Response to reviewers' comments

Thanks to the reviewers for giving us very useful comments to improve our manuscript entitled "Modeling the impact of heterogeneous reactions of chlorine on summertime nitrate formation in Beijing, China" (acp-2018-1270). Detailed responses to reviewers' comment are list below:

8 Report #1

10 <u>Comment 1:</u> Qiu et al., developed a revised regional CMAQ model, by updating 11 several chloride contained reactions, to study the impact of chloride-related 12 heterogeneous chemistry on summer particulate nitrate formation. This work is 13 meaningful and improved the understanding of regional nitrate pollution as well as 14 chloride chemistry. Several comments should be addressed before publishing in ACP.

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16 <u>Response #1</u>: Thanks for the positive comments on this manuscript. The suggestions
 17 are addressed in detail in the following.

19 <u>Comment 2:</u> Section 3.2, the uptake coefficient of O_3 on chloride-contain aerosol 20 were increased by a factor of 10 and showed the importance of O3 heterogeneous 21 reaction and the Cl2 budget. The increasing of O_3 uptake coefficient may also strongly 22 affect the O3 lifetime, an intercomparison of modeled and observed O3 need to be 23 conducted to verify the rationality and prove the significance.

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Response #2: In previous response #3 from the reviewer #1, we have performed the 25 comparison of predicted O_3 concentration in HET case with observation, as Table R1. 26 We further compare the predicted O₃ concentration in uptake coefficient increased by a 27 factor of 10 (represent as Sim* in Table R1) with observation and simulation in HET 28 case. The results show that O₃ concentrations in 'Sim*' column are slight lower than 29 the 'Sim' column, which may indicate that the O₃ consumption by heterogeneous 30 reaction is generally larger than the production owing to the Cl₂ formation. Moreover, 31 these differences are mainly found in nighttime because the weak photolysis limit the 32 generated Cl_2 to transfer to O_3 . 33

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Table R1 Comparison of simulated 1-h NO₂, PM_{2.5} and O₃ concentrations with observations averaged from 11 to 15, June, 2017 (Obs: observation, Sim:simulation).

NO ₂				O3			PM _{2.5}						
Sites	Obs	Sim	NMB	NME	Obs	Sim	Sim*	NMB	NME	Obs	Sim	NMB	NME
WSXG	49	54	11%	55%	99	122	121.6	23%	63%	40	38	-6%	53%
DL	21	17	-20%	68%	111	108	107.5	-2%	41%	32	29	-10%	52%
DS	47	53	13%	54%	100	114	113.8	15%	56%	44	41	-7%	53%
TT	40	48	20%	64%	98	130	129.4	33%	60%	37	37	1%	58%
NZG	51	66	28%	62%	111	121	120.5	9%	57%	42	39	-7%	52%
GY	55	65	17%	57%	107	116	115.4	9%	75%	36	33	-8%	54%
WL	52	41	-21%	54%	92	112	111.6	22%	73%	35	33	-7%	54%
XC	43	31	-28%	47%	100	108	107.3	8%	52%	33	29	-12%	55%
HR	26	11	-56%	70%	124	105	104.4	-15%	47%	27	22	-19%	51%
СР	42	28	-34%	58%	96	91	90.3	-5%	77%	33	32	-1%	54%
ATZX	56	62	10%	55%	105	107	106.2	1%	68%	33	31	-4%	54%
GC	56	42	-25%	58%	106	107	106.4	0%	59%	43	37	-14%	52%

- 38
- WSXG: Wanshouxigong; DL: Dingling; DS: Dongsi; TT:Tiantan; NZG:Nongzhanguan;

40 GY: Guanyuan; WL: Wanliu; XC:Xincheng; HR:Huairou; CP:Changping;
41 ATZX:Aotizhongxin; GC:Gucheng

- 42 Sim*: O₃ concentration in uptake coefficient increased by a factor of 10
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44 **Comment 3:** Figure 3(e, f) do not show the higher OH plot in Het case, so the 45 statement in line 366 can't get the conclusion that high ANO3 is attribute to the elevated 46 OH, although other studies presented the same conclusion. By the way, the photolysis 47 of ClNO2 also release NO2 and enhance the reaction of OH + NO2. It would be good 48 if the increasing of daytime NO2 and OH by ClNO2 photolysis can be quantified and 49 used to assess the contribution from NO2 and OH.

