

1 **Trends in secondary inorganic aerosol pollution in China and its responses to**
2 **emission controls of precursors in wintertime**

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33 **ABSTRACT:** The Chinese government recently proposed ammonia (NH₃) emissions
34 reductions (but without a specific national target) as a strategic option to mitigate PM_{2.5}
35 pollution. We combined a meta-analysis of nationwide measurements and air quality
36 modelling to identify efficiency gains by striking a balance between controlling NH₃
37 and acid gas (SO₂ and NO_x) emissions. We found that PM_{2.5} concentrations decreased
38 from 2000 to 2019, but annual mean PM_{2.5} concentrations still exceeded 35 μg m⁻³ at
39 74% of 1498 monitoring sites in 2015-2019. Secondary inorganic aerosols (SIA) were
40 the dominant contributor to ambient PM_{2.5} concentrations. The concentration of PM_{2.5}
41 and its components were significantly higher (16%-195%) on hazy days than on non-
42 hazy days. Compared with mean values of other components, this difference was more
43 significant for the secondary inorganic ions SO₄²⁻, NO₃⁻, and NH₄⁺ (average increase
44 98%). While sulfate concentrations significantly decreased over the time period, no
45 significant change was observed for nitrate and ammonium concentrations. Model
46 simulations indicate that the effectiveness of a 50% NH₃ emission reduction for
47 controlling SIA concentrations decreased from 2010 to 2017 in four megacity clusters
48 of eastern China, simulated for the month of January under fixed meteorological
49 conditions (2010). Although the effectiveness further declined in 2020 for simulations
50 including the natural experiment of substantial reductions in acid gas emissions during
51 the CoVID-19 pandemic, the resulting reductions in SIA concentrations were on
52 average 20.8% lower than that in 2017. In addition, the reduction of SIA concentrations
53 in 2017 was greater for 50% acid gas reductions than for the 50% NH₃ emissions
54 reduction. Our findings indicate that persistent secondary inorganic aerosol pollution in
55 China is limited by acid gases emissions, while an additional control on NH₃ emissions
56 would become more important as reductions of SO₂ and NO_x emissions progress.

57

58 **Keywords:** Air pollution, Particulate matter, Second inorganic aerosols, Anthropogenic
59 emission, Ammonia.

60

61 **1. Introduction**

62 Over the past two decades, China has experienced severe PM_{2.5} (particulate matter
63 with aerodynamic diameter $\leq 2.5 \mu\text{m}$) pollution (Huang et al., 2014; Wang et al., 2016),
64 leading to adverse impacts on human health (Liang et al., 2020) and the environment
65 (Yue et al., 2020). In 2019, elevated PM_{2.5} concentrations accounted for 46% of polluted
66 days in China and PM_{2.5} was officially identified as a key year-round air pollutant
67 (MEEP, 2019). Mitigation of PM_{2.5} pollution is therefore the most pressing current
68 challenge to improve China's air quality.

69 The Chinese government has put a major focus on particulate air pollution control
70 through a series of policies, regulations, and laws to prevent and control severe air
71 pollution. Before 2010, the Chinese government mainly focused on controlling SO₂
72 emissions via improvement of energy efficiency, with less attention paid to NO_x
73 abatement (CSC, 2007, 2011, 2016). For example, the 11th Five-Year Plan (FYP) (2006-
74 2010) set a binding goal of a 10% reduction for SO₂ emission (CSC, 2007). The 12th
75 FYP (2011-2015) added NO_x regulation and required 8% and 10% reductions for SO₂
76 and NO_x emissions, respectively (CSC, 2011) This was followed by further reductions
77 in SO₂ and NO_x emissions of 15% and 10%, respectively, in the 13th FYP (2016-2020)
78 (CSC, 2016). In response to the severe haze events of 2013, the Chinese State Council
79 promulgated the toughest-ever 'Atmospheric Pollution Prevention and Control Action
80 Plan' in September 2013, aiming to reduce ambient PM_{2.5} concentrations by 15-20% in
81 2017 relative to 2013 levels in metropolitan regions (CSC, 2013). As a result of the

82 implementation of stringent control measures, emissions reductions markedly
83 accelerated from 2013-2017, with decreases of 59% for SO₂, 21% for NO_x, and 33%
84 for primary PM_{2.5} (Zheng et al., 2018). Consequently, significant reductions in annual
85 mean PM_{2.5} concentrations were observed nationwide (Zhang et al., 2019; Yue et al.,
86 2020), in the range 28-40% in the metropolitan regions (CSC, 2018a). To continue its
87 efforts in tackling air pollution, China promulgated the Three-Year Action Plan (TYAP)
88 in 2018 for Winning the Blue-Sky Defense Battle (CSC, 2018b), which required a
89 further 15% reduction in NO_x emissions by 2020 compared to 2018 levels.

90 Despite a substantial reduction in PM_{2.5} concentrations in China, the proportion of
91 secondary aerosols during severe haze periods is increasing (An et al., 2019), and can
92 comprise up to 70% of PM_{2.5} concentrations (Huang et al., 2014). Secondary inorganic
93 aerosols (SIA, the sum of sulfate (SO₄²⁻), nitrate (NO₃⁻), and ammonium (NH₄⁺)) were
94 found to be of equal importance to secondary organic aerosols, with 40-50%
95 contributions to PM_{2.5} in eastern China (Huang et al., 2014; Yang et al., 2011). The acid
96 gases (i.e., NO_x, SO₂), together with NH₃, are crucial precursors of SIA via chemical
97 reactions that form particulate ammonium sulfate, ammonium bisulfate, and
98 ammonium nitrate (Ianniello et al., 2010). In addition to the adverse impacts on human
99 health via fine particulate matter formation (Liang et al., 2020; Kuerban et al., 2020),
100 large amounts of NH₃ and its aerosol-phase products also lead to nitrogen deposition
101 and consequently to environmental degradation (Ortiz-Montalvo et al., 2014; Zhan et
102 al., 2021).

103 Following the successful controls on NO_x and SO₂ emissions, attention is turning
104 to NH₃ emissions as a possible means of further PM_{2.5} control (Bai et al., 2019; Kang
105 et al., 2016), particularly as emissions of NH₃ increased between the 1980s and 2010s.
106 Some studies have found that NH₃ limited the formation of SIA in winter in the eastern

107 United States (Pinder et al., 2007) and Europe (Megaritis et al., 2013). Controls on NH₃
108 emissions have been proposed in the TYAP, although mandatory measures and binding
109 targets have not yet been set (CSC, 2018b). Nevertheless, this proposal means that
110 China will enter a new phase of PM_{2.5} mitigation, with attention now given to both acid
111 gas and NH₃ emissions. However, in the context of effective control of PM_{2.5} pollution
112 via its SIA component, two key questions arise: 1) what are the responses of the
113 constituents of SIA to implementation of air pollution control policies, and 2) what is
114 the relative efficiency of NH₃ versus acid gas emission controls to reduce SIA pollution?

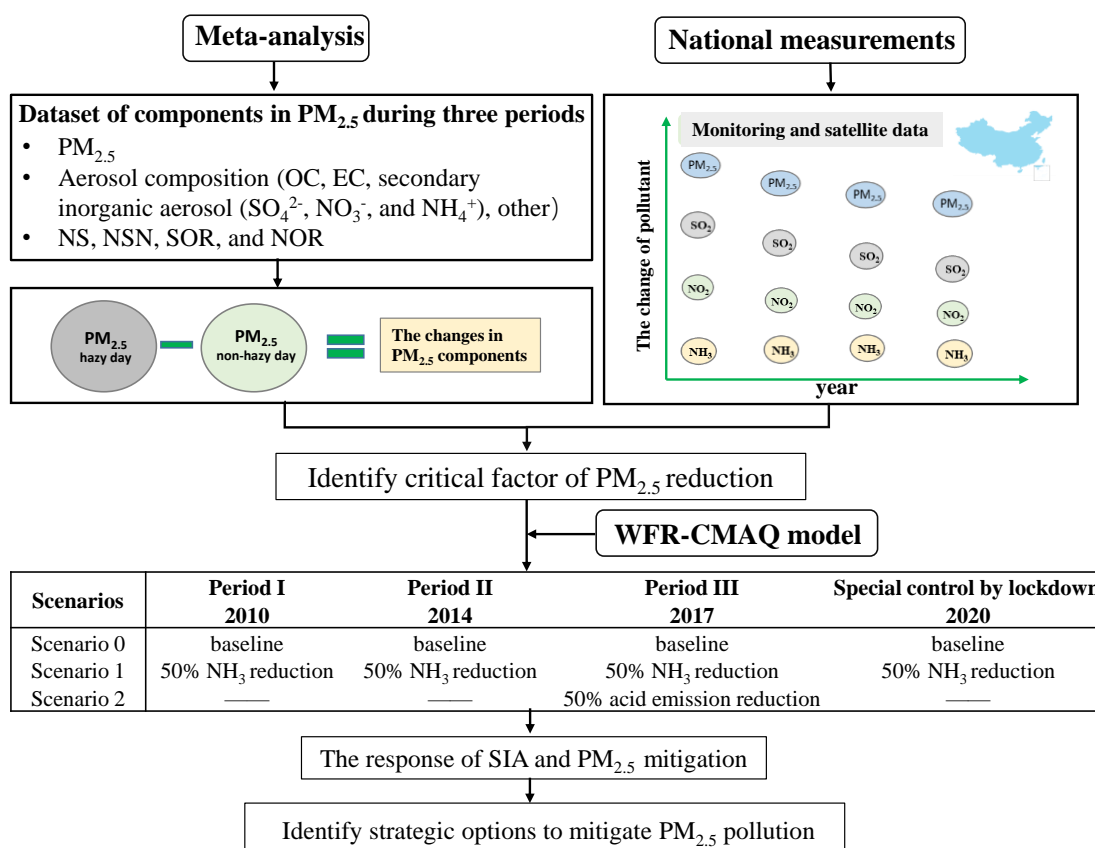
115 To fill this evidence gap and provide useful insights for policy-making to improve
116 air quality in China, this study adopts an integrated assessment framework. With respect
117 to the emission control policy summarized above, China's PM_{2.5} control can be divided
118 into three periods: period I (2000–2012), in which PM_{2.5} was not the targeted pollutant;
119 period II (2013–2016), the early stage of targeted PM_{2.5} control policy implementation;
120 and period III (2017–2019), the latter stage with more stringent policies. Therefore, our
121 research framework consists of two parts: (1) assessment of trends in annual mean
122 concentrations of PM_{2.5}, its chemical components and SIA gaseous precursors from
123 meta-analyses and observations; (2) quantification of SIA responses to emissions
124 reductions in NH₃ and acid gases using the Weather Research and Forecasting and
125 Community Multiscale Air Quality (WRF/CMAQ) models.

