

Photochemical aging of aerosols contributes significantly to the production of atmospheric formic acid

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Abstract

Formic acid (HCOOH) is one of the most abundant organic acids in the atmosphere and affects atmospheric acidity and aqueous chemistry. **However, the HCOOH sources are not well understood.** In a recent field study, we measured atmospheric HCOOH concentrations at a coastal site in South China. The average concentrations of HCOOH were 191 ± 167 ppt in marine air masses and 996 ± 433 ppt in coastal air masses. A strong linear correlation between HCOOH concentrations and the surface area densities of submicron particulate matter was observed in coastal air masses. Post-campaign laboratory experiments confirmed that the photochemical aging of ambient aerosols promoted by heterogeneous reactions with ozone produced a high concentration of HCOOH at a rate of 0.185 ppb h^{-1} under typical ambient conditions at noon time. HCOOH production was strongly affected by nitrate photolysis, as this efficiently produces OH radicals that oxidise organics to form HCOOH. We incorporated this particle-phase source into a photochemical model and **the net HCOOH production rate increased by about three times compared with the default Master Chemical Mechanism (MCM).** These findings demonstrate that the photochemical aging of aerosols is an important source of HCOOH that **should** be included in atmospheric chemistry-transport models.

1. Introduction

Organic acids are ubiquitous in the troposphere and constitute a significant fraction of the total organics in both the gas and particle phases (Chebbi and Carlier, 1996). They also participate in the aqueous-phase chemistry of clouds, contribute to secondary organic aerosol

33 (SOA) formation through reactions within the condensed phase (Carlton et al., 2007; Ervens et
34 al., 2004; Lim et al., 2010), and are proposed to enhance the formation of new particles in the
35 atmosphere (Zhang et al., 2004). Formic acid (HCOOH) is among the most abundant organic
36 acids in the atmosphere (Khare et al., 1999) and accounts for over 60% of the free acidity in
37 precipitation in remote areas and more than 30% of that in polluted areas (Andreae et al., 1988;
38 Keene et al., 1983; Keene & Galloway, 1988; Khare et al., 1999; Stavrou et al., 2012). This
39 contribution is increasingly important due to the decline in the concentrations of anthropogenic
40 nitrogen oxides (NO_x) and sulfur dioxide. HCOOH serves as a significant sink of in-cloud
41 hydroxyl radicals ($\cdot\text{OH}$) and stabilised Criegee intermediates (SCIs) (Jacob, 1986), and thus
42 influences aqueous-phase chemistry by affecting pH-dependent reaction rates, oxidant
43 concentrations, and solubilities (Vet et al., 2014). HCOOH also plays a role in the formation of
44 cloud condensation nuclei (Yu, 2000), due to its comparatively higher hygroscopicity at low
45 critical supersaturations when incorporated into aerosols (Novakov and Penner, 1993). This, in
46 turn, affects total indirect radiative forcing. Additionally, HCOOH may be involved in halogen
47 chemistry through its heterogeneous reaction with solid sodium chloride in sea-salt aerosols
48 (Xia et al., 2018).

49 Considering the abovementioned roles of HCOOH in atmospheric chemistry, it is essential
50 to understand its sources and sinks. The main sources of HCOOH include direct emissions
51 from terrestrial vegetation (Andreae et al., 1988), biomass and biofuel burning (Akagi et al.,
52 2011; Goode et al., 2000; Yokelson et al., 2009), fossil-fuel combustion (Kawamura et al., 2000;
53 Zervas et al., 2001a, b) and soil emissions (Sanhueza and Andreae, 1991). **The secondary gas-**
54 **phase formation mechanisms of HCOOH are mainly** the oxidation of volatile organic
55 compounds (VOCs), including ozonolysis of terminal alkenes (Neeb et al., 1997), alkyne
56 oxidation (Bohn et al., 1996), OH-initiated isoprene oxidation (Paulot et al., 2009),
57 monoterpene oxidation (Larsen et al., 2001), keto-enol tautomerisation (Andrews et al., 2012;
58 Shaw et al., 2018) and $\cdot\text{OH}$ oxidation of methyldioxy radicals ($\text{CH}_3\text{O}_2\cdot$) (Bossolasco et al.,
59 2014). HCOOH is primarily removed from the atmosphere through wet and dry deposition,
60 with a minor sink of being photo-oxidation by $\cdot\text{OH}$ (Atkinson et al., 2006). **However, with**
61 **these mechanisms included, HCOOH concentrations** remain significantly underestimated by
62 **previous models** (Baboukas et al., 2000; Bannan et al., 2017; Chaliyakunnel et al., 2016; Le
63 Breton et al., 2012; Millet et al., 2015; Yuan et al., 2015), indicating a substantial missing source
64 of HCOOH.

65 **In addition to gas-phase production pathways, HCOOH can also be generated through**
66 **heterogeneous or condensed-phase processes.** Aqueous reactions of formaldehyde (HCHO)
67 (Chameides and Davis, 1983; Jacob, 1986), glyoxal (Carlton et al., 2007) and other species
68 with $\cdot\text{OH}$ (aq) can produce HCOOH, particularly in moderately acidic environments (Jacob,
69 1986). **A recent chamber experiment has revealed that formaldehyde can be efficiently**
70 **converted to HCOOH through a multiphase pathway that involves its hydrated form,**

71 methanediol. This pathway has been shown to generate up to four times more formic acid
72 compared to all other known chemical sources combined in a chemistry-climate model, and
73 the modified model largely reproduced observed ambient concentrations of HCOOH (Franco
74 et al., 2021). Moreover, Gao et al. (2022) proposed a new bidirectional deposition-emission
75 process, in which HCOOH deposits rapidly in night-time dew and is re-emitted subsequently
76 from the dew as it evaporates on the following day.

77 Many laboratory chamber studies have demonstrated that the photochemical aging of
78 organic aerosols can also produce HCOOH (Henry and Donahue, 2012; Malecha and
79 Nizkorodov, 2016; Mang et al., 2008; Pan et al., 2009; Walser et al., 2007; Zhang et al., 2021).
80 The photochemical aging of aerosols occurs through the reactive uptake of oxidants onto
81 particle surfaces, altering their chemical compositions and physical properties (George et al.,
82 2015). In the condensed organic phase, HCOOH can be produced through the photodegradation
83 of SOA (Henry and Donahue, 2012; Malecha and Nizkorodov, 2016). Additionally, the
84 oxidants such as $\cdot\text{OH}$, nitrogen dioxide (NO_2), and nitrite ions/nitrous acid (HONO) produced
85 from the photolysis of particulate nitrate (NO_3^-) can also efficiently oxidise organics to produce
86 HCOOH (Zhang et al., 2021). Apart from laboratory experiments, Paulot et al. (2011) observed
87 a marked positive correlation between HCOOH concentrations and submicron organic aerosol
88 masses in field measurements conducted in three distinct areas: coastal, urban, and polar, and
89 suggested that aerosol aging produces HCOOH. The aforementioned results show that there is
90 a need for a quantitative assessment of the contribution of the photochemical aging of aerosols
91 to HCOOH production in the ambient atmosphere.

92 In this study, we measured HCOOH concentrations at near-ground level at a coastal site in
93 Hong Kong, China, for two months during autumn 2021. We examined the characteristics of
94 HCOOH concentrations and their correlation with related species' concentrations or other
95 parameters. We then conducted a chamber study to measure the rate of HCOOH production
96 during the aging of ambient aerosols and extrapolated the results to the real atmosphere. We
97 incorporated this HCOOH-formation mechanism into a model using a parameterisation
98 involving fine particulate matter concentration, surface area density, light intensity, and ozone
99 (O_3) concentration, and then performed simulations to evaluate the HCOOH production rate
100 from the photochemical aging of aerosols. Furthermore, we showed that NO_3^- photolysis acted
101 as a crucial source of $\cdot\text{OH}$ during the aging process. Our results enhanced the understanding of
102 HCOOH sources and model simulations of ambient HCOOH concentrations.