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51 <u>Response #3</u>: We have supplemented the spatial distribution of diurnal OH 52 concentration (panel (a)) and difference ((HET/BASE)/BASE, panel (b)) averaged from 53 11-15 June 2017. Which can reflect that the high ANO3 is attribute to the elevated OH. 54 We add the Figure R1 in SI.

As the reviewer's representation, the photolysis of $CINO_2$ also release NO_2 and enhance the reaction of $OH + NO_2$. However, it's hard to quantify its contribution to NO_2 in the model. But we think this process can be ignored because the NO_2 generated by $CINO_2$ is rather low (because the $CINO_2$ is rather low, less than 1ppb). By contrast, the atmospheric NO_2 concentration is reaching up to $20{\sim}40$ ppb.

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Figure R1. Spatial distributions of episode-average OH concentration (a) and the
difference (b) from 11 to 15 June 2017. Unit: 10⁶ molecules cm⁻³

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65 <u>Comment 4:</u> L89, high Cl- do not represent high ClNO2 yield, which may be subject
66 to the aerosol liquid water content, organics and so on, please change to "When ClNO2
67 yield is high...".

- 72 **<u>Comment 5:</u>** Line 229, Eqs. 8, change to γOH.
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74 **<u>Response #5</u>**: Revised.

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- 75 Comment 6: Line 321, Bracket is not complete. 76 77 **Response #6**: Revised 78 79 **Comment 7:** Line 328, Tham et al., 2019 change to 2018. 80 81 82 **Response #7**: Revised. 83 **Comment 8:** Line 718-719, the font size is bigger than other words. 84 85 Response #8: Revised. 86 87 88 **Comment 9:** Line 332-336, the long sentence is difficult to read. Please divide into two sentences, "... and the larger constant N2O5 leads to..." change to ". The 89 application of large and fixed N2O5 uptake coefficient ... " 90 91 92 **Response #9**: We have revised following the reviewer's option. 93 94 Comment 10: Line 377, subscript of "PM2.5". 95 Response #10: Revised. 96 97 **Comment 11:** Figure 4 should add the base case percentage like HET in the left edge. 98 99 **Response #11**: The left edge in Figure 4 presents the contribution of heterogeneous 100 reaction and HNO₃ partitioning to nitrate formation, which is concentration 101 contribution, not the percentage. 102 103 104 Report #2 105 106 **Comment 1:** There is one thing I would like to mention, although it will not change 107 the results and the conclusion of the present study. 108 In Response 35, the authors stated that "HONO photolysis affects the OH level just a 109 110 few hours in the morning and can be neglected", which is not correct. It has been well established that HONO is produced both in nighttime and in the daytime, and HONO 111 is the predominant source of OH radical during daytime (not just a few hours in the 112 morning) in the polluted environment, e.g. Fu et al., 2019 (Fig 2 and 5) and the reference 113 therein. 114 115 Fu, X., Wang, T., Zhang, L., Li, Q., Wang, Z., Xia, M., Yun, H., Wang, W., Yu, C., Yue, 116 D. and Zhou, Y., 2019. The significant contribution of HONO to secondary pollutants 117 during a severe winter pollution event in southern China. Atmospheric Chemistry and 118
- 119 Physics, 19(1), pp.1-14.
- 120

121 **<u>Response #1</u>**: Thanks for the reviewer correcting our addressing. In original CMAQ, 122 HONO is produced by the reaction NO₂ with H_2O . However, this study improves this 123 reaction to produce ClNO and nitrate, not involve HONO. From Fu's work, there are

some other HONO sources, which may increase the OH level in daytime. But we don't

- intend to improve these to our study since our work focus on the impact of chlorine heterogeneous reaction on the nitrate.