126 **2. Materials and methods**

127 **2.1. Research framework**

128 This study developed an integrated assessment framework to analysis the trends of
129 secondary inorganic aerosol and strategic options to reduce SIA and PM_{2.5} pollution in
130 China (Fig. 1). The difference in PM_{2.5} chemical components between hazy and non-
131 hazy days was first assessed by meta-analysis of published studies. These were

132 interpreted in conjunction with the trends in air concentrations of PM_{2.5} and its
 133 secondary inorganic aerosol precursors (SO₂, NO₂, and NH₃) derived from surface
 134 measurements and satellite observations. The potential of SIA and PM_{2.5} concentration
 135 reductions from precursor emission reductions was then evaluated using the Weather
 136 Research and Forecasting and Community Multiscale Air Quality (WRF/CMAQ)
 137 models.



138

139 **Fig. 1.** Integrated assessment framework for Chinese PM_{2.5} mitigation strategic options.

140 OC is organic carbon, EC is elemental carbon, NO₃⁻ is nitrate, SO₄²⁻ is sulfate, and NH₄⁺
 141 is ammonium. NS is the slope of the regression equation between [NH₄⁺] and [SO₄²⁻],
 142 NSN is the slope of the regression equation between [NH₄⁺] and [SO₄²⁻ + NO₃⁻], SOR
 143 is sulfur oxidation ratio, and NOR is nitrogen oxidation ratio. SIA is Secondary
 144 inorganic aerosols. WRF-CMAQ is Weather Research and Forecasting and Community
 145 Multiscale Air Quality models.

2.2. Meta-analysis of PM_{2.5} and its chemical components

Meta-analyses can be used to quantify the differences in concentrations of PM_{2.5} and its secondary inorganic aerosol components (NH₄⁺, NO₃⁻, and SO₄²⁻) between hazy and non-hazy days and to identify the major pollutants on non-hazy days (Wang et al., 2019b); this provides evidence for effective options on control of precursor emissions (NH₃, NO₂, and SO₂) for reducing occurrences of hazy days.

To build a database of atmospheric concentrations of PM_{2.5} and chemical components between hazy and non-hazy days, we conducted a literature survey using the Web of Science and the China National Knowledge Infrastructure for papers published between January 2000 and January 2020. The keywords included: (1) "particulate matter," or "aerosol," or "PM_{2.5}" and (2) "China" or "Chinese". Studies were selected based on the following conditions:

(1) Measurements were taken on both hazy and non-hazy days.

(2) PM_{2.5} chemical components were reported.

(3) If hazy days were not defined in the screened articles, the days with PM_{2.5} concentrations > 75 µg m⁻³ (the Chinese Ambient Air Quality Standard Grade II for PM_{2.5} (CSC, 2012)) were treated as hazy days.

(4) If an article reported measurements from different monitoring sites in the same city, e.g. Mao et al. (2018) and Xu et al. (2019), then each measurement was considered an independent study.

(5) If there were measurements in the same city for the same year, e.g. Tao et al. (2016) and Han et al. (2017), then each measurement was treated as an independent study.

One hundred articles were selected based on the above conditions with the lists provided in the Supporting Material dataset. For each selected study, we documented the study sites, study periods, seasons, aerosol types, and aerosol species mass concentrations (in µg m⁻³) over the entire study period (2000–2019) (the detailed data

171 are provided in the dataset). In total, the number of sites contributing data to the meta-
172 analysis was 267 and their locations are shown in Fig. S1. If relevant data were not
173 directly presented in studies, a GetData Graph Digitizer (Version 2.25,
174 <http://www.getdatagraph-digitizer.com>) was used to digitize concentrations of PM_{2.5}
175 chemical components from figures. The derivations of other variables such as sulfur
176 and nitrogen oxidation ratios are described in Supplementary Information Method 1.

177 Effect sizes were developed to normalize the combined studies' outcomes to the
178 same scale. This was done through the use of log response ratios (lnRR) (Nakagawa et
179 al., 2012; Ying et al., 2019). The variations in aerosol species were evaluated as follows:

$$180 \ln RR = \ln \left(\frac{X_p}{X_n} \right) \quad (1)$$

181 where X_p and X_n represent the mean values of the studied variables of PM_{2.5} components
182 on hazy and non-hazy days, respectively. The mean response ratio was then estimated
183 as:

$$184 RR = \exp \left[\frac{\sum \ln RR(i) \times W(i)}{\sum W(i)} \right] \quad (2)$$

185 where $W(i)$ is the weight given to that observation as described below. Finally, variable-
186 related effects were expressed as percent changes, calculated as $(RR-1) \times 100\%$. A 95%
187 confidence interval not overlapping with zero indicates that the difference is significant.
188 A positive or negative percentage value indicates an increase or decrease in the response
189 variables, respectively.

190 We used inverse sampling variances to weight the observed effect size (RR) in the
191 meta-analysis (Benitez-Lopez et al., 2017). For the measurement sites where standard
192 deviations (SD) or standard errors (SE) were absent in the original study reports, we
193 used the "Bracken, 1992" approach to estimate SD (Bracken et al., 1992). The variation-
194 related chemical composition of PM_{2.5} was assessed by random effects in meta-analysis.
195 Rosenberg's fail safe-numbers (N_{fs}) were calculated to assess the robustness of findings

196 on PM_{2.5} to publication bias (Ying et al., 2019) (See Table S1). The results (effects)
197 were considered robust despite the possibility of publication bias if $N_{fs} > 5 \times n + 10$,
198 where n indicates the number of sites. The statistical analysis of the concentrations of
199 PM_{2.5} and secondary inorganic ions for three periods used a non-parametric statistical
200 method since concentrations were not normally distributed based on the Kruskal-Wallis
201 test (Kruskal and Walls, 1952). For each species, the Kruskal-Wallis one-way analysis
202 of variance (ANOVA) on ranks among three periods was performed with pairwise
203 comparison using Dunn’s method (Dunn, 1964).

204 **2.3. Data collection of air pollutant concentrations**

205 To assess the recent annual trends in China of PM_{2.5} and of the SO₂ and NO₂
206 gaseous precursors to SIA, real-time monitoring data of these pollutants at 1498
207 monitoring stations in 367 cities during 2015–2019 were obtained from the China
208 National Environmental Monitoring Center (CNEMC) (<http://106.37.208.233:20035/>).
209 This is an open-access archive of air pollutant measurements from all prefecture-level
210 cities since January 2015. Successful use of data from CNEMC to determine
211 characteristics of air pollution and related health risks in China has been demonstrated
212 previously (Liu et al., 2016; Kuerban et al., 2020). The geography stations are shown
213 in Fig. S1. The annual mean concentrations of the three pollutants at all sites were
214 calculated from the hourly time-series data according to the method of Kuerban et al.
215 (2020). Information about sampling instruments, sampling methods, and data quality
216 controls for PM_{2.5}, SO₂, and NO₂ is provided in Supplementary Method 2. Surface NH₃
217 concentrations over China for the 2008–2016 (the currently available) were extracted
218 from the study of Liu et al. (2019). Further details are in Supplementary Method 2.