103 2. Methods

104 2.1. Field observations

105 Ambient measurements of the atmospheric concentrations of HCOOH and related
106 species/parameters were conducted from 13 August to 31 October 2021 at the Hong Kong

107 Environmental Protection Department's Cape D'Aguilar Super Site (CDSS; 22.21°N,
108 114.25°E), which is situated a few hundred meters away from the nearest coastline of the South
109 China Sea (Fig. S1). During the study period, the air quality of this coastal site was initially
110 predominantly influenced by marine air masses from the South China Sea and subsequently by
111 the coastal air masses transporting regional anthropogenic pollution from East China. The site
112 was also affected by biogenic emissions from local vegetation and ship emissions transported
113 mainly from about 8 kilometres away. There were no other significant anthropogenic sources
114 nearby.

115 The species quantified were HCOOH, HONO, trace gases (nitric oxide, NO₂, carbon
116 monoxide (CO), sulfur dioxide (SO₂), and O₃), volatile organic compounds (VOCs),
117 oxygenated VOCs, aerosol mass concentrations (1-µm particulate matter (PM₁), PM_{2.5}, and
118 PM₁₀ concentrations), aerosol size distributions, aerosol ionic compositions, NO₂ photolysis
119 frequency (jNO₂), and meteorological parameters (temperature (T), relative humidity (RH),
120 wind direction, and wind speed). A detailed description of the measurement of HCOOH
121 concentration is provided below, and information on other measurements is summarised in
122 Table S1.

123 The concentration of HCOOH was measured at 172.91 atomic mass unit (amu) using an
124 iodide-adduct time-of-flight chemical ionisation mass spectrometer (I⁻-ToF-CIMS, Aerodyne
125 Research), as we used in our previous study at the same site (Xia et al., 2022). A comprehensive
126 description of the use of the I⁻-ToF-CIMS can be found in previous studies (Lee et al., 2014;
127 Aljawhary et al., 2013). Briefly, humidified iodomethane-containing N₂ air was passed through
128 an inline ioniser (containing polonium-210) to generate iodide ions (I⁻) and iodide-water ions,
129 which served as the reagent ions, and the HCOOH produced was detected as IHCOOH⁻. The
130 background concentration of HCOOH was determined every 2 days by injecting zero air and
131 was found to be 60.9 ppt. HCOOH calibration was performed three times on-site and once in
132 the laboratory immediately following the field campaign using diluted gas standards generated
133 by a permeation tube (KIN-TEK) with a permeation rate of 90.87 ng min⁻¹. HCOOH sensitivity
134 varies with RH as water competes with HCOOH for I⁻ (Lee et al., 2014). Thus, the HCOOH
135 sensitivity was measured at various RHs, as shown in Fig. S2. The sensitivity remained stable
136 at a given RH, with a variation of less than 5% throughout the campaign.

137 The ToF-CIMS was housed in an air-conditioned shelter at an indoor air T maintained at
138 25–28 °C. The shelter was located approximately 15 m away from the CDSS station. The
139 sampling tube was a 0.5-m long perfluoroalkoxy-Teflon tube (1/2 in. outer diameter), the inlet
140 of which was situated on the sidewall of the shelter, 1.5 m above the ground. To achieve laminar
141 flow in the sampling tube, a flow rate of 25 Lpm was adopted, with a residence time of 0.1 s.
142 The ToF-CIMS drew ~2 Lpm sample air, and the remaining airflow was discarded. The
143 sampling tube was replaced with a new tube every 2 days to reduce inlet artifacts. We
144 investigated possible inlet artifacts by injecting known concentrations of HCOOH into a used

145 sampling inlet and found that the artifacts had a negligible effect on the measured HCOOH
146 concentration (difference < 3%). The remaining instruments were housed in the CDSS station,
147 with their sampling inlets located ~1.5 m above the roof.

148 2.2. Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) and Extended 149 Aerosol Inorganic Model IV (E-AIM IV) models

150 Hourly 24-h backward trajectories were obtained using the HYSPLIT model
151 (<https://www.ready.noaa.gov/hypub-bin/trajsrc.pl>). The input parameters were Global Data
152 Assimilation System 1° for the meteorology data; 22.21°N and 114.25°E for the location; and
153 60 m for the endpoint height. Air masses were classified as marine or coastal, based on their
154 source regions. A unique period was identified as a haze period, during which there was a
155 transition from marine air masses to coastal air masses under stagnant conditions (wind speed
156 < 3 m s⁻¹). A detailed classification can be found in Fig. S3.

157 The aerosol water content (AWC) and pH were predicted by the E-AIM IV online in batch
158 mode (<http://www.aim.env.uea.ac.uk/aim/model4/model4d.php>). The thermodynamic model
159 was constrained by hourly field-measured molar concentrations of NH₄⁺, sodium ions (Na⁺),
160 sulfate ions (SO₄²⁻), NO₃⁻, and chloride ions (Cl⁻) in PM_{2.5}; gas-phase ammonia concentrations;
161 ambient T; and RH. Initial concentrations of protons (H⁺) and hydroxide ions were estimated
162 based on the ion balance of the major water-soluble ions. The input of Na⁺ into the model
163 (Na⁺_(eq)) was calculated as a sum of the equivalent concentrations of Na⁺, potassium ions,
164 magnesium ions, and calcium ions (Eq. (1)). The model also considered water dissociation and
165 allowed all possible solids to form in the system. Parameters p, q, r, and s (options in batch
166 mode) were set to 3, meaning that the input portions of ammonium (NH₄⁺), NO₃⁻, SO₄²⁻, and
167 Cl⁻ always remained in the condensed phase and did not produce corresponding gas-phase
168 species, such as nitric acid (HNO₃) and hydrochloric acid. The aqueous-phase NO₃⁻
169 concentration was calculated by dividing the moles of aqueous NO₃⁻ by the AWC. Aerosol pH
170 was calculated as the negative logarithm of the concentration of H⁺.

$$171 [Na^+]_{eq} = [Na^+] + [K^+] + 2[Mg^{2+}] + 2[Ca^{2+}] \quad (1)$$

172 2.3. Laboratory experiments

173 We illuminated ambient aerosols collected on filters or aqueous solutions in a dynamic
174 chamber to mimic HCOOH formation in the atmosphere. The overall experimental setup is
175 illustrated in Fig. S4. The dynamic chamber has a dimension of 25-cm length × 15-cm width ×
176 4-cm height with the top side sealed by a transparent Teflon film. Each aerosol filter or solution
177 sample was placed in a quartz Petri dish (inner diameter: 35 mm, inner height: 7 mm) at the
178 chamber's centre. Aerosol filter sampling details can be found in Text S1. A high-pressure
179 xenon (Xe) lamp was used to simulate sunlight and its spectral irradiance is displayed in Fig.
180 6d. Compared with standard air mass 1.5 solar irradiation (AM 1.5) corresponding to a solar
181 zenith angle of 48.2°, the Xe lamp exhibited a smaller flux at 300–326 nm but a larger flux at
182 326–420 nm. An air stream from a zero-air generator (EnviroNics, model 7000) served as the

183 carrier gas that delivered reaction products to the chamber's outflow. The weather conditions
184 that prevailed during the field campaign were mimicked by maintaining the reactor's T at
185 approximately 28 °C and adjusting the RH in the chamber to 70% by passing the carrier gas
186 through a water bubbler. Prior to the introduction of a sample into the chamber, the background
187 HCOOH concentration was monitored for 10 min with the light on and zero air added. After 1
188 h of irradiation, 100 ppb of O₃ was introduced via a dynamic calibrator (EnviroNics® Series
189 6100) and monitored using an O₃ analyser (Thermo Scientific Model 49i). An AM 1.5 filter
190 (which removes light below 360 nm) and a 300–800 nm filter (which allows the passage of
191 300-800 nm light) were applied to the Xe lamp to investigate the effect of the irradiation
192 wavelength on the formation of HCOOH.

193 We conducted an experiment to explore the role of nitrate photolysis on the proposed
194 particle-phase mechanism. A sample solution was prepared by mixing formaldehyde (HCHO,
195 Sigma-Aldrich, 37 wt% in water) and sodium nitrate (NaNO₃, Honeywell, 99.5% purity). The
196 resulting solution contained 0.15 wt% HCHO and 0.2 M NaNO₃, and was adjusted to pH 2.7
197 by the addition of sulfuric acid (H₂SO₄, Sigma-Aldrich, 98% purity), as this was the E-AIM
198 model's prediction of the average aerosol acidity during the entire campaign. We assumed
199 that ·OH produced by NO₃⁻ photolysis was the rate-limiting species due to its significantly
200 lower abundance relative to organics and HCHO was taken as a representative example of
201 potential precursors of HCOOH. However, it is important to note that the production rate of
202 HCOOH from the oxidation of organics may be overestimated in the solution experiment
203 because ·OH generated by nitrate photolysis can also react with other oxidizable species in the
204 ambient atmosphere.