1	Modeling the impact of heterogeneous reactions of chlorine on
2	summertime nitrate formation in Beijing, China
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19 20	Abstract:
21	A comprehensive chlorine heterogeneous chemistry is incorporated into the
22	Community Multiscale Air Quality (CMAQ) model to evaluate the impact of chlorine-
23	related heterogeneous reaction on diurnal and nocturnal nitrate formation and quantify
24	the nitrate formation from gas-to-particle partitioning of HNO_3 and from different
25	heterogeneous pathways. The results show that these heterogeneous reactions increase
26	the atmospheric Cl_2 and $ClNO_2$ level (~100%), which further affect the nitrate
27	formation. Sensitivity analyses of uptake coefficients show that the empirical uptake
28	coefficient for the O ₃ heterogeneous reaction with chlorinated particles may lead to the
29	large uncertainties in the predicted Cl_2 and nitrate concentrations. The $\mathrm{N}_2\mathrm{O}_5$ uptake
30	coefficient with particulate Cl ⁻ concentration dependence performs better to capture the
31	concentration of $CINO_2$ and nocturnal nitrate concentration. The reaction of OH and
32	NO_2 in daytime increases the nitrate by ${\sim}15\%$ when the heterogeneous chlorine
33	chemistry is incorporated, resulting in more nitrate formation from HNO3 gas-to-
34	particle partitioning. By contrast, the contribution of the heterogeneous reaction of

N₂O₅ to nitrate concentrations decreases by about 27% in the nighttime when its 35 reactions with chlorinated particles are considered. However, the generated gas-phase 36 ClNO₂ from the heterogeneous reaction of N₂O₅ and chlorine-containing particles 37 further reacts with the particle surface to increase the nitrate by 6%. In general, this 38 study highlights the potential of significant underestimation of daytime and 39 40 overestimation of nighttime nitrate concentrations for chemical transport models without proper chlorine chemistry in the gas and particle phases. 41

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43 1. Introduction

44 In recent years, nitrate has become the primary component of PM_{2.5} (particulate matter with aerodynamic diameter less than 2.5µm) in Beijing with sustained and rapid 45 reduction of SO₂ and primary particulate matter emissions (Ma et al., 2018; Li et al., 46 2018; Wen et al., 2018). Observations showed that the relative contributions of 47 secondary nitrate in PM_{2.5} could reach up to approximately 50% during some severe 48 49 haze pollution days (Li et al., 2018). The mechanism of secondary nitrate formation can be summarized as two major pathways: (1) Gas-to-particle partitioning of HNO₃, which 50 happens mostly in the daytime. The reaction of OH with NO₂ produces gaseous HNO₃, 51 which subsequently partition into the particle phase. The existence of NH₃ or basic 52 53 particles enhances this process by NH₃–NH₄⁺ gas-particle equilibrium (Kleeman et al., 2005; Seinfeld and Pandis, 2006); (2) Hydrolysis of N₂O₅, which is more important at 54 nighttime. N₂O₅ forms from the reactions of NO₂, O₃ and NO₃ and hydrolyzes to 55 produce particulate nitrate. They can be summarized as reactions R1-R5 (Brown and 56 Stutz 2012): 57

 $OH + NO_2 \rightarrow HNO_3$ (R1) 58

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$$HNO_3 + NH_3 \rightarrow NH_4^+ + NO_3^-$$
 (R2)
60 $NO_2 + O_3 \rightarrow NO_2 + O_2$ (R3)

$$NO_2 + O_3 \rightarrow NO_3 + O_2$$
(R3)
61
$$NO_3 + NO_2 \leftrightarrow N_2O_5$$
(R4)

62
$$N_2O_5 + H_2O(aq) \rightarrow 2H^+ + 2NO_3^-$$
 (R5)

63 In addition to reactions R1 and R5, gas phase reactions of NO₃ with HO₂ and VOCs, N₂O₅ with water vapor (Tuazon et al., 1983) and the heterogeneous reaction of NO₂ 64 with water-containing particle (Goodman et al., 1999) produce HNO₃ or nitrate as well. 65 Theses reactions are listed in Table 2 as reactions R8, R9 and R10. 66

67 However, chemistry transport models (CTMs) incorporated with these mechanisms still can't accurately capture the spatiotemporal distributions of nitrate in some studies in polluted northern China. For example, Chang et al. (2018) showed that the simulated nitrate concentrations derived from the default CMAQ (version 5.0.2) were significantly higher than the observations in summer at two sites adjacent to Beijing. Fu et al. (2016) also found that default CMAQ (version 5.0.1) overestimated the simulated nitrate concentrations in the Beijing-Tianjin-Hebei region.