219 **2.4. WRF/CMAQ model simulations**

220 The Weather Research and Forecasting model (WRFv3.8) and the Models-3

221 community multi-scale air quality (CMAQv5.2) model were used to evaluate the
222 impacts of emission reductions on SIA and PM_{2.5} concentrations over China. The
223 simulations were conducted at a horizontal resolution of 12 km × 12 km. The simulation
224 domain covered the whole of China, part of India and east Asia. In the current study,
225 focus was on the following four regions in China: Beijing-Tianjin-Hebei (BTH),
226 Yangtze River Delta (YRD), Pearl River Delta (PRD), and Sichuan Basin (SCB). The
227 model configurations used in this study were the same as those used in Wu et al. (2018a)
228 and are briefly described here. The WRFv3.8 model was applied to generate
229 meteorological inputs for the CMAQ model using the National Center for
230 Environmental Prediction Final Operational Global Analysis (NCEP-FNL) dataset
231 (Morrison et al., 2009). Default initial and boundary conditions were used in the
232 simulations. The carbon-bond (CB05) gas-phase chemical mechanism and AERO6
233 aerosol module were selected in the CMAQ configuration (Guenther et al., 2012).
234 Anthropogenic emissions for 2010, 2014 and 2017 were obtained from the Multi-
235 resolution Emission Inventory (<http://meicmodel.org>) with 0.25° × 0.25° spatial
236 resolution and aggregated to 12km×12km resolution (Zheng et al., 2018; Li et al., 2017).
237 Each simulation was spun-up for six days in advance to eliminate the effects of the
238 initial conditions.

239 The years 2010, 2014 and 2017 were chosen to represent the anthropogenic
240 emissions associated with the periods I, II, III, respectively. January was selected as the
241 typical simulation month because wintertime haze pollution frequently occurs in this
242 month (Wang et al., 2011; Liu et al., 2019b). The sensitivity scenarios of emissions in
243 January can therefore help to identify the efficient option to control haze pollution.

244 The Chinese government has put a major focus on acid gas emission control
245 through a series of policies in the past three periods (Fig S2). The ratio decreases of

246 anthropogenic emissions SO₂ and NO_x in January for the years 2010, 2014, 2017 and
247 2020 are presented in SI [Tables S2 and S3](#), respectively. The emissions from
248 surrounding countries were obtained from the Emissions Database for Global
249 Atmospheric Research (EDGAR): HTAPV2. The scenarios and the associated
250 reductions of NH₃, NO_x and SO₂ for selected four years in three periods can be found
251 in [Fig. 1](#).

252 The sensitivities of SIA and PM_{2.5} to NH₃ emissions reductions were determined
253 from the average PM_{2.5} concentrations in model simulations without and with an
254 additional 50% NH₃ emissions reduction. The choice [of 50% additional NH₃ emissions](#)
255 [reduction is based on the feasibility and current upper bound of NH₃ emissions](#)
256 [reduction expected to be realized in the near future \(Liu et al., 2019a; Table S4\).](#) Zhang
257 [et al. \(2020\) found that the mitigation potential of NH₃ emissions from cropland](#)
258 [production and livestock production in China can reach up to 52% and 58%,](#)
259 [respectively.](#) To eliminate the influences of varying meteorological conditions, all
260 simulations were conducted under the fixed meteorological conditions of 2010.

261 During the COVID-19 lockdown in China, emissions of primary pollutants were
262 subject to unprecedented reductions due to national restrictions on traffic and industry;
263 in particular, emissions of NO_x and SO₂ reduced by 46% and 24%, respectively,
264 averaged across all Chinese provinces ([Huang et al., 2021](#)). We therefore also ran
265 simulations applying the same reductions in NO_x and SO₂ (based on 2017 MEIC) that
266 were actually observed during the COVID-19 lockdown as a case of special control in
267 2020.

268 [2.5 Model performance](#)

269 The CMAQ model has been extensively used in air quality studies ([Zhang et al.,](#)
270 [2019; Backes et al., 2016](#)) [and the validity of the chemical regime in the CMAQ model](#)

271 [had been confirmed by our previous studies \(Zhang et al., 2021a; Wang et al., 2020a,](#)
272 [2021a\). In this study, we used surface measurements from previous publications \(e.g.,](#)
273 [\(Xiao et al., 2020, 2021; Geng et al., 2019; Xue et al., 2019\) and satellite observations](#)
274 [to validate the modelling meteorological parameters by WRF model and air](#)
275 [concentrations of PM_{2.5} and associated chemical components by CMAQ model. The](#)
276 [meteorological measurements used for validating the WRF model performances were](#)
277 [obtained from the National Climate Data Center \(NCDC\)](#)
278 [\(ftp://ftp.ncdc.noaa.gov/pub/data/noaa/\). For validation of the CMAQ model, monthly](#)
279 [mean concentrations of PM_{2.5} were obtained from Tracking Air pollution in China \(TAP,](#)
280 [http://tapdata.org.cn/\) database. We also collected ground-based observations from](#)
281 [previous publications to validate the modeling concentrations of SO₄²⁻, NO₃⁻, and NH₄⁺.](#)
282 [The detailed information of the monitoring sites are presented in Table S5. Further](#)
283 [information about the modelling is given in Supplementary Method 3 and Figs. S3-S7](#)
284 [and Table S5.](#)

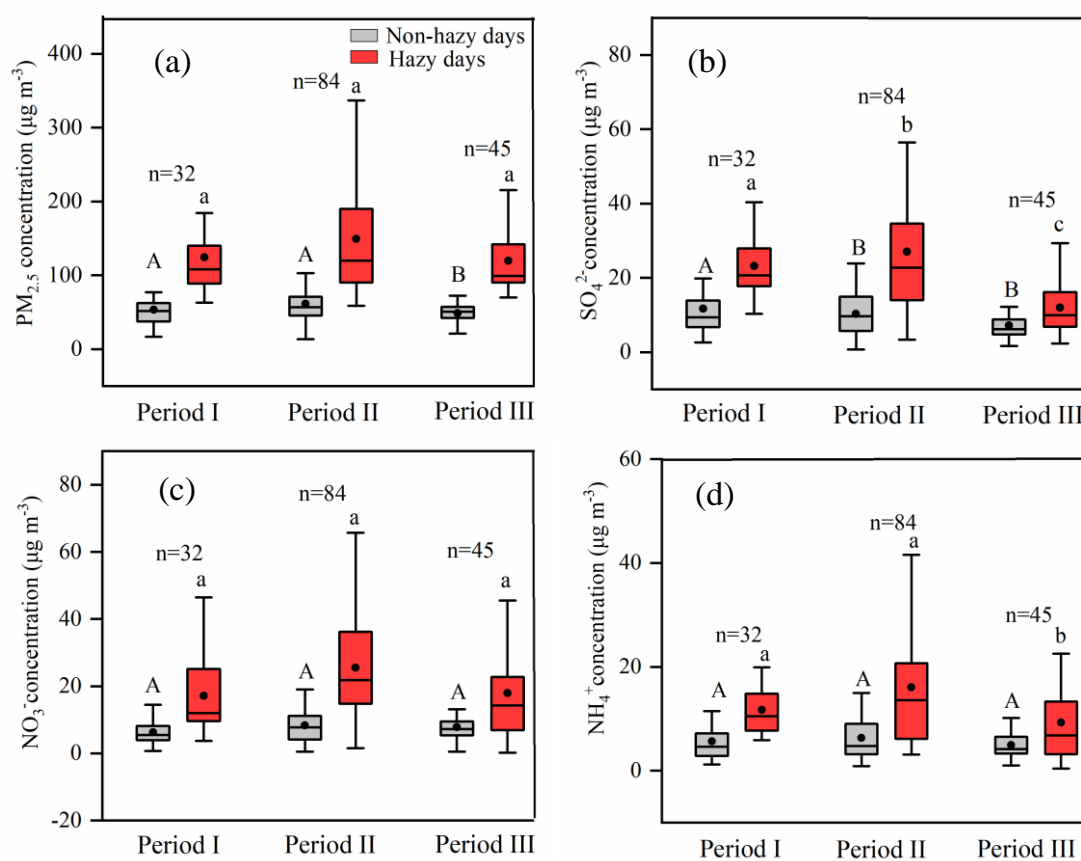
285 **3. Results and discussion**

286 **3.1. Characteristics of PM_{2.5} and its chemical components from the meta-analysis** 287 **and from nationwide observations**

288 The meta-analysis based on all published analyses of PM_{2.5} and chemical
289 component measurements during 2000–2019 reveals the changing characteristics of
290 PM_{2.5}. [To assess the annual trends in PM_{2.5} and its major chemical components, we](#)
291 [made a three-period comparison using the measurements at sites that include both PM_{2.5}](#)
292 [and secondary inorganic ions SO₄²⁻, NO₃⁻, and NH₄⁺ \(Fig. 2\). The PM_{2.5} concentrations](#)
293 [on both hazy and non-hazy days showed no significant trend from period I and period](#)
294 [II based on Kruskal-Wallis test. However, the observed concentrations of PM_{2.5} showed](#)
295 a downward trend from Period I to Period III on the non-hazy days, decreasing by [8.2%](#)

296 (Fig. 2a), despite no significant decreasing trend on the hazy days (Fig. 2a). In addition,
 297 the annual mean PM_{2.5} concentrations from the nationwide measurements showed
 298 declining trends during 2015-2019 averaged across all China and for each of the BTH,
 299 YRD, SCB, and PRD megacity clusters of eastern China (Fig. 3a, d).

300 These results reflect the effectiveness of the pollution control policies (Fig. S2)
 301 implemented by the Chinese government at the national scale. Nevertheless, PM_{2.5}
 302 remained at relatively high levels. Over 2015–2019, the annual mean PM_{2.5}
 303 concentrations at 74% of the 1498 sites (averaging $51.9 \pm 12.4 \mu\text{g m}^{-3}$, Fig. 3a) exceeded
 304 the Chinese Grade-II Standard (GB 3095–2012) of $35 \mu\text{g m}^{-3}$ (MEPC, 2012), indicating
 305 that PM_{2.5} mitigation is a significant challenge for China.