205 The average rate of production (ppb·s⁻¹) of HCOOH (P_{HCOOH}) during the 1-h irradiation
206 was calculated by the following equation (Eq. (2)), derived from (Peng et al., 2022):

$$207 \quad P_{\text{HCOOH}} = \int_0^{60} (C_{\text{HCOOH}} - C_{\text{HCOOH-bkg}}) dt \times \frac{Q}{V} / 60 \quad (2)$$

208 where Q is the carrier gas flow rate (4 L min⁻¹); V is the reactor chamber volume (1.875 L);
209 and C_{HCOOH} and C_{HCOOH-bkg} (ppb) are the concentrations of HCOOH in the chamber after and
210 before adding the sample, respectively. The photolytic loss of HCOOH was ignored, as the
211 cross-section of HCOOH was beyond the spectral range of the Xe lamp (Burkholder et al.,
212 2020).

213 We attempted to extrapolate the laboratory results to account for the field-observed
214 concentrations of HCOOH. As photochemical aging occurs on aerosol surfaces and a strong
215 correlation between the surface area (Sa) and the concentration of HCOOH observed in the
216 field, the extrapolation was conducted based on Sa. The Sa in the chamber was calculated as
217 the Sa of the filter divided by the chamber's volume, assuming that only the first layer of the
218 aerosols was illuminated. Although this might have resulted in an underestimation of the Sa
219 density in the chamber, this assumption was reasonable, because particles in the lower layers

220 would receive less light than those in the uppermost layer due to the light-screening effect of
221 the first layer (Ye et al., 2017). The aging process was also influenced by light intensity, and as
222 we discovered that the major oxidant was generated by the photolysis of particulate NO_3^- (See
223 **Results**, section 2), the light intensity was normalised based on the photolytic frequency of
224 aqueous NO_3^- ($J_{\text{NO}_3^-(\text{aq})}$) due to the absence of an absorption coefficient for particulate NO_3^- .
225 Although there is a redshift of the particulate NO_3^- absorption wavelength compared with the
226 aqueous-phase NO_3^- absorption wavelength, our results should be reliable because we used
227 $J_{\text{NO}_3^-(\text{aq})}$ as a reference for normalisation rather than for calculating an accurate $J_{\text{NO}_3^-(\text{aq})}$ (Du
228 and Zhu, 2011; Zhu et al., 2008). The $J_{\text{NO}_3^-(\text{aq})}$ under the Xe lamp was $8.85 \times 10^{-6} \text{ s}^{-1}$ and the
229 daytime average $J_{\text{NO}_3^-(\text{aq})}$ in the ambient air at our site was $1.12 \times 10^{-5} \text{ s}^{-1}$ (Text S2). Both
230 $J_{\text{NO}_3^-(\text{aq})}$ values were calculated assuming a quantum yield equal to 1. The normalised HCOOH
231 production rate in the ambient air ($P_{\text{HCOOH-nml}}$) was calculated using the following equation (Eq
232 (3)):

$$233 \quad P_{\text{HCOOH-nml}} = P_{\text{HCOOH}} \times \frac{S_{\text{amb}}}{S_{\text{cha}}} \times 1.266 \quad (3)$$

234 where S_{amb} represents the field-measured Sa density; S_{cha} denotes the Sa density calculated
235 for the chamber; and 1.266 is the ratio of the ambient $J_{\text{NO}_3^-(\text{aq})}$ to the chamber $J_{\text{NO}_3^-(\text{aq})}$. For the
236 aging process involving O_3 , the photolytic rate constant of O_3 generating O^1D ($J_{\text{O}_3 \rightarrow \text{O}^1\text{D}}$) in the
237 chamber ($1.31 \times 10^{-5} \text{ s}^{-1}$) was also normalised to the average daytime $J_{\text{O}_3 \rightarrow \text{O}^1\text{D}}$ ($1.84 \times 10^{-5} \text{ s}^{-1}$)
238 under ambient conditions (Text S2). For the results of the aqueous solution, the
239 concentrations of HCHO and NO_3^- were also normalised.

240 2.4. Chemical box model

241 The Framework for 0-D Atmospheric Modeling (F0AM version 4.2.1) (Wolfe et al., 2016)
242 coupled with Master Chemical Mechanism (MCM v3.3.1, <http://mcm.york.ac.uk>) (Jenkin et
243 al., 2015) was employed to **evaluate the HCOOH production rate from photochemical aging of**
244 **aerosols compared with other sources** at the field site. The **physical loss due to dilution process**
245 was treated as a first-order loss with a rate coefficient of $1/86,400 \text{ s}^{-1}$, consistent with previous
246 studies (Li et al., 2014; Yuan et al., 2015). The dry deposition rate was determined based on the
247 deposition velocity and the boundary layer height (BLH). For HCOOH, a deposition velocity
248 of 1 cm s^{-1} was employed (Müller et al., 2018). A sensitivity analysis of the deposition velocity
249 was also conducted. The diurnal profile of BLH was acquired from a previous study at another
250 coastal site in Hong Kong (Su et al., 2017). Wet deposition was not considered as there was no
251 rainfall **except on 3 October and 7 October. We simulated the averaged diurnal cycle for the**
252 **whole campaign with field-observed relevant species constrained hourly in the model.**
253 **Simulations were also performed daily for a 2-week period, from 24 September to 7 October.**
254 **The details of input data are described in Text S3.** The model was executed for three replicates
255 to stabilise the intermediate species it generated, and the results from the final run were used

256 for further analysis.

257 The local sources of HCOOH at this site mainly consist of ship and biogenic emissions.
258 The box model used in this study did not account for the contribution of ship emissions since
259 the modelling period was dominated by continental outflows which is unfavourable for the
260 transportation of HCOOH from ship emissions to our site as evidenced by the relatively low
261 concentrations of NO_x in the modelling period. To estimate the biogenic emissions, we used
262 the algorithm of the Model of Emissions of Gases and Aerosols from Nature version 2.1
263 (MEGAN v2.1) (Guenther et al., 2012), assuming instantaneous dilution into the whole box. It
264 is worth noting that this approach may introduce some uncertainties in estimating biogenic
265 emission at a specific location. The specific parameters used can be found in Text S3. For
266 secondary formation mechanisms, we followed Yuan et al. (2015) by enhancing the secondary
267 production of HCOOH by modifying the HCOOH yields obtained by the ozonolysis of alkenes
268 and other unsaturated species, and by adding chemical processes (vinyl alcohol oxidation;
269 photo-tautomerisation of vinyl alcohol; and the reactions of HCHO + HO₂ and CH₃O₂· + ·OH)
270 to the MCM. However, the model used in this study was unable to account for the downward
271 transport of HCOOH produced in clouds through a newly proposed multiphase pathway
272 (Franco et al., 2021) due to its inability to assess the contribution of vertical mixing and aqueous
273 phase chemistry.

274 3. Results and Discussion

275 3.1. Field measurements of HCOOH concentrations

276 The field site was exposed to two distinct types of air masses; initially, it was largely
277 exposed to marine air masses, and later to coastal air masses. Marine air masses ($T = 29.4 \pm$
278 2.0 °C, $RH = 85.8 \pm 7.0$ %) were warmer and more humid than coastal air masses ($T = 25.7 \pm$
279 2.3 °C, $RH = 77.0 \pm 6.0$ %), and exhibited low concentrations of O₃ (15.0 ± 8.9 ppb) and high
280 concentrations of NO_x (6.2 ± 4.5 ppb). Conversely, coastal air masses were characterised by
281 high concentrations of O₃ (53.6 ± 14.2 ppb) and low concentrations of NO_x (1.9 ± 1.6 ppb).
282 The high concentrations of NO_x in the marine air masses are attributable to the emissions from
283 ocean-going container ships that passed the site approximately 8 km to the south. A haze event
284 occurred from 24 September to 2 October, due to a transition from marine to coastal air masses
285 under stagnant conditions. O₃ concentrations steadily increased during the first 5 days, peaked
286 on 29 September, and remained high until the end of the haze period (Fig. 1). Therefore, the
287 potential HCOOH formation mechanism was analysed separately for these three distinct
288 periods.