In recent fields studies, it was found that high particulate chlorine emissions might 74 have a significant impact on the oxidation capacity of the urban atmosphere and thus 75 76 could affect nitrate concentrations. According to the field measurements in June 2017 in Beijing (Zhou et al., 2018), the 2-min averaged concentrations of reactive molecular 77 chlorine (Cl₂) and nitryl chloride (ClNO₂) reached up to 1000 pptv and 1200 pptv, 78 respectively, during some severe air pollution periods in summer. The Cl₂ 79 concentrations were significantly higher than those observed in North American coastal 80 cities affected by onshore flow and the lower atmosphere in the remote Arctic region 81 (Spicer et al., 1998; Glasow et al., 2010; Liu et al., 2017). During these pollution events, 82 the corresponding concentrations of N₂O₅ (2-min average) and nitrate (5-min average) 83 rose from 40 pptv and 1 μ g m⁻³ to 700 pptv and 5 μ g m⁻³. To explain the high levels of 84 85 CINO₂, some studies suggested that reaction R5 should be revised to account for CINO₂ production from the heterogeneous reaction of N₂O₅ on chloride-containing particles 86 87 (CPS) (Osthoff et al., 2008; Thornton et al., 2010), as shown in reaction R6:

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 $N_2O_5 + (1-\phi) H_2O + \phi Cl^{-} \rightarrow \phi ClNO_2 + (2-\phi) NO_3^{-}$ (R6)

89 where ϕ represents the molar yield of ClNO₂. When Cl⁻ is enough, this reaction leads 90 to lower nitrate concentrations than reaction R5. By incorporating this reaction into WRF-Chem, Li et al. (2016) found that the improved model performed better to match 91 92 the observed nitrate concentrations in Hongkong during 15 November and 5 December 2013. However, ClNO₂ could affect the formation of nitrate indirectly by increasing the 93 atmospheric OH after a series of chemical reactions, which are briefly summarized into 94 95 three steps: (1) the photolysis of ClNO₂ produces chlorine radicals (Cl[•]); (2) the reaction of Cl[•] with VOCs produces peroxy radical (HO₂ and RO₂); and (3) the increased HO₂ 96 and RO₂ prompt the formation of OH through HOx cycle and lead to increased HNO₃ 97 production (Young et al., 2014; Jobson et al., 1994). The overall impact of R6 on nitrate 98 remains to be investigated. 99

100 Another related but unresolved issue is the sources of the high concentrations Cl_2 , 101 which could not be explained by the N₂O₅ heterogeneous reaction with Cl^- and the

subsequent reactions of ClNO₂ in the gas phase. It has been reported that the reactions 102 of gaseous O₃, OH, HO₂, ClNO₂, hypochlorous acid (HOCl), chlorine nitrate (ClONO₂) 103 with CPS can produce Cl₂, which can subsequently photolyze to produce Cl[•] 104 (Knipping et al., 2000; George et al., 2010; Pratte et al., 2006; Deiber et al., 2004; Faxon 105 et al., 2015). However, these heterogeneous reactions on CPS are generally missing in 106 most of the current CTMs and it is unclear whether these reactions will be able to 107 explain the observed Cl₂ concentrations and the overall impact of these reactions on 108 nitrate. 109

110 Previously, biomass burning, coal combustion, and waste incineration were identified as the main sources of gaseous and particulate chlorine compounds in China 111 from International Global Atmospheric Chemistry Program's Global Emissions 112 Inventory Activity (GEIA) based on the year 1990 and a localized study by Fu et al. 113 based on the year 2014 (Keene et al., 1999; Fu et al., 2018). However, recent source 114 115 apportionment results of $PM_{2.5}$ in Beijing showed that the contribution of coal combustion had extremely decreased from 22.4% in 2014 to 3% in 2017 with the 116 117 replacement of natural gas (obtained from official website of Beijing Municipal Bureau of Statistics, available at http://edu.bjstats.gov.cn/). Another important source-cooking 118 has received attention as its increasing contribution to PM2.5 (accounting for 33% of the 119 residential sector; obtained from the official source apportionment analysis of PM_{2.5} in 120 Beijing in 2017; see http://www.bjepb.gov.cn/bjhr-b/index/index.html). Moreover, the 121 high content of particulate sodium chloride was measured from the source 122 characterization studies of PM_{2.5} released from the cooking activities (Zhang et al., 123 124 2016). Thus, it is necessary to compile an updated emission inventory for Beijing to include the emissions from cooking and other sources (coal burning, solid waste 125 incineration, biomass burning, etc.) in order to explore the emissions of the chlorine 126 species on atmospheric nitrate formation. 127