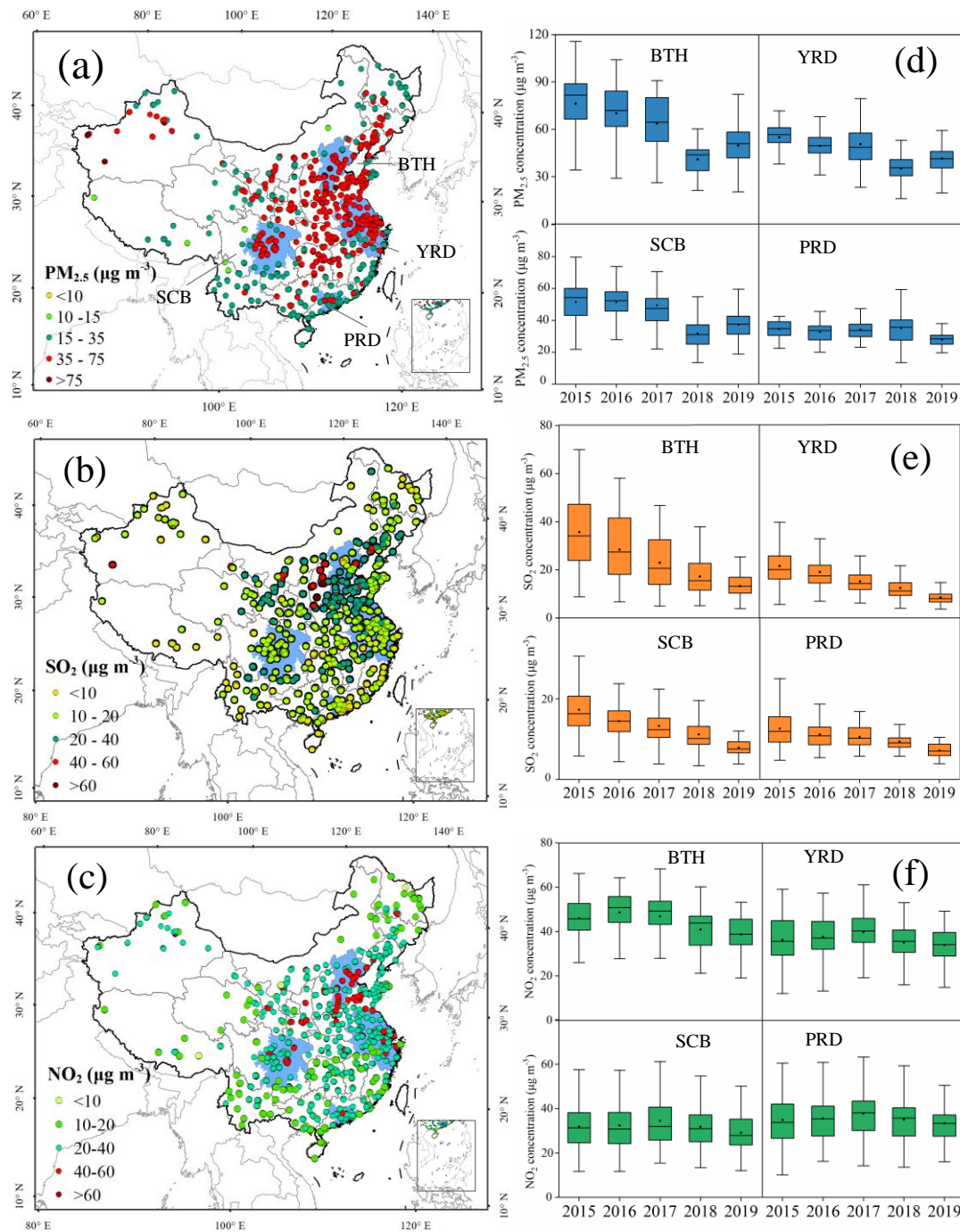


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308 **Fig. 2.** Comparisons of [observed](#) concentrations of (a) PM_{2.5}, (b) SO₄²⁻, (c) NO₃⁻, and
 309 (d) NH₄⁺ between non-hazy and hazy days in Period I (2000–2012), Period II (2013–

310 2016), and Period III (2017–2019). Bars with different letters denote significant
311 differences among the three periods ($P < 0.05$) (upper and lowercase letters for non-
312 hazy and hazy days, respectively). The upper and lower boundaries of the boxes
313 represent the 75th and 25th percentiles; the line within the box represents the median
314 value; the whiskers above and below the boxes represent the 90th and 10th percentiles;
315 the point within the box represents the mean value. [Comparison of the pollutants among](#)
316 [the three-periods using Kruskal-Wallis and Dunn’s test. The \$n\$ represents independent](#)
317 [sites; more detail on this is presented in Section 2.2.](#)
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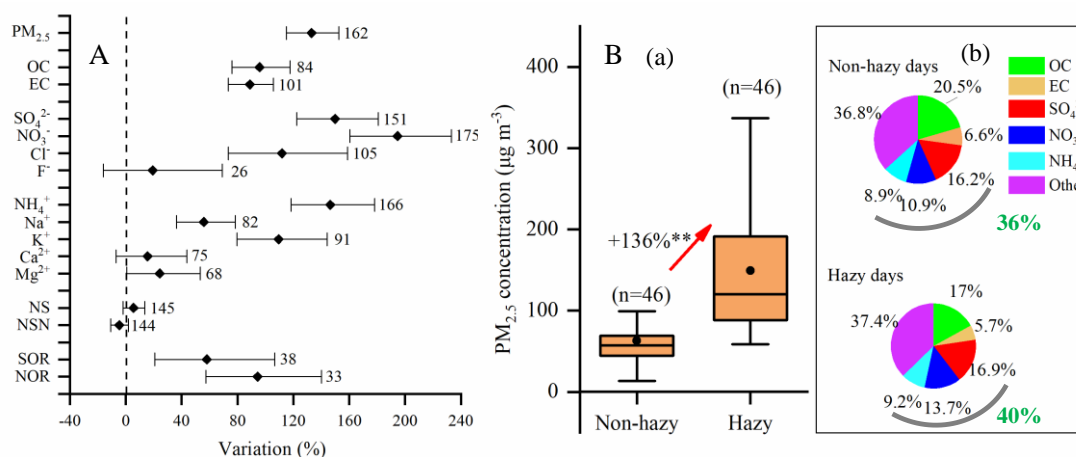
320 **Fig. 3.** Left: spatial patterns of annual mean observed concentration of (a) PM_{2.5}, (b)
 321 SO₂, (c) NO₂ at 1498 sites, averaged for 2015–2019. Right: the annual observed
 322 concentrations of (d) PM_{2.5}, (e) SO₂, and (f) NO₂ for 2015-2019 in four megacity
 323 clusters (BTH: Beijing-Tianjin-Hebei, YRD: Yangtze River Delta, SCB: Sichuan Basin,
 324 PRD: Pearl River Delta). The locations of the regions are indicated by the blue shading
 325 on the map. The upper and lower boundaries of the boxes represent the 75th and 25th

326 percentiles; the line within the box represents the median value; the whiskers above and
327 below the boxes represent the 90th and 10th percentiles; the point within the box
328 represents the mean value.

329 To further explore the underlying drivers of PM_{2.5} pollution, we analyzed the
330 characteristics of PM_{2.5} chemical components and their temporal changes in China. The
331 concentrations of PM_{2.5} and all its chemical components (except F⁻ and Ca²⁺) were
332 significantly higher on hazy days than on non-hazy days (Fig. 4A). Compared with
333 other components this difference was more significant for secondary inorganic ions (i.e.,
334 SO₄²⁻, NO₃⁻, and NH₄⁺). Sulfur oxidation ratio (SOR) and nitrogen oxidation ratio
335 (NOR) were also 58.0% and 94.4% higher on hazy days than on non-hazy days,
336 respectively, implying higher oxidations of gaseous species to sulfate- and nitrate-
337 containing aerosols on the hazy days (Sun et al., 2006; Xu et al., 2017).

338 To provide quantitative information on differences in PM_{2.5} and its components
339 between hazy days and non-hazy days, we made a comparison using 46 groups of data
340 on simultaneous measurements of PM_{2.5} and chemical components. [The 46 groups refer](#)
341 [to independent analyses from the literature that compare concentrations of PM_{2.5} and](#)
342 [major components \(SO₄²⁻, NO₃⁻, NH₄⁺, OC, and EC\) on hazy and non-hazy days](#)
343 [measured across different sets of sites.](#) As shown in Fig.4B (a), PM_{2.5} concentrations
344 significantly increased (by 136%) on the hazy days ($149.2 \pm 81.6 \mu\text{g m}^{-3}$) relative to
345 those on the non-hazy days ($63.2 \pm 29.8 \mu\text{g m}^{-3}$). By contrast, each component's
346 proportions within PM_{2.5} differed slightly, with 36% and 40% contributions by SIA on
347 non-hazy days and hazy days, respectively (Fig. 4B(b)). This is not surprising because
348 concentrations of PM_{2.5} and SIA both significantly increased on the hazy days ($60.1 \pm$
349 $37.4 \mu\text{g m}^{-3}$ for SIA) relative to the non-hazy days ($22.4 \pm 12.1 \mu\text{g m}^{-3}$ for SIA). Previous
350 studies have found that increased SIA formation is the major influencing factor for haze

351 pollution in wintertime and summertime (mainly in years since 2013) in major Chinese
 352 cities in eastern China (Huang et al., 2014; Wang et al., 2019a; Li et al., 2018). Our
 353 results extend confirmation of the dominant role of SIA to PM_{2.5} pollution over a large
 354 spatial scale in China and to longer temporal scales.