289 Ambient HCOOH concentrations significantly varied during the three periods. The average
290 HCOOH concentration in marine air masses was 191 ± 167 ppt; this was higher than those over
291 the remote ocean, due to local emission sources, but significantly lower than those in urban
292 environments (Table 1). In contrast, the ambient HCOOH concentrations in coastal air masses

293 were substantially higher, averaging 996 ± 433 ppt, comparable with other measurements at
294 rural or urban background sites. During the haze period, the concentrations of HCOOH
295 displayed a pattern similar to the concentrations of O_3 , with the daytime peak concentration
296 increasing from 674 to 2790 ppt. A pronounced diurnal variation in the concentration of
297 HCOOH was observed throughout the entire campaign, as illustrated in Fig. 2, consistent with
298 other studies (Millet et al., 2015; Yuan et al., 2015). HCOOH concentrations rapidly increased
299 after sunrise, peaking at approximately 1 pm (local time), and then quickly decreasing in the
300 late afternoon, due to the weaker sunlight and lower BLH than earlier in the day.

301 HCOOH is widely recognised as a secondary photochemical product. Table 2 presents the
302 Pearson correlation coefficients (r) between the concentration of HCOOH and those of other
303 air pollutants or other meteorological parameters during the three distinct periods. The
304 concentration of HNO_3 was strongly correlated with the concentration of HCOOH throughout
305 the entire field campaign, consistent with other studies (Bannan et al., 2017; Millet et al., 2015).
306 This finding suggests that HCOOH is predominantly generated through secondary
307 photochemical mechanisms at this site, as HNO_3 is a secondary photochemical product
308 resulting from the reaction between $\cdot OH$ and NO_2 . The positive linear relationship between the
309 concentrations of O_3 and HCOOH also implies the secondary source of HCOOH.

310 A previous laboratory study revealed that HCOOH can be produced by the photochemical
311 aging of aerosols (Malecha and Nizkorodov, 2016), which may be an important process in
312 ambient air. In the coastal air masses and haze period, there was a strong correlation between
313 the concentrations of HCOOH and PM, particularly between the concentrations of HCOOH
314 and PM_{10} . This was also observed by Paulot et al. (2011) and suggests that HCOOH may be
315 produced from PM. The Sa of PM_{10} was also highly correlated with the concentration of
316 HCOOH in both coastal air masses and haze periods, indicating that HCOOH is mainly
317 produced from reactions on aerosol surfaces. However, in the marine air masses, the
318 concentration of HCOOH was not related to aerosols due to the low particle concentrations in
319 such masses. To further explore the potential role of aerosol aging in HCOOH production, we
320 plotted the correlation of HCOOH concentrations with $Sa \times O_3$, $Sa \times NO_3^-$, and $Sa \times O_3 \times NO_3^-$
321 for the coastal air masses (Fig. 3). We found that the correlation coefficient significantly
322 increased when Sa was combined with the concentration of O_3 or NO_3^- or with the
323 concentrations of both species, compared with these three factors being considered separately.
324 This finding suggests that the HCOOH observed in the coastal air masses was not
325 predominantly derived from gas-phase O_3 oxidation of VOCs; rather, it was derived from
326 heterogeneous or condensed-phase reactions on aerosol surfaces. The results during the haze
327 period were similar. Therefore, photochemical aerosol aging may play a key role in HCOOH
328 production as the aging process involves the reactive uptake of oxidants onto particle surfaces.

329 3.2. Laboratory experiments

330 Figure 4a presents the results of a typical aerosol-filter irradiation experiment. Upon turning

331 on the light, HCOOH was instantaneously produced, indicating a rapid transfer from the
332 condensed-phase to the gas phase through photochemical reactions. Within 3 minutes, the
333 HCOOH concentration reached 11.1 ppb, but when the light was turned off, the HCOOH
334 concentration quickly returned to nearly background concentrations. This suggests that
335 HCOOH was produced predominantly via photochemical reactions. The HCOOH
336 concentration exhibited a logarithmic decay after its first peak concentration and this decay
337 also occurred continued after its second peak concentration, which may be attributable to either
338 the evaporative loss of HCOOH or the photochemical loss of oxidants (Ye et al., 2017). When
339 the AM 1.5 filter was added, the HCOOH concentration decreased by approximately 48.1%
340 within 5 min, and after the filter was removed, the HCOOH concentration returned to the
341 logarithmic decay line. This suggests that there was only minor evaporation of HCOOH from
342 the condensed-phase due to the increased temperature of aerosol surfaces under light irradiation.
343 However, the addition of the 300–800 nm filter reduced the HCOOH concentration by only
344 13.2%, indicating that the photochemical production of HCOOH primarily occurs at
345 wavelengths lower than 360 nm. Given the agreement between the wavelength at which NO_3^-
346 absorbs light (290–350 nm) and the wavelength of HCOOH production (< 360 nm), and the
347 high correlation between the ambient HCOOH concentration and the product of Sa density and
348 NO_3^- concentration (as shown in Fig. 3), we infer that $\cdot\text{OH}$ produced from NO_3^- photolysis
349 were the major oxidants in the particle phase and thus drove HCOOH production. The
350 production of HCOOH was also found to be dependent on O_3 , as the concentration of HCOOH
351 increased by 64.7% after the addition of 100 ppb of O_3 .

352 We next extrapolated the production rate of HCOOH observed in the chamber to ambient
353 conditions, using the method described in Section 2.3, to assess the role played by the
354 photochemical aging of aerosols in HCOOH production. Table 3 summarises the HCOOH
355 concentrations and production rates observed in the chamber experiments, and the normalised
356 HCOOH production rates in ambient air under light and light + O_3 conditions, respectively.
357 The average $P_{\text{HCOOH-nml}}$ without the addition of O_3 was determined to be 0.106 ppb h^{-1} . The
358 addition of 100 ppb of O_3 increased $P_{\text{HCOOH-nml}}$ by 0.079 ppb h^{-1} , indicating that the
359 heterogeneous reaction between O_3 and aerosols made a non-negligible contribution to HCOOH
360 production. To quantify the contribution of this particle-phase pathway, we established a
361 relationship between $P_{\text{HCOOH-nml}}$ and three parameters: $\text{PM}_{2.5}$ concentration (cPM), which
362 represents the reactant concentration; Sa, which represents the available reaction area; and
363 $j\text{NO}_2$, which represents the light intensity. After multiplying these three factors, we discovered
364 a strong linear correlation between $P_{\text{HCOOH-nml}}$ and $\text{cPM} \times \text{Sa} \times j\text{NO}_2$ (Fig. 4b). The intercept
365 was set to zero, as there should be no HCOOH production when $\text{cPM} \times \text{Sa} \times j\text{NO}_2$ is zero.
366 Based on the correlation, we derived an equation (Eq. (4)) for calculating $P_{\text{HCOOH-nml}}$ with
367 assumption that $P_{\text{HCOOH-nml}(\text{O}_3)}$ increased linearly with O_3 concentration. This equation was then
368 incorporated into a box model to assess the importance of this production pathway compared

369 to other sources (See Section 3.3).

$$370 P_{\text{HCOOH-nml}} = 0.0091x + 0.010_3 \times 0.0064x, \quad x = \text{cPM} \times \text{Sa} \times j\text{NO}_2 \quad (4)$$

371 Photolysis of particulate NO_3^- is an important source of $\cdot\text{OH}$ (Zellner et al., 1990; Mack
372 and Bolton, 1999). To investigate the potential production of HCOOH from this source, an
373 irradiation experiment was conducted on a solution (Fig. 4c). The concentration of HCOOH
374 increased linearly with time and did not reach a stable state after 90 min of illumination. This
375 differs from the aerosol experiments and might have been due to the continuous evaporation of
376 water from the solution caused by the heating effect of the light source, which would have
377 concentrated the solution. To determine the appropriate time to calculate P_{HCOOH} , we also
378 plotted the time series of HONO concentrations. This showed that the HONO concentration
379 stabilised after 1 h of irradiation, suggesting that NO_3^- photolysis also reached a steady state.
380 As $\cdot\text{OH}$ produced from NO_3^- photolysis were the only oxidants present in the system, the actual
381 HCOOH production rate at the initial HCHO concentration should have followed the same
382 trend as the HONO concentration. Therefore, we chose 1 h after turning on the light as the
383 appropriate time to quantify P_{HCOOH} , and found that at this time, P_{HCOOH} in the chamber was
384 21.9 ppt s^{-1} .