In this study, a CMAQ model with an improved chlorine heterogeneous chemistry is applied to simulate summer nitrate concentration in Beijing. Sensitivity simulations are conducted to evaluate the contributions of HNO₃ gas-to-particle partitioning and heterogonous production to aerosol nitrate. The results of this work can improve our understandings on nitrate formation and provide useful information on nitrate pollution control strategies in Beijing.

135 **2. Emissions, chemical reactions and model description**

136 2.1 Emissions

Generally, the conventional emission inventories of air pollutants in China only include 137 138 the common chemical species, such as SO₂, NO_X, VOCs, PM_{2.5}, PM₁₀, NH₃, BC, and OC (Wang et al., 2014). Chloride compound emissions were not included. However, 139 the emissions of chlorine species are vital for studying the chlorine chemical 140 mechanism. Recently, the inorganic hydrogen chloride (HCl) and fine particulate 141 chloride (PCl) emission inventories for the sectors of coal combustion, biomass burning, 142 and waste incineration were developed for the year of 2014 (Qiu et al., 2016, Fu et al., 143 2018, Liu et al., 2018). However, the gaseous chlorine emission was not estimated in 144 these studies. In addition, these studies did not account for the rapid decrease of coal 145 consumption in recent years in Beijing, from 2000 Mt in 2014 to 490 Mt in 2017. More 146 importantly, the cooking source, as one of the major contributors to particulate chlorine 147 in Beijing, is not included in current chlorine emission inventories. Thus, a new 148 emission inventory of reactive chlorine species, which includes HCl, Cl₂ and PCl, were 149 developed in this study for the year of 2017. 150

151 The emission factor method (equation (1)) is applied to calculate the emissions of 152 these reactive chlorine species from coal combustion, biomass burning, municipal solid 153 waste incineration and industrial processes:

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$$E_{i,j} = A_i \times EF_{i,j} \tag{1}$$

where $E_{i,j}$ represents the emission factor of pollutant j in sector i; A represents the 155 activity data; EF represents the emission factor. EF for PCl is estimated by $EF_{i,PCl} =$ 156 157 $EF_{i,PM2.5} \times f_{Cl,i}$, where $f_{Cl,i}$ represents the mass fraction of PCl in primary PM_{2.5}. Activity data are obtained from the Beijing Municipal Bureau of Statistics (available at 158 http://tjj.beijing.gov.cn/). The Cl₂ emission factor for coal combustion is calculated 159 based on the content of Cl in coal, which had been measured by Deng et al (2017). The 160 PM_{2.5} emission factors and mass fractions of PCl in PM_{2.5} to calculate the emissions of 161 Cl had been described in detail by Fu et al. (2018). PCl in PM_{2.5} for coal combustion 162 and biomass burning are taken as 1% and 9.0%, respectively, based on local 163 164 measurements in Beijing.

165 Emissions of PCl from cooking, including contributions from commercial and 166 household cooking, are estimated using equation (2):

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$$E_{PCl} = \left[N_f \times V_f \times H_f \times EF_{f,PCl} + V_c \times H_c \times N_c \times n \times EF_{c,PCl} \times (1-\eta)\right] \times 365$$
(2)

where N_f is the number of households, V_f is the volume of exhaust gas from a household 168 stove (2000 m³ h⁻¹); H_f is the cooking time for a family (0.5 h day⁻¹); $EF_{f,PCl}$ and 169 $EF_{c,PCl}$ are the emission factors (kg m⁻³) of PCl for household and commercial cooking, 170 respectively; *Hc* is the cooking time in a commercial cooking facility (6 h day⁻¹); *Nc* is 171 the number of restaurants, schools and government departments. Vc is the volume of 172 exhaust gas from a commercial cooking stove (8000 m³ h⁻¹); n is the number of stoves 173 for each unit, which equals to 6 for a restaurant and is calculated as one stove per 150 174 students for each school. η is the removal efficiency of fume scrubbers (30%). $EF_{c,PCl}$ 175 is the emission factor (kg m⁻³) of PCl in commercial cooking. These constants are all 176 based on Wu et al. (2018). The PCl fraction in PM_{2.5} from cooking is take as 10%, based 177 178 on local measurements. HCl and Cl₂ emissions from cooking are not considered in this study. 179