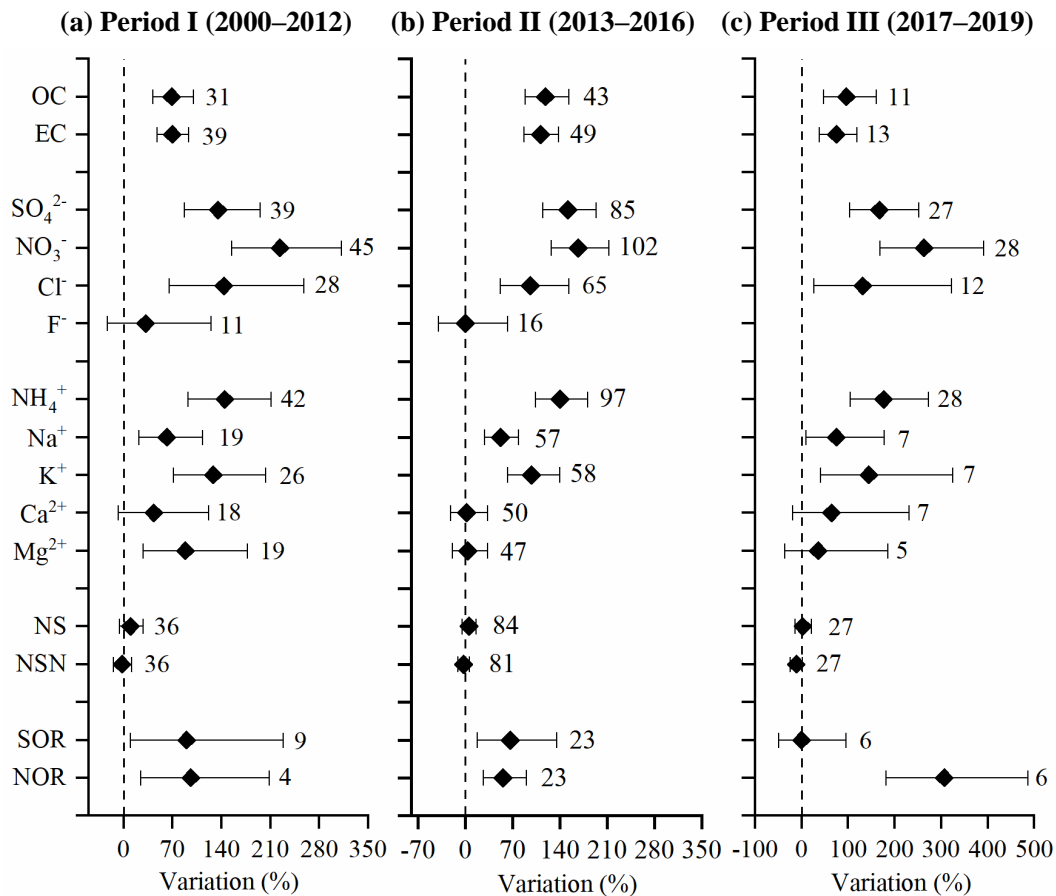


355
 356 **Fig. 4.** (A) Variations in PM_{2.5} concentration, aerosol component concentration, NS,
 357 NSN, SOR, and NOR from non-hazy to hazy days in China during 2000–2019. (B) (a)
 358 Summary of differences in PM_{2.5} concentration between non-hazy and hazy days in
 359 China; (b) the average proportions of components of PM_{2.5} on non-hazy and hazy days.
 360 NS is the slope of the regression equation between [NH₄⁺] and [SO₄²⁻], NSN is the slope
 361 of the regression equation between [NH₄⁺] and [SO₄²⁻ + NO₃⁻], SOR is sulfur oxidation
 362 ratio, and NOR is nitrogen oxidation ratio. The variations are considered significant if
 363 the confidence intervals of the effect size do not overlap with zero. ** denotes significant
 364 difference ($P < 0.01$) between hazy days and non-hazy days. The upper and lower
 365 boundaries of the boxes represent the 75th and 25th percentiles; the line within the box
 366 represents the median value; the whiskers above and below the boxes represent the 90th
 367 and 10th percentiles; the point within the box represents the mean value. Values
 368 adjacent to each confidence interval indicate number of measurement sites. [The \$n\$](#)
 369 [represents independent sites; more detail on this is presented in Section 2.2.](#)

370 The effect values of SIA on the hazy days were significantly higher than those on
371 non-hazy days for all three periods (I, II, and III) (Fig. 5), indicating the persistent
372 prevalence of the SIA pollution problem over the past two decades. Considering
373 changes in concentrations, SO_4^{2-} showed a downward trend from Period I to Period III
374 on the non-hazy days and hazy day, decreasing by 38.6% and 48.3%, respectively (Fig.
375 2b). These results reflect the effectiveness of the SO_2 pollution control policies (Ronald
376 et al., 2017). In contrast, there were no significant downward trends in concentrations
377 of NO_3^- and NH_4^+ on either hazy or non-hazy days (Fig. 2c, d), but the mean NO_3^-
378 concentration in Period III decreased by 10.5% compared with that in Period II,
379 especially on hazy days (-16.8%). These results could be partly supported by decreased
380 NO_x emissions and tropospheric NO_2 vertical column densities between 2011 and 2019
381 in China owing to effective NO_x control policies (Zheng et al., 2018; Fan et al., 2021).
382 The lack of significantly downward trends in NH_4^+ concentrations may be due to the
383 fact that the total NH_3 emissions in China changed little and remained at high levels
384 between 2000 and 2018, i.e., slightly decreased from 2000 (10.3 Tg) to 2012 (9.3 Tg)
385 (Kang et al., 2016) and then slightly increased between 2013 and 2018 (Liu et al., 2021).
386 The similar trends are also found in Quzhou in China, which is a long-term in situ
387 monitoring site (in Quzhou County, North China Plain, operated by our group) during
388 the period 2012-2020 from previous publications (Xu et al., 2016; Zhang et al., 2021b,
389 noted that data during 2017-2020 are unpublished before) (Fig. S8). Zhang et al. (2020)
390 found that the clean air actions implemented in 2017 effectively reduced wintertime
391 concentrations of PM_{10} (particulate matter with diameter $\leq 10 \mu\text{m}$), SO_4^{2-} and NH_4^+ in
392 Beijing compared with those in 2007, but had no apparent effect on NO_3^- . Our findings
393 are to some extent supported by the nationwide measurements. Annual mean SO_2
394 concentrations displayed a clear decreasing trend with a 53% reduction in 2019 relative

395 to 2015 for the four megacity clusters of eastern China (Fig. 3b, e), whereas there were
396 only slight reductions in annual mean NO₂ concentrations (Fig. 3c, f). In contrast,
397 annual mean NH₃ concentrations showed an obvious increasing trend in in both
398 northern and southern regions of China, and especially in the BTH region (Fig. S9).

399 Overall, the above analyses indicate that SO₄²⁻ concentrations responded
400 positively to air policy implementations at the national scale, but that reducing NO₃⁻
401 and NH₄⁺ remains a significant challenge. China has a history of around 10-20 years
402 for SO₂ and NO_x emission control and has advocated NH₃ controls despite to date no
403 mandatory measures and binding targets having been set (Fig. S2). Nevertheless, PM_{2.5}
404 pollution, especially SIA such as NO₃⁻ and NH₄⁺, is currently a serious problem (Fig. 4
405 and 5a, b). Some studies have reported that PM_{2.5} pollution can be effectively reduced
406 if implementing synchronous NH₃ and NO_x/SO₂ controls (Liu et al., 2019b). Therefore,
407 based on the above findings, we propose that NH₃ and NO_x/SO₂ emission mitigation
408 should be simultaneously strengthened to mitigate haze pollution.