385 We also attempted to extrapolate the results to ambient air, similar to the aerosol filter
386 experiments. To do so, in addition to normalising Sa and light intensity, we needed to normalise
387 the HCHO and NO_3^- concentrations. The average concentration of gas-phase HCHO ($\text{HCHO}_{(\text{g})}$)
388 measured on 28 September was 2.35 $\mu\text{g cm}^{-3}$. By using a ratio of 0.03 between the
389 concentration of HCHO in the particle phase ($\text{HCHO}_{(\text{p})}$) and the concentration of HCHO in the
390 gas phase (Toda et al., 2014), the concentration of $\text{HCHO}_{(\text{p})}$ was calculated to be 0.07 $\mu\text{g cm}^{-3}$,
391 which is comparable to the concentrations that have been reported in previous studies (Klippel
392 and Warneck, 1980; Toda et al., 2014). Based on the aqueous volume of aerosol calculated by
393 the E-AIM model (0.02 $\mu\text{l m}^{-3}$), the HCHO mass concentration in the aqueous phase was found
394 to be 3.5 g L^{-1} . The NO_3^- concentration on the aerosol surface was determined to be 0.98 mol
395 L^{-1} . Therefore, the P_{HCOOH} in ambient air attributable to the aqueous oxidation of HCHO was
396 estimated to be 0.41 ppb h^{-1} , which is 285% higher than the P_{HCOOH} attributable to the
397 photochemical aging of ambient particles. This greater-than-100% contribution could be
398 attributable to the simplicity of the solution system. In ambient air, other oxidisable species,
399 such as halides (Peng et al., 2022; M. Xia et al., 2022) could also react with the $\cdot\text{OH}$ produced
400 from NO_3^- photolysis, which competes with HCHO. In summary, NO_3^- photolysis appears to
401 be a critical source of oxidants during the photochemical aging process of aerosols.

402 3.3. Box model simulation

403 We next use a box model (see Methods 2.4) to evaluate the production and loss of HCOOH
404 in four scenarios (Table 4) which include the default MCM mechanism (S1), modified with
405 gas-phase reactions following Yuan et al. (2015) (S2), further addition of the photochemical
406 aging source (S3), and further adding of a biogenic source (S4). Figure 5 presents a

407 comprehensive analysis of HCOOH budget of campaign-averaged diurnal profile. In the base
408 case (Default MCM), the highest net production rate of HCOOH ($P_{\text{HCOOH-net}}$) is 0.018 ppb h^{-1} ,
409 significantly lower than the observed average rate of increase in HCOOH concentrations of
410 0.095 ppb h^{-1} from 6 am to 1 pm. Despite an increase of $P_{\text{HCOOH-net}}$ to 0.031 ppb h^{-1} in modified
411 case, the modelled value still lower than the observed rate of change. In both cases, the reaction
412 of CH_2OO Criegee intermediate biradicals with H_2O was identified as the major source of
413 HCOOH, accounting for about 68% of the currently known sources. CH_2OO is formed from
414 seven excited biradicals that originate from the O_3 oxidation of various alkenes and unsaturated
415 compounds (Saunders et al., 2003). Among these, CH_2OOE is the largest contributor to the
416 production of CH_2OO (Fig. S5) and is generated by the ozonolysis of isoprene.

417 Incorporating the photochemical production of HCOOH from particles into the F0AM
418 model (S3) resulted in substantial improvements in predictions. The peak $P_{\text{HCOOH-net}}$ increased
419 to 0.073 ppb h^{-1} , which is more than double that of the modified case. Among the secondary
420 production mechanisms considered, the production of HCOOH from particles was found to be
421 the largest source, accounting for 52.1% of the secondary production (Fig. 5c). The biogenic
422 emissions (S4) also played an important role, contributing to 34.4% of the total production (Fig.
423 5d). After considering all sources, including primary emissions and secondary productions, the
424 modelled $P_{\text{HCOOH-net}}$ (0.094 ppb h^{-1}) was much closer to the observed increase rate of HCOOH
425 (0.095 ppb h^{-1}).

426 Figure 6 presents the observed and modelled concentrations of HCOOH using different
427 mechanisms for the averaged diurnal profile throughout the whole campaign. It is evident that
428 the predicted HCOOH concentration increased substantially after incorporating the
429 productions from the photochemical aging of aerosols. However, the modelled HCOOH
430 concentration is still lower than the observed value in Scenario 4, where all sources are included.
431 The discrepancy may be explained by the inadequate treatment of physical processes in the box
432 model, such as deposition, convection, and advection. The primary loss of HCOOH is via
433 deposition, owing to its high solubility in water. To account for uncertainty in the deposition
434 velocity (V_d) of HCOOH, we conducted a sensitivity test of HCOOH production to various V_d
435 values in Scenario 4 (Fig. S6). The results revealed that a smaller deposition velocity results in
436 higher modelled HCOOH concentrations. The daytime peak concentration increased by about
437 20% when V_d decreased from 1.00 to 0.50 cm s^{-1} . The field-observed V_d of HCOOH ranges
438 from 0.43 cm s^{-1} to 1.10 cm s^{-1} (Müller et al., 2018), and thus given the high humidity at the
439 study site, the observed V_d of HCOOH of 0.5 cm s^{-1} should have been close to the lower limit.
440 Although the modelled concentration with $V_d = 0.50 \text{ cm s}^{-1}$ were similar to observed values,
441 the model failed to accurately predict the timing of the initial increase, peak, and sharp decrease
442 in HCOOH concentrations, indicating that other physical process, such as vertical mixing, also
443 influence the HCOOH concentration. Therefore, to account for the complex physical processes,
444 we employed a bimodal physical loss rate due to vertical dilution that varied with time of day

445 (1/21,600 s⁻¹ in daytime and a much smaller value of 1/518,400 s⁻¹ at night), as suggested by
446 Yuan et al. (2015). It is clear that the model performed better in predicting the diurnal pattern
447 using the bimodal physical loss rate compared to a constant value (Fig. 6).

448 After applying a bimodal physical loss rate, the model also predicts better in the continuous
449 simulation of two weeks (Fig. S7), except for 30 September and 1 October. The observed
450 HCOOH concentrations on these two days were significantly lower than the modelled values
451 due to a lower nitrate proportion (13.5%) on 30 September compared to other days during the
452 model period (22.2%). Therefore, the simplified parameterization using PM_{2.5} may
453 overestimate the production of HCOOH from photochemical aging of aerosols in areas with
454 high PM concentrations but a low nitrate proportion. An improved parameterization using the
455 concentrations of nitrate and organics should be developed in future studies.

456 To evaluate the role of aerosol photochemical aging on HCOOH production in a broader
457 context, we also examined the HCOOH data over the remote marine boundary layer obtained
458 from the Atmospheric Tomography Mission (ATom) aircraft campaign which was conducted
459 around the globe during April-May 2018 (Wofsy et al., 2021) (Text S5). Our results show that
460 the photochemical aging of aerosols was insignificant in remote ocean areas due to the low PM
461 and nitrate concentrations found there. However, when these regions are affected by aged fire
462 plumes containing higher levels of organics and nitrate, photochemical aging of aerosols
463 accounts for the substantial sources of HCOOH. These results suggest the photochemical aging
464 appears to be important in relatively polluted atmospheres.

465 4. Conclusion and implications

466 This study **demonstrates** high rates of HCOOH production from the photochemical aging
467 of real ambient particles and the potential importance of this pathway in the formation of
468 HCOOH under ambient conditions at a coastal site in Hong Kong. Incorporating aerosol
469 photochemical aging significantly improved the performance of a widely used chemical model,
470 which underscores the significance of condensed-phase photochemistry and the necessity of
471 incorporating its mechanisms into atmospheric models. The substantial production of HCOOH
472 from condensed-phase photochemical reactions altered both the composition and the volatility
473 of SOA. Moreover, other low-molecular-weight organic acids, such as acetic acid, may be
474 produced via this mechanism, as observed in our irradiation experiments. Improving the
475 constraints on this photochemical aging of aerosols will not only aid the understanding of the
476 budget of these organic acids but also affect their SOA chemistry.

