ on acidic solutions ( <i>i.e.</i> , 60-93 wt% H <sub>2</sub> SO <sub>4</sub> at 253-273 k)	Gomez et al., 2015; Zhang et al., 2015
$3.3 \times 10^{-3}$	Knote et al., 2014 and references therein; Waxman et al., 2013; Waxman et al., 2013
$2.9 \times 10^{-3}$	Fu et al., 2008
$1.0 \times 10^{-3}$	Knote et al., 2014 and references therein; this study; Li et al., 2014

710

#### 711 **4. Conclusion**

712 A photochemical box model coupled with MCM (v3.2) (PBM-MCM) and further  
 713 improvements on the evolution of semi- and non-volatility oxidation products to a  
 714 condensed particle-phase, was used to investigate the production and heterogeneous  
 715 processes of Gly and Mgly, as well as the SOA-precursor relationship at a receptor site  
 716 (*i.e.*, the Heshan site) for the first time in the PRD region. Compared to the  
 717 measurements, the initial model configuration overestimated the Gly and Mgly  
 718 concentrations by a factor of 3.3 and 3.5, respectively. This discrepancy occurred  
 719 largely due to the absence of irreversible uptake and reversible partitioning. Model  
 720 investigation regarding the production of Gly and Mgly revealed that the oxidation of

721 aromatics by OH radicals was the most important contributor to the formation of Gly  
722 and Mgly, with mean contributions of ~80% and ~94%, respectively, with toluene and  
723 *m,p*-xylene acting as the most important precursors for Gly and Mgly. For SOA  
724 formation, the heterogeneous processes of Gly and Mgly probably can explain ~21%  
725 of SOA mass in PRD. Toluene and *m,p*-xylene were the main precursors for SOA<sub>het</sub>  
726 formation, while *o*-xylene was the most important precursor of SOA<sub>gp</sub>. Overall, this  
727 study evaluated the formation and heterogeneous processes of Gly and Mgly in a  
728 polluted subtropical environment and highlighted the important role of intermediate  
729 products that are produced from photochemical oxidation of VOCs in SOA formation.  
730 The results of this study are expected to provide a better understanding of the evolution  
731 of VOC precursors, intermediate products, and heterogeneous process of the  
732 dicarbonyls, and the developed model modules can provide a robust tool for  
733 investigating SOA formation in the PRD and other regions in China.

734

#### 735 ***Author contributions***

736 In this study, the model was developed by ZL and QX. The whole structure for the  
737 paper was designed by ZL, XW and ZW. QX, ZL and ZW wrote the manuscript. ZW  
738 provided the observed data. All the authors have made substantial contributions to the  
739 work reported in the manuscript. ZL and QX contribute equally to this article.

740

#### 741 ***Data availability***

742 The underlying research data and the newly developed MCM scheme of Gly and Mgly  
743 in this study are available to the community and can be accessed by request to Zhenhao  
744 Ling (lingzhh3@mail.sysu.edu.cn) of Sun Yat-sen University.

745

#### 746 ***Competing interests***

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

748

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