409

410 **Fig. 5.** Variations in PM_{2.5} composition, NS, NSN, SOR, and NOR from non-hazy to

411 hazy days in (a) Period I (2000–2012), (b) Period II (2013–2016), (c) Period III (2017–

412 2019). NS is the slope of the regression equation between [NH₄⁺] and [SO₄²⁻], NSN is

413 the slope of the regression equation between [NH₄⁺] and [SO₄²⁻ + NO₃⁻], SOR is sulfur

414 oxidation ratio, and NOR is nitrogen oxidation ratio. The variations are statistically

415 significant if the confidence intervals of the effect size do not overlap with zero. Values

416 adjacent to each confidence interval indicate number of measurement sites. [The *n*](#)

417 [represents independent sites; more detail on this is presented in Section 2.2.](#)

418 3.2. Sensitivities from model simulations

419 To further examine the efficiencies of NH₃ and acid gas emission reductions on

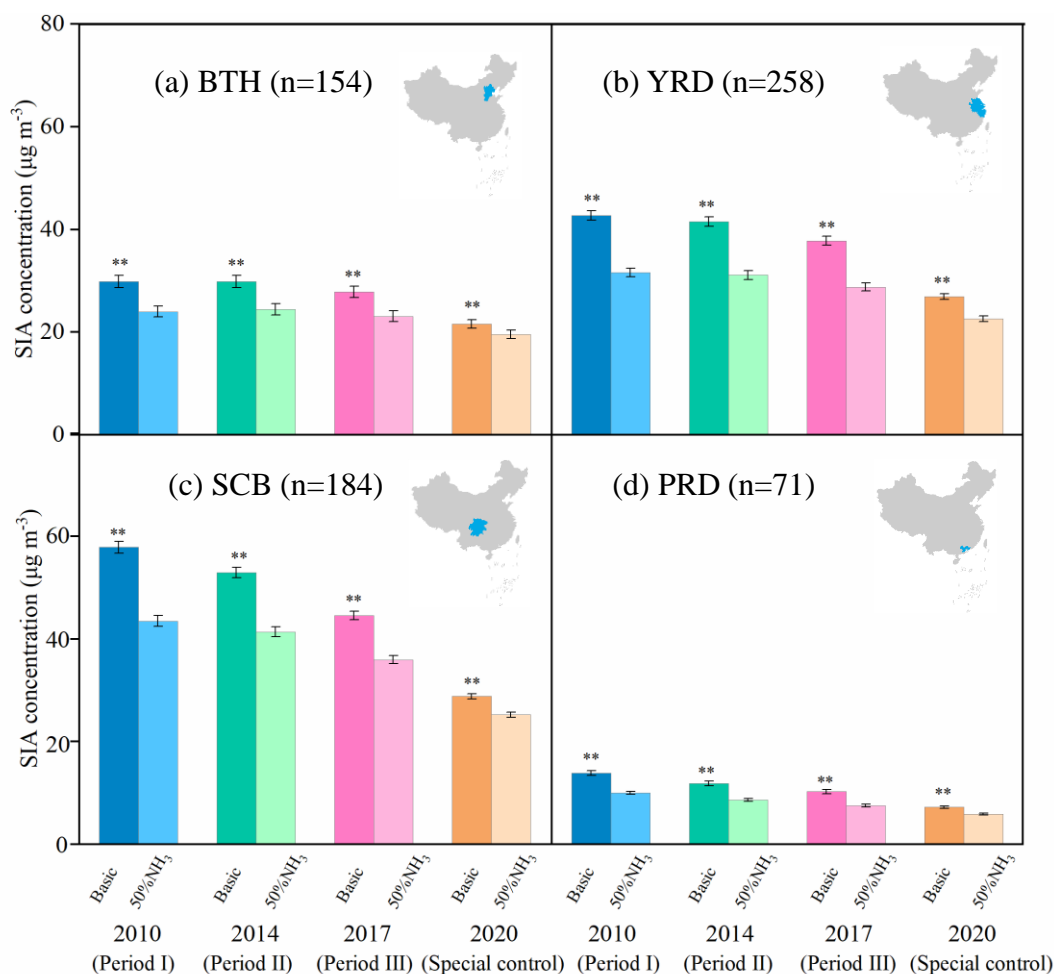
420 SIA and PM_{2.5} mitigation, the decreases of mean SIA and PM_{2.5} concentrations with and

421 without additional 50% NH₃ reductions were simulated using the WRF/CMAQ model.

422 Fig. 6 and Fig S10 shows that, compared to 2010, SIA and PM_{2.5} concentrations in
423 January in 2017 were significantly decrease in the BTH, YRD, SCB, and PRD megacity
424 clusters, respectively, in the simulations without additional NH₃ emission reductions.
425 Across the four megacity clusters, the reduction in SIA and PM_{2.5} is largest in the SCB
426 region from 2010 to 2017 and smallest in the PRD region.

427 When simulating the effects of an additional 50% NH₃ emissions reductions in
428 January in each of the years 2010, 2014 and 2017, the SIA concentrations in the BTH,
429 YRD, SCB and PRD megacity clusters decreased by $25.9 \pm 0.3\%$, $24.4 \pm 0.3\%$, and
430 $22.9 \pm 0.3\%$, respectively (Fig. 6 and Fig. S11). The reductions of PM_{2.5} in 2010, 2014
431 and 2017 were $9.7 \pm 0.1\%$, $9.0 \pm 0.1\%$, and $9.2 \pm 0.2\%$ in the megacity clusters,
432 respectively. (Figs. S10 and S12). Whilst these results confirm the effectiveness of NH₃
433 emission controls, it is important to note that the response of SIA concentrations is less
434 sensitive to additional NH₃ emission controls along the timeline of the SO₂ and NO_x
435 anthropogenic emissions reductions associated with the series of clean air actions
436 implemented by the Chinese government from 2010 to 2017 (Zheng et al., 2018). Given
437 the feasibility and current upper bound of NH₃ emission reductions options in the near
438 future (50%) (Liu et al., 2019b), further abatement of SIA concentrations merely by
439 reducing NH₃ emissions is limited in China. In other words, the controls on acid gas
440 emissions should continue to be strengthened beyond their current levels.

441

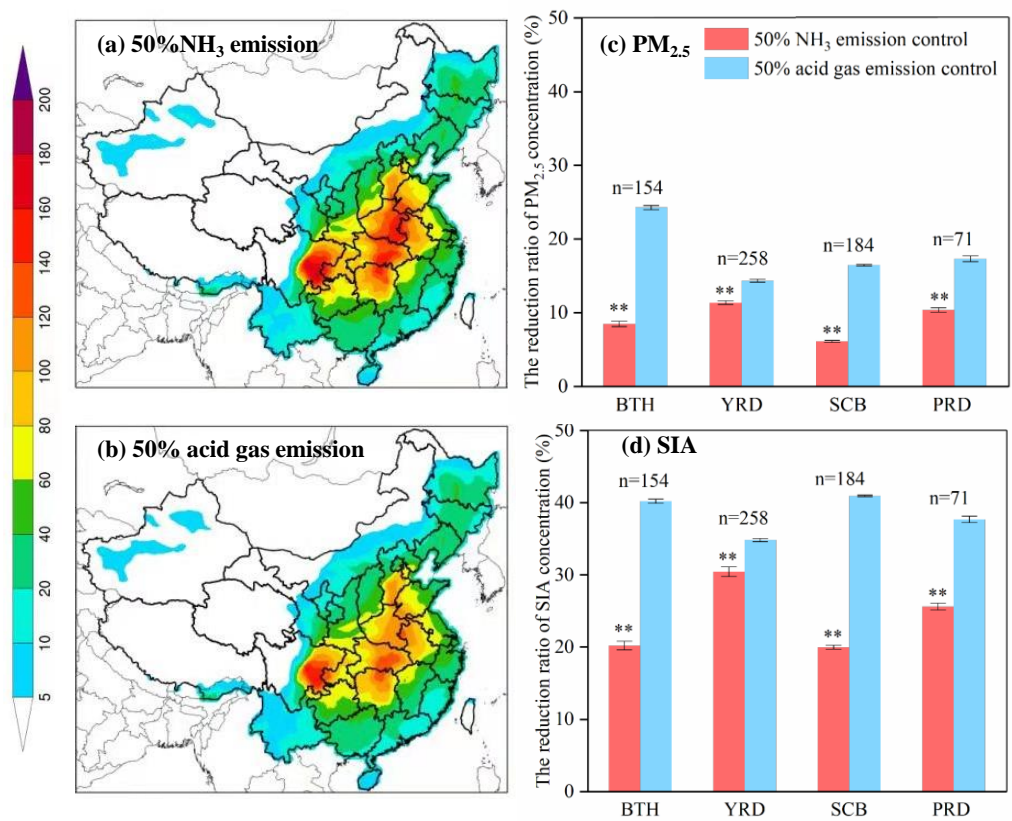


442

443 **Fig. 6.** Simulated SIA concentrations (in $\mu\text{g m}^{-3}$) without (basic) and with 50%
 444 ammonia (NH_3) emissions reductions in January for the years 2010, 2014, 2017 and
 445 2020 in four megacity clusters (BTH: Beijing-Tianjin-Hebei, YRD: Yangtze River
 446 Delta, SCB: Sichuan Basin, PRD: Pearl River Delta). Inset maps indicate the location
 447 of each region. ** denotes significant difference without and with 50% ammonia
 448 emission reductions ($P < 0.05$). n is the number of calculated samples by grid extraction.
 449 Error bars are standard errors of means. (Period I (2000–2012), Period II (2013–2016),
 450 and Period III (2017–2019); Special control is the restrictions in economic activities
 451 and associated emissions during the COVID-19 lockdown period in 2020.)