The sectoral emissions of HCl, Cl₂ and PCl are summarized in Table 1. The 180 estimated HCl, Cl₂ and PCl emissions in Beijing are 1.89 Gg, 0.07Gg and 0.63Gg 181 respectively. The Cl emissions estimated for 2014 by Fu et al. (2018) were used for 182 other areas. This simplification is a good approximation because replacing coal with 183 natural gas only occurred in Beijing, and reduction of coal consumption in surrounding 184 regions was generally less than 15%. In addition, strict control measures for biomass 185 burning, cooking and municipal solid waste incineration have not been implemented in 186 187 most regions yet. Emissions of conventional species for this study period are developed in a separate study that is currently under review and are summarized in Table S1. 188

189

190 2.2 Chlorine-related heterogeneous reactions

191 The heterogeneous reactions in original CMAQ (version 5.0.1) are not related to 192 chlorine species. In this study, the original heterogeneous reactions of N₂O₅ and NO₂ 193 (R5 and R10 in Table 2) are replaced with a revised version which includes production 194 of ClNO₂ from CPS (R6 and R11 in Table 2). In reaction R6, the molar yield of ClNO₂ 195 (ϕ_{ClNO_2}) is represented as equation (3) (Bertram and Thornton, 2009):

$$\phi_{ClNO_2} = \left(1 + \frac{[H_20]}{483 \times [Cl^-]}\right)^{-1}$$
(3)

where $[H_2O]$ and $[Cl^-]$ are the molarities of liquid water and chloride (mol m⁻³).

197 In addition, laboratory observations confirmed that the heterogeneous uptake of 198 some oxidants (such as O₃ and OH) and reactive chlorine species (such as ClNO₂, HOCl, and ClONO₂) could also occur on CPS to produce Cl₂. These reactions are implemented in the model and summarized in Table 2 as R13-R18. Note that the products from the heterogeneous uptake of ClNO₂ on CPS vary with particle acidity (Riedel et al., 2012; Rossi, 2003). It generates Cl₂ under the condition of pH lower than 2 but produces nitrate and chloride under higher pH conditions. The reaction rates of the heterogeneous reactions are parameterized as first-order reactions, with the rate of change of gas phase species concentrations determined by equations (4) (Ying et al., 2015):

$$\frac{dC}{dt} = -\frac{1}{4}(v\gamma A)C = -k^{\mathrm{I}}C\tag{4}$$

where *C* represents the concentration of species; ν represents the thermal velocity of the gas molecules (m s⁻¹); *A* is the CMAQ-predicted wet aerosol surface area concentration (m² m⁻³); γ represents the uptake coefficient. For all gas phases species (except ClNO₂) involved in the heterogeneous reactions (R6 and R11-R18), a simple analytical solution can be used to update their concentrations from time t₀ to t₀+ Δ t: [C]_{t0+ Δ t}=[C]_{t0} exp(-k^I Δ t), where Δ t is the operator-splitting time step for heterogeneous reactions.

The rate of change of ClNO₂ includes both removal and production terms, as shown in equation (5):

$$\frac{d[\text{CINO}_2]}{dt} = -k_i^{\text{I}}[\text{CINO}_2] + k_6^{\text{I}}\phi_{\text{CINO}2}[N_2O_5]$$
(5)
= $-k_i^{\text{I}}[\text{CINO}_2] + k_6^{\text{I}}\phi_{\text{CINO}2}[N_2O_5]_{t0}\exp(-k_6^{\text{I}}t)$

Assuming ϕ_{CINO2} is a constant, an analytical solution for equation (5) can be obtained, as shown in equation (6):

$$[\text{CINO}_{2}]_{t0+\Delta t} = [\text{CINO}_{2}]_{t0} \exp(-k_{i}^{I}\Delta t) + \frac{k_{6}^{I}\phi_{\text{CINO2}}[N_{2}O_{5}]_{t0}}{k_{i}^{I}-k_{6}^{I}} [\exp(-k_{6}^{I}\Delta t) - \exp(-k_{i}^{I}\Delta t)]$$
(6)

where k_i^{I} represents the pseudo first-order rate coefficient of either reaction R17 or R18, depending on pH.