477 Our solution irradiation experiments demonstrated the **crucial role** of NO₃⁻ photolysis in
478 HCOOH production via the production of ·OH. This suggests that NO₃⁻ photolysis not only
479 influences the aerosol-based production of inorganic species (such as HONO (Ye et al., 2017)
480 and halogens (Peng et al., 2022; M. Xia et al., 2022)), but also the aerosol-based production of
481 water-soluble organics. **Therefore, future investigations into the involvement of NO₃⁻**

482 photodissociation in aerosol aging processes are necessary to enhance our understanding of the
483 underlying mechanisms in the photochemical aging of aerosols.

484 Data availability

485 The data that support the findings of this study are openly available in Zenodo at
486 <https://10.5281/zenodo.8415792>. Other raw data are also available from the corresponding
487 author, upon reasonable request.

488 Authors' contributions

489 T. W. arranged the field campaign and designed the laboratory irradiation experiment. Y. J. and
490 M. X. conducted the field campaign and photochemical filter experiments. M. X. revised the
491 model code. Y. J. conducted the photochemical solution experiments, analysed the data, ran the
492 model, and wrote the draft manuscript. T. W. and M. X. revised the manuscript.

493 Competing interests

494 One author (Tao Wang) is a member of the editorial board of Atmospheric Chemistry and
495 Physics. The authors have no other competing interests to declare.

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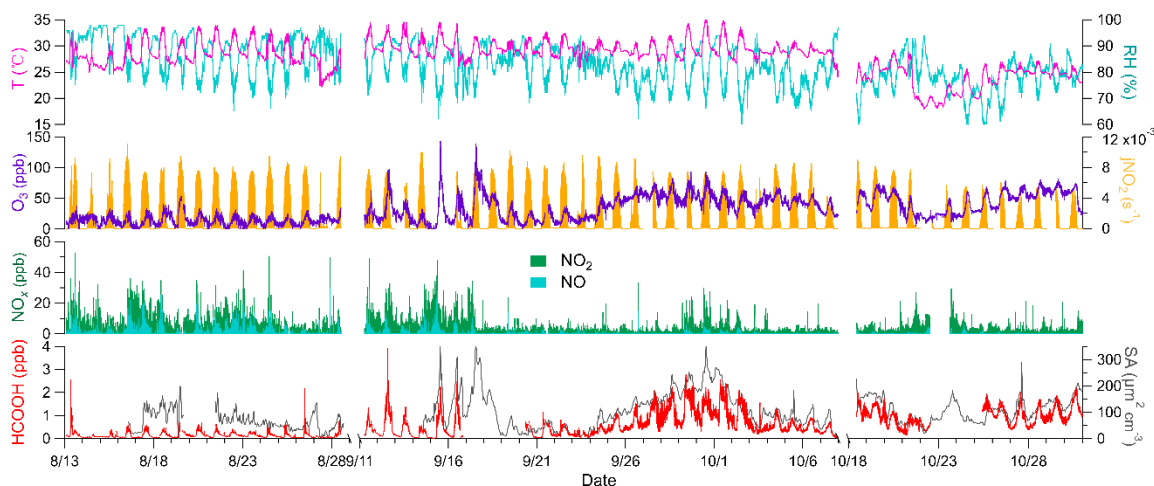
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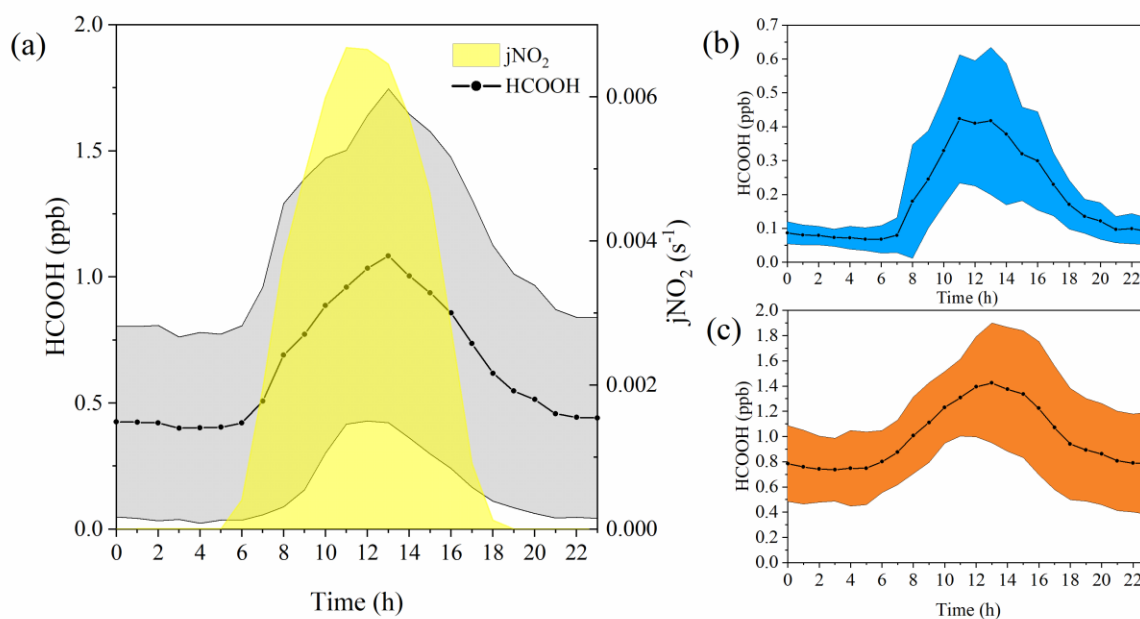
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788 **Fig. 1** Times series of temperature (T); relative humidity (RH); nitrogen dioxide photolysis
 789 frequency ($j\text{NO}_2$); concentrations of ozone (O_3), nitrogen oxides (NO_x), and formic acid
 790 (HCOOH); and surface area density (SA) during the field campaign. Data were not collected
 791 from August 29 to September 10 due to the persistently clean weather conditions, similar to
 792 those during the initial stage of the campaign. The gap in data collection from October 8 to
 793 October 17 was due to a rainstorm. Any other short gaps were caused by instrument
 794 maintenance.

795 **Table 1** Summary of worldwide field-observed formic acid (HCOOH) concentrations

Location	Type	Time	HCOOH (ppb)	Reference
Pasadena, USA	urban	2010.06-07	2.0 ± 1.0	(Yuan et al., 2015)
Kensington, London	urban background	2012.01-02 2012.07-08	0.63 (winter) 1.33 (summer)	(Bannan et al., 2017)
Shanghai, China	suburban	2017.06.18-12.23	2.08 ± 1.89	(Xu et al., 2020)
Yorkville, USA	rural	2016.08.15-10.13	1.17 ± 0.85	(Nah et al., 2018)
North Pacific Pacific and Atlantic	marine and marine	2008.07.29-08.19 2017.09-10 2018.04-05	30 ± 39.8 ppt < 0.1	(Miyazaki et al., 2014) (Chen et al., 2021)
Colorado, USA	forest	2016.02.01-03.01 2016.04.15-05.15 2016.07.15-08.15 2016.10.01-11.01	55 ± 57 ppt (winter) 30 ± 24 ppt (spring) 1.2 ± 0.91 (summer) 0.81 ± 0.48 (autumn)	(Fulgham et al., 2019)
Alabama, USA	deciduous forest	2013.06-07	2.5 (peak average daytime)	(Millet et al., 2015)
Hong Kong, China	coastal	2021.08.13-10.31	0.58 ± 0.53	This study