452 To further verify the above findings, we used the reductions of emissions of acid
 453 gases (46% and 23% for NO_x and SO_2 , respectively, in the whole China) during the

454 COVID-lockdown period as a further scenario (Huang et al., 2021). The model
455 simulations suggest that the effectiveness of reductions in SIA and PM_{2.5} concentrations
456 by a 50% NH₃ emission reduction further declined in 2020 ($15 \pm 0.2\%$ for SIA, and
457 $5.1 \pm 0.2\%$ for PM_{2.5}), but the resulting concentrations of them were lower ($20.8 \pm 0.3\%$
458 for SIA, and $15.6 \pm 0.3\%$ for PM_{2.5}) when compared with that in 2017 under the same
459 scenario of an additional 50% NH₃ emissions reduction (and constant meteorological
460 conditions) (Fig. 6), highlighting the importance of concurrently NH₃ mitigation when
461 acid gas emissions are strengthened. To confirm the importance of acid gas emissions,
462 another sensitivity simulation was conducted for 2017, in which the acid gas (NO_x and
463 SO₂) emissions were reduced by 50% (Fig. 7). We found that reductions in SIA
464 concentrations are $13.4 \pm 0.5\%$ greater for the 50% reductions in SO₂ and NO_x
465 emissions than for the 50% reductions in NH₃ emissions. These results indicate that to
466 substantially reduce SIA pollution it remains imperative to strengthen emission controls
467 on NO_x and SO₂ even when a 50% reduction in NH₃ emission is targeted and achieved.



468
 469 **Fig. 7.** Left: the spatial distributions of simulated PM_{2.5} concentrations (in µg m⁻³) in
 470 January 2017 with (a) 50% reductions in ammonia (NH₃) emissions and (b) 50%
 471 reductions in acid gas (NO_x and SO₂) emissions. Right: the % decreases in PM_{2.5} (c)
 472 and SIA (d) concentrations for the simulations with compared to without the NH₃ and
 473 acid gas emissions reductions in four megacity clusters (BTH: Beijing-Tianjin-Hebei,
 474 YRD: Yangtze River Delta, SCB: Sichuan Basin, PRD: Pearl River Delta). ** denotes
 475 significant differences without and with 50% ammonia emission reductions (*P* < 0.05).
 476 *n* is the number of calculated samples by grid extraction. Error bars are standard errors
 477 of means.

478 3.3. Uncertainty analysis and limitations

479 Some limitations should be noted in interpreting the results of the present study: this
 480 study examined period-to-period changes in PM_{2.5} chemical components based on a
 481 meta-analysis and the efficiencies of NH₃ and acid gas emission reductions on PM_{2.5}

482 mitigation. Some uncertainties may still exist in meta-analysis of nationwide
483 measurements owing to differences in monitoring, sample handling and analysis
484 methods as well as lack of long-term continuous monitoring sites (Fig. 2). For example,
485 the measurements of PM_{2.5} were mainly taken using the TEOM method, which is
486 associated with under-reading of PM due to some nitrate volatilization at its operational
487 temperature. To test whether the use of data during 2000–2019 could bias annual trends
488 of PM_{2.5} and chemical components, we summarize measurements of PM_{2.5} at a long-
489 term monitoring site (in Quzhou County, North China Plain, operated by our group)
490 during the period 2012-2020 from previous publications (Xu et al., 2016; Zhang et al.,
491 2021, noted that data during 2017-2020 are unpublished before). The PM_{2.5} and SO₄²⁻
492 show the same decreasing trend. The concentration of NO₃⁻ and NH₄⁺ do not show
493 significant change (Fig. S8). The results are consistent with the trend for the whole of
494 China obtained from the meta-analysis.

495 WRF-CMAQ model performance also has some uncertainty. We performed the
496 validations of WRF and CMAQ models. The simulations of temperature at 2 m above
497 ground (T2), wind speed (WS), and relative humidity (RH) versus observed values at
498 400 monitoring sites in China are shown in Fig. S7. The meteorological measurements
499 were obtained from the National Climate Data Center (NCDC)
500 (<ftp://ftp.ncdc.noaa.gov/pub/data/noaa/>). The comparisons showed that the model
501 performed well at predicting meteorological parameters with *R* values of 0.94, 0.64 and
502 0.82 for T2, WS and RH, respectively. However, the WS was overestimated (22.3%
503 NMB) in most regions of China, which is also reported in previous studies (Gao et al.,
504 2016; Chen et al., 2019). This may be related to the underlying surface parameters set
505 in the WRF model configurations.

506 In addition, the simulations of PM_{2.5} and associated chemical components by the

507 CMAQ model have potential biases in the spatial pattern, although the CMAQ model
508 has been extensively used in air quality studies (Zhang et al., 2019; Backes et al., 2016)
509 and the validity of the chemical regime in the CMAQ model had been confirmed by
510 our previous studies (Zhang et al., 2021a; Wang et al., 2020a, 2021a). Since nationwide
511 measurements of PM_{2.5} and associated chemical components are lacking in 2010 in
512 China, we undertook our own validation of PM_{2.5} and its components (such as SO₄²⁻,
513 NO₃⁻, and NH₄⁺) using a multi-observation dataset that includes those monitoring data
514 and satellite observations at a regional scale that were available.

515 First, the simulated monthly mean PM_{2.5} concentration in January 2010 was
516 compared with corresponding data obtained from the Tracking Air pollution in China
517 (TAP, <http://tapdata.org.cn/>) database. The satellite historical PM_{2.5} predictions are
518 reliable (average $R^2 = 0.80$ and RMSE = 11.26 $\mu\text{g m}^{-3}$) in a validation against the in-
519 situ surface observations on a monthly basis (Wei et al., 2020, 2021). The model well
520 captured the spatial distributions of PM_{2.5} concentrations in our studied regions of BTH,
521 YRD, PRD, and SCB (Fig. S3a), with correlation coefficient (R) between simulated and
522 satellite observed PM_{2.5} concentrations of 0.96, 0.80, 0.60, and 0.85 for BTH, YRD,
523 PRD, and SCB, respectively.

524 Second, we also collected ground-based observations from previous publications
525 (Xiao et al., 2020, 2021; Geng et al., 2019; Xue et al., 2019) to validate the modeling
526 concentrations of SO₄²⁻, NO₃⁻, and NH₄⁺. Detailed information about the monitoring
527 sites is presented in Table S5. The distributions of the simulated monthly mean
528 concentrations of SO₄²⁻, NO₃⁻, and NH₄⁺ in January 2010 over China is compared with
529 collected surface measurements are shown in Fig. S4a, b, and c, respectively, with their
530 linear regression analysis presented in Fig. S4d. The model showed underestimation in
531 simulating SO₄²⁻ and NO₃⁻ in the BTH region, which might be caused by the uncertainty

532 in the emission inventory. The lack of heterogeneous pathways for SO_4^{2-} formation in
533 the CMAQ model might also be an important reason for the negative bias between
534 simulations and measurements (Yu et al., 2005; Cheng et al., 2016). The model
535 overestimated NO_3^- concentration in the SCB region, but can capture the spatial
536 distribution of NO_3^- in other regions. The overestimation of NO_3^- has been a common
537 problem in regional chemical transport models such as CMAQ, GEOS-CHEM and
538 CAMx (Yu et al., 2005; Fountoukis et al., 2011; Zhang et al., 2012; Wang et al., 2013),
539 due to the difficulties in correctly capturing the gas and aerosol-phase nitrate
540 partitioning (Yu et al., 2005). The modeling of NH_4^+ concentrations show good
541 agreement with the observed values. Generally, the evaluation results indicate that the
542 model reasonably predicted concentrations of SO_4^{2-} , NO_3^- , and NH_4^+ in $\text{PM}_{2.5}$.

543 Third, we performed a comparison of the time-series of the observed and simulated
544 hourly $\text{PM}_{2.5}$ and its precursors (SO_2 and NO_2) during January 2010. The model well
545 captures the temporal variations of the $\text{PM}_{2.5}$ in Beijing, with an NMB value of 0.05 ug
546 m^{-3} , NME of 28%, and R of 0.92 (Fig. 5a). The predicted daily concentrations of NO_2
547 and SO_2 during January 2010 also show good agreement with the ground measurements
548 in Beijing, with NMB and R values of 0.12 ug m^{-3} and 0.89 for NO_2 , and -0.04, 0.95
549 for SO_2 , respectively (Fig. 5b). The variations of daily $\text{PM}_{2.5}$ concentrations between
550 simulation and observation at 4 monitoring sites (Shangdianzi, Chengdu, Institute of
551 Atmospheric Physics, Chinese Academy of Sciences (IAP-CAS), and Tianjin) from 14
552 to 30 January 2010 also matched well, with NMB values ranging from -0.05 to 0.12 ug
553 m^{-3} , and R values exceeding 0.89 (Fig S5c).

554 We also compared the simulated and observed concentrations of $\text{PM}_{2.5}$, NO_2 , and
555 SO_2 in China in pre-COVID period (1–26 January 2020) and during the COVID-
556 lockdown period (27 January–26 February). As shown in Fig. S6, both the simulations

557 and observations suggested that the PM_{2.5} and NO₂ concentrations substantially
558 decreased during the COVID-lockdown, mainly due to the sharp reduction in vehicle
559 emissions (Huang et al., 2021; Wang et al., 2021b). For SO₂, the concentrations
560 decreased very little and even increased at some monitoring sites. The model
561 underestimated the concentrations of PM_{2.5}, NO₂, and SO₂, with NMB values of -21.4%,
562 -22.1%, and -9.6%, respectively. This phenomenon is reasonable as the simulations for
563 the two periods in 2020 used the meteorology for 2010 whereas measured changes are
564 strongly influenced by the actual meteorological conditions.