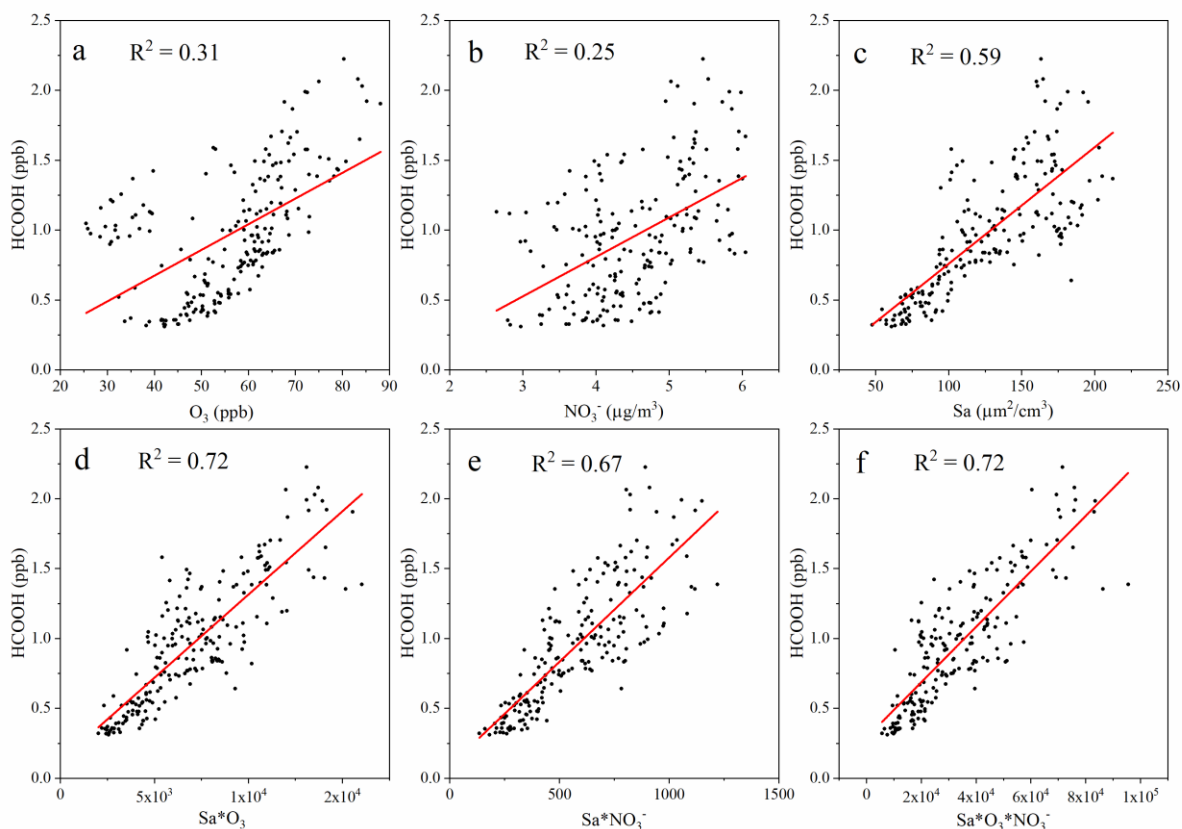
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797
 798 **Fig. 2** Diurnal variation in the concentrations of formic acid (HCOOH) in different periods. (a)
 799 Diurnal cycle of HCOOH concentrations and frequency of nitrogen dioxide photolysis ($j\text{NO}_2$)
 800 for the whole campaign; (b) and (c) diurnal cycle of HCOOH concentrations in marine and
 801 coastal air masses, respectively. The shading represents the standard deviations of the
 802 measurements.

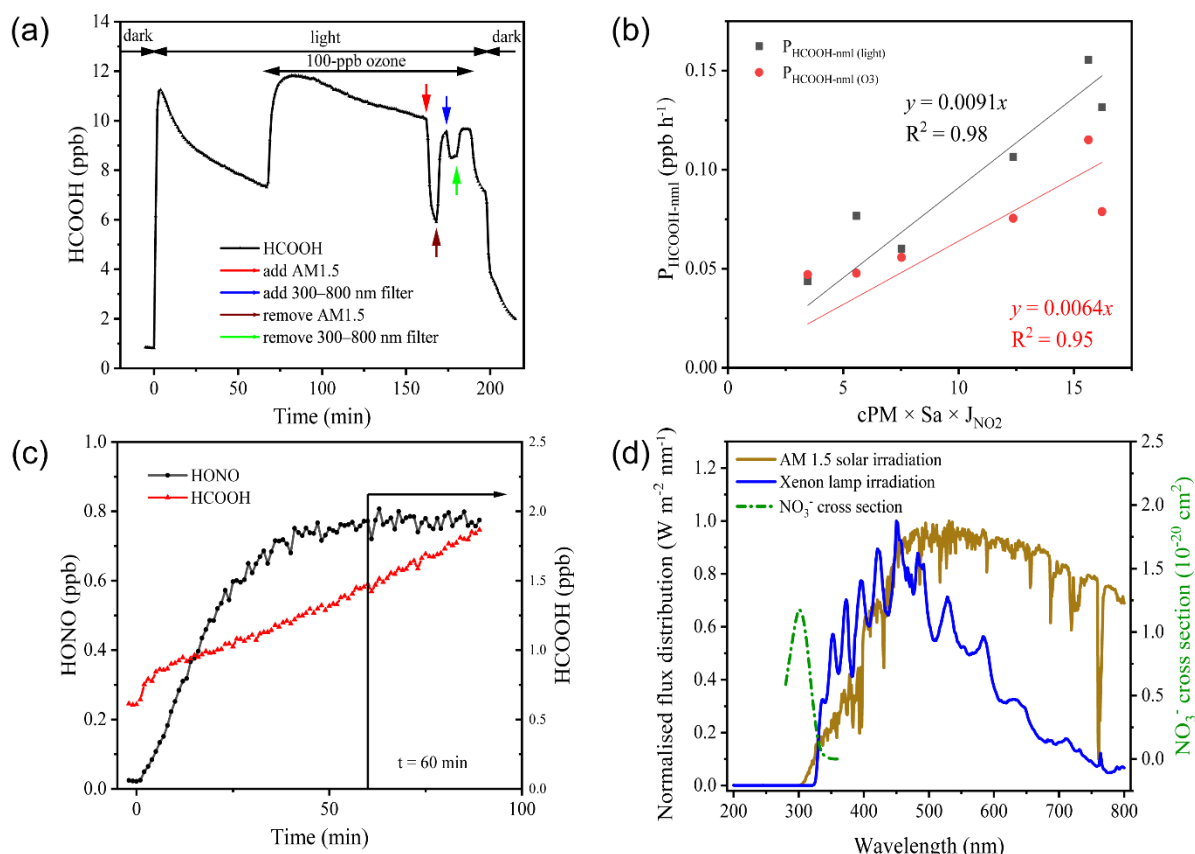
803 **Table 2** Pearson correlation coefficient (r) matrix between the concentration of formic acid and
 804 other air pollutants, and related meteorological parameters, during three distinct periods.

Parameter	Coastal	Haze	Marine	Parameter	Coastal	Haze	Marine
$j\text{NO}_2$	0.41	0.58	0.65	Sa	0.73	0.68	-0.03
T	-0.27	0.70	0.72	Sa \times NO_3^-	0.85	0.56	0.15
RH	-0.56	-0.51	-0.65	Sa \times O_3	0.83	0.74	0.31
PM_1	0.79	0.66	0.05	HNO_3	0.75	0.59	0.69
$\text{PM}_{2.5}$	0.69	0.63	0.19	Cl^-	-0.41	-0.44	0.09
PM_{10}	0.68	0.55	0.26	NO_3^-	0.67	-0.10	0.57
HONO	-0.03	0.26	-0.34	SO_4^{2-}	0.66	0.65	0.10
CH_3COOH	0.89	0.88	-0.27	Na^+	-0.28	-0.50	0.37
NO	-0.12	0.44	0.13	NH_4^+	0.72	0.64	0.24
NO_2	-0.24	0.36	-0.39	K^+	0.53	0.32	0.15
NO_x	-0.22	0.40	-0.27	Mg^{2+}	-0.30	-0.38	0.47
O_3	0.69	0.65	0.68	Ca^{2+}	-0.11	0.09	0.04
SO_2	0.64	0.66	0.41	HCl	0.18	0.51	0.55
CO	0.63	0.51	0.13	isoprene	0.03	0.61	0.63
NH_3	0.37	0.46	0.16	benzene	0.63	0.55	0.05



806

807 **Fig. 3** Scatter plot of the concentration of formic acid (HCOOH) and (a) the concentration of
 808 ozone (O_3); (b) the mass concentration of nitrate ions (NO_3^-) in $PM_{2.5}$; (c) the surface
 809 density (Sa) of PM_1 ($\mu m^2 cm^{-3}$); (d) the product of Sa and the concentration of O_3 ; (e) the
 810 product of Sa and the concentration of NO_3^- ; and (f) the product of Sa, the concentration of O_3 ,
 811 and the concentration of NO_3^- in coastal air masses.