565 **3.4. Implication and outlook**

566 Improving air quality is a significant challenge for China and the world. A key
567 target in China is for all cities to attain annual mean PM_{2.5} concentrations of 35 µg m⁻³
568 or below by 2035 (Xing et al., 2021). However, this study has shown that 74% of 1498
569 nationwide measurement sites have exceeded this limit value in recent years (averaged
570 across 2015-2019). Our results indicated that acid gas emissions still need to be a focus
571 of control measures, alongside reductions in NH₃ emissions, in order to reduce SIA (or
572 PM_{2.5}) formation. Model simulations for the month of January underpin the finding that
573 the relative effectiveness of NH₃ emission control decreased over the period from 2010
574 to 2017. However, simulating the substantial emission reductions in acid gases due to
575 the lockdown during the COVID-19 pandemic, with fossil fuel-related emissions
576 reduced to unprecedented levels, indicated the importance of ammonia emission
577 abatement for PM_{2.5} air quality improvements when SO₂ and NO_x emissions have
578 already reached comparatively low levels. Therefore, a strategic and integrated
579 approach to simultaneously undertaking acid gas emissions and NH₃ mitigation is
580 essential to substantially reduce PM_{2.5} concentrations. However, the mitigation of acid

581 gas and NH₃ emissions pose different challenges due to different sources they originate
582 from.

583 The implementation of further reduction of acid gas emissions is challenging. The
584 prevention and control of air pollution in China originally focused on the control of acid
585 gas emissions (Fig.S2). The controls have developed from desulfurization and
586 denitrification technologies in the early stages to advanced end-of-pipe control
587 technologies. By 2018, over 90% of coal-fired power plants had installed end-of-pipe
588 control technologies (CEC, 2020). The potential for further reductions in acid gas
589 emissions by end-of-pipe technology might therefore be limited. Instead, addressing
590 total energy consumption and the promotion of a transition to clean energy through a
591 de-carbonization of energy production is expected to be an inevitable requirement for
592 further reducing PM_{2.5} concentrations (Xing et al., 2021). In the context of improving
593 air quality and mitigating climate change, China is adopting a portfolio of low-carbon
594 policies to meet its Nationally Determined Contribution pledged in the Paris Agreement.
595 Studies show that if energy structure adjusts and energy conservation measures are
596 implemented, SO₂ and NO_x will be further reduced by 34% and 25% in Co-Benefit
597 Energy scenario compared to the Nationally Determined Contribution scenario in 2035
598 (Xing et al., 2021). Although it has been reported that excessive acid gas emission
599 controls may increase the oxidizing capacity of the atmosphere and increase other
600 pollution, PM_{2.5} concentrations have consistently decreased with previous acid gas
601 control (Huang et al., 2021). In addition, under the influence of low-carbon policies,
602 other pollutant emissions will also be controlled. Opportunities and challenges coexist
603 in the control of acid gas emissions.

604 In contrast to acid gas emissions, NH₃ emissions predominantly come from
605 agricultural sources. Although the Chinese government has recognized the importance

606 of NH₃ emissions controls in curbing PM_{2.5} pollution, NH₃ emissions reductions have
607 only been proposed recently as a strategic option and no specific nationwide targets
608 have yet been implemented (CSC, 2018b). The efficient implementation of NH₃
609 reduction options is a major challenge because NH₃ emissions are closely related to
610 food production, and smallholder farming is still the dominant form of agricultural
611 production in China. The implementation of NH₃ emissions reduction technologies is
612 subject to investment in technology, knowledge and infrastructure, and most farmers
613 are unwilling or economically unable to undertake additional expenditures that cannot
614 generate financial returns (Gu et al., 2011; Wu et al., 2018b). Therefore, economically
615 feasible options for NH₃ emission controls need to be developed and implemented
616 nationwide.

617 We propose the following three requirements that need to be met to achieve
618 effective reductions of SIA concentrations and hence of PM_{2.5} concentrations in China.

619 First, binding targets to reduce both NH₃ and acid gas emissions should be set. The
620 targets should be designed to meet the PM_{2.5} standard, and NH₃ concentrations should
621 be incorporated into the monitoring system as a government assessment indicator. In
622 this study, we find large differences in PM_{2.5} concentration reductions from NH₃
623 emissions reduction in the four megacity regions investigated. At a local scale (i.e., city
624 or county), the limiting factors may vary within a region (Wang et al., 2011). Thus,
625 local-specific environmental targets should be considered in policy-making.

626 Second, further strengthening of the controls on acid gas emissions are still needed,
627 especially under the influence of low-carbon policies, to promote emission reductions
628 and the adjustment of energy structures and conservation. Ultra-low emissions should
629 be requirements in the whole production process, including point source emissions,
630 diffuse source emissions, and clean transportation (Xing et al., 2021; Wang et al.,

631 [2021a](#)). The assessment of the impact of ultra-low emissions is provided in [Table S6](#).

632 In terms of energy structure, it is a requirement to eliminate outdated production
633 capacity and promote low-carbon new energy generation technologies.

634 Third, a requirement to promote feasible NH₃ reduction options throughout the
635 whole food production chain, for both crop and animal production. Options include the
636 following. 1) Reduction of nitrogen input at source achieved, for example, through
637 balanced fertilization based on crop needs instead of over-fertilization, and promotion
638 of low-protein feed in animal breeding. 2) Mitigation of NH₃ emissions in food
639 production via, for example, improved fertilization techniques (such as enhanced-
640 efficiency fertilizer (urease inhibitor products), fertilizer deep application, fertilization-
641 irrigation technologies ([Zhan et al., 2021](#)), and coverage of solid and slurry manure. 3)
642 Encouragement for the recycling of manure back to croplands, and reduction in manure
643 discarding and long-distance transportation of manure fertilizer. Options for NH₃
644 emissions control are provided in [Table S4](#). Although the focus here has been on
645 methods to mitigate NH₃ emissions, it is of course critical simultaneously to minimize
646 N losses in other chemical forms such as nitrous oxide gas emissions and aqueous
647 nitrate leaching ([Shang et al., 2019](#); [Wang et al., 2020b](#)).

648 **4. Conclusions**

649 The present study developed an integrated assessment framework using meta-
650 analysis of published literature results, analysis of national monitoring data, and
651 chemical transport modelling to provide insight into the effectiveness of SIA precursor
652 emissions controls in mitigating poor PM_{2.5} air quality in China. We found that PM_{2.5}
653 concentration significantly decreased in 2000-2019 due to acid gas control policies, but
654 PM_{2.5} pollution still severe. Compared with other components, this difference was more
655 significant higher (average increase 98%) for secondary inorganic ions (i.e., SO₄²⁻, NO₃⁻,

656 [and NH₄⁺\) on hazy days than on-hazy days.](#) This is mainly caused by the persistent SIA
657 pollution during the same period, with sulfate concentrations significantly decreased
658 and no significant changes observed for nitrate and ammonium concentrations. The
659 reductions of SIA concentrations in January in megacity clusters of eastern China by
660 additional 50% NH₃ emission controls decreased from 25.9 ± 0.3% in 2010 to 22.9 ±
661 0.3% in 2017, and to 15 ± 0.2% in the COVID lockdown in 2020 for simulations
662 representing reduced acid gas emissions to unprecedented levels, but the SIA
663 concentrations decreased by 20.8 ± 0.3% in 2020 compared with that in 2017 under the
664 same scenario of an additional 50% NH₃ emissions reduction. In addition, the reduction
665 of SIA concentration in 2017 was 13.4 ± 0.5% greater for 50% acid gas (SO₂ and NO_x)
666 reductions than for the NH₃ emissions reduction. These results indicate that acid gas
667 emissions need to be further controlled concertedly with NH₃ reductions to substantially
668 reduce PM_{2.5} pollution in China.

669 Overall, this study provides new insight into the responses of SIA concentrations
670 in China to past air pollution control policies and the potential balance of benefits in
671 including NH₃ emissions reductions with acid gas emissions controls to curb SIA
672 pollution. The outcomes from this study may also help other countries seeking feasible
673 strategies to mitigate PM_{2.5} pollution.

674

675 **Data availability**

676 All data in this study are available from the from the corresponding authors (Wen Xu,
677 wenxu@cau.edu.cn; Shaocai Yu, shaocaiyu@zju.edu.cn) upon request.

678 **Author contributions**

679 W.X., S.Y., and F.Z. designed the study. F.M., Y.Z., W.X., and J.K. performed the
680 research. F.M., Y.Z., W.X., and J.K. analyzed the data and interpreted the results. Y.Z.

681 conducted the model simulations. L.L. provided satellite-derived surface NH₃
682 concentration. F.M., W.X., Y.Z., and M.R.H. wrote the paper, S. R., M.W., K.W., J.K.,
683 Y.Z., Y.H., P.L., J.W., Z.C., X.L., M.R.H., S.Y. and F.Z. contributed to the discussion
684 and revision of the paper.

685 **Declaration of Competing Interest**

686 The authors declare that they have no known competing financial interests or personal
687 relationships that could have appeared to influence the work reported in this paper.

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