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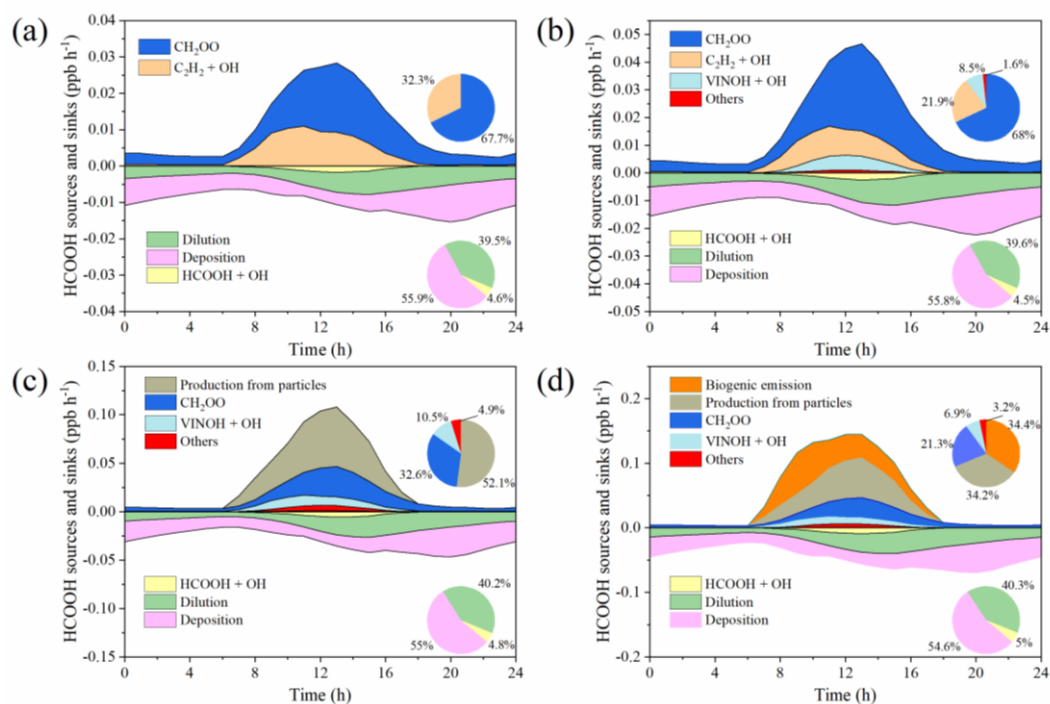
813 **Fig. 4** Results of the irradiation experiments. (a) Typical variation in formic acid (HCOOH)
 814 concentrations during irradiation (in aerosols collected on 2 November 2020). AM 1.5 and
 815 300–800 nm filters were added after the addition of 100-ppb ozone. (b) The correlation between
 816 $P_{\text{HCOOH-nml}}$ and 2.5- μm particulate matter ($\text{PM}_{2.5}$) concentration (cPM) \times surface area (Sa) \times
 817 nitrogen dioxide photolytic frequency (j_{NO_2}). (c) Variations in the concentrations of HCOOH
 818 and nitrous acid (HONO) as a function of time after illumination of an aqueous solution of
 819 formaldehyde and sodium nitrite at $\text{pH} = 2.7$. The vertical black line indicates the time at which
 820 HONO stabilised. (d) Comparison of the irradiation spectrum of the xenon lamp used in this
 821 study and standard air mass 1.5 solar irradiation (AM 1.5).

822 **Table 3** Summary of formic acid (HCOOH) concentrations and production rates observed in
 823 chamber experiments and normalised HCOOH production rates in ambient air under light and
 824 light + ozone (O_3) conditions, respectively. $\text{HCOOH}_{(\text{O}_3)}$ denotes the increased concentration of
 825 HCOOH after the addition of 100 ppb O_3 .

Date	$\text{HCOOH}_{(\text{light})}$ (ppt)	$\text{HCOOH}_{(\text{O}_3)}$ (ppt)	$P_{\text{HCOOH}(\text{light})}$ (ppt s ⁻¹)	$P_{\text{HCOOH}(\text{O}_3)}$ (ppt s ⁻¹)	$P_{\text{HCOOH-nml}}$ nml (light) (ppb h ⁻¹)	$P_{\text{HCOOH-nml}}$ nml (O ₃) (ppb h ⁻¹)
2020.10.07	8420.2	4670.0	299.4	166.0	1.70E-01	1.33E-01
2020.10.08	6787.7	2899.0	241.3	103.1	1.31E-01	7.89E-02

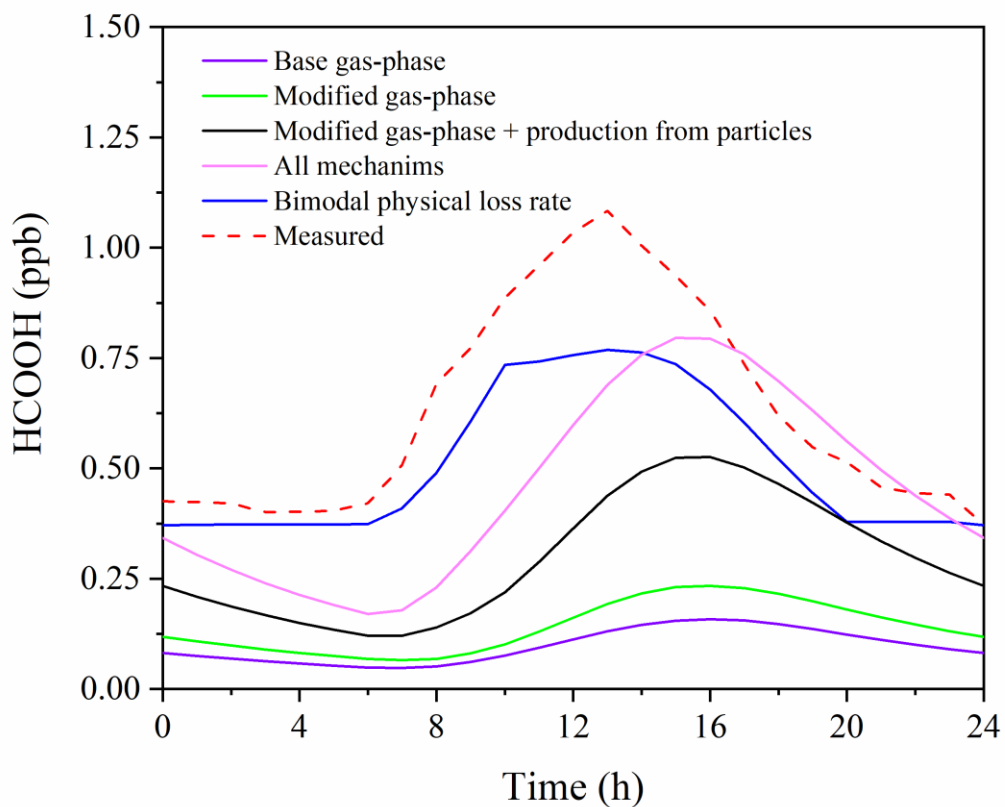
2020.10.26	4660.9	3077.5	165.7	109.4	6.01E-02	5.57E-02
2020.11.02	6656.3	3507.6	236.7	124.7	1.55E-01	1.15E-01
2020.11.03	4490.8	2266.7	159.7	80.6	1.06E-01	7.55E-02
2020.11.04	4943.1	2191.5	175.8	77.9	7.67E-02	4.77E-02
2020.11.05	3088.0	2368.6	109.8	84.2	4.37E-02	4.71E-02
Average	5578.1	2997.3	198.3	106.6	1.06E-01	7.89E-02

826



827

828 **Fig. 5** Model-calculated profiles of sources and sinks of formic acid (HCOOH) on **averaged**
829 **diurnal profile during the whole campaign in four scenarios described in Table 4.** Upper right
830 inset: the contribution from various sources to HCOOH concentrations. Bottom right inset: the
831 contribution from different sinks to HCOOH concentrations. CH₂OO = formaldehyde oxide, a
832 Criegee intermediate (biradical); VINOH = vinyl alcohol.



833

834 **Fig. 6** Comparison of measured and modelled diurnal profiles of HCOOH during the whole
 835 campaign.

836 **Table 4** The mechanisms included in different model scenarios.

scenario	Default MCM	Modified MCM	Particle-phase pathway	Biogenic emissions
1	√			
2		√		
3		√	√	
4		√	√	√

837