The uptake coefficients γ of gaseous species are obtained from published laboratorial studies. In the original CMAQ, the uptake coefficient of N₂O₅ is determined as a function of the concentrations of (NH₄)₂SO₄, NH₄HSO₄ and NH₄NO₃ (Davis et al., 2008). In this study, the PCl and NO₃⁻ dependent parameterization by Bertram and Thornton (2009) (equation (7)) is used:

$$\gamma_{N_2O_5} = \begin{cases} 0.02, & \text{for frozen aerosols} \\ 3.2 \times 10^{-8} K_f \left[1 - \left(1 + \frac{6 \times 10^{-2} [\text{H}_2\text{O}]}{[\text{NO}_3^-]} + \frac{29 [\text{Cl}^-]}{[\text{NO}_3^-]} \right)^{-1} \right] \end{cases}$$
(7)

In the above equation, $K_{\rm f}$ is parameterized function based on molarity of water: $K_f = 1.15 \times 10^6 (1 - e^{-0.13[{\rm H_2}O]})$. NO₃⁻ and Cl⁻ concentrations are also in molarity. The uptake coefficient of OH is expressed in equation (8) as a function of the concentration of PCl following the IUPAC (International Union of Pure and Applied Chemistry, available at http://iupac.poleether.fr/htdocs/datasheets/pdf/O-H_halide_solutions_VI.A2.1.pdf).

$$\gamma = \gamma_{\rm OH} = \min(0.04 \times \frac{[Cl^-]}{1000 \times M}, 1)$$
 (8)

where *M* represents the volume of liquid water in aerosol volume ($m^3 m^{-3}$). For frozen particles, the uptake coefficient is limited to 0.02, as used in the original CMAQ model.

The uptake coefficients of O₃, NO₃, NO₂, HOCl, ClNO₂, and ClONO₂ are treated 232 as constants. Among of them, the γ values of NO₃, NO₂, HOCl and ClONO₂ are set as 233 3×10^{-3} , 1×10^{-4} , 1.09×10^{-3} and 0.16 based on laboratory measurements (Rudich et al., 234 1996; Abbatt et al., 1998; Pratte et al., 2006; Gebel et al., 2001). A preliminary value 235 of 10^{-3} in the daytime and 10^{-5} at nighttime is chosen for the O₃ uptake coefficient. The 236 daytime γ_{O_3} is based on the analysis of Cl₂ production rate in a hypothesized 237 geochemical cycle of reactive inorganic chlorine in the marine boundary layer by Keene 238 et al. (1990). The lower nighttime value was also recommended by Keene et al. (1990) 239 who noted that Cl₂ production in the marine boundary layer are lower at night. The 240 uptake coefficient of ClNO₂ depends on the particle acidity, with the value of 2.65×10^{-10} 241 ⁶ for reaction R17 and 6×10^{-3} for reaction R18 (Robert et al., 2008). 242

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244 2.4 CMAQ model configuration

These heterogeneous reactions of chlorine are incorporated into a revised CMAQ based on the CMAQ version 5.0.1 to simulate the distribution of nitrate concentration in Beijing from 11 to 15 June 2017. The revised CMAQ model without heterogeneous reactions of chlorine has been described in detail by Ying et al. (2015) and Hu et al. (2016, 2017). In summary, the gas phase chemical mechanism in the revised CMAQ model is based on the SAPRC-11 (Cater et al., 2012) with a comprehensive inorganic chlorine chemistry. Reactions of Cl radical with several major VOCs, which lead to

production of HCl, are also included. The aerosol module is based on AERO6 with an 252 updated treatment of NO₂ and SO₂ heterogeneous reaction and formation of secondary 253 organic aerosol from isoprene epoxides. Three-level nested domains with the 254 resolutions of 36km, 12km, and 4km using Lambert Conformal Conic projection 255 $(173 \times 136, 135 \times 228 \text{ and } 60 \times 66 \text{ grid cells})$ are chosen in this work (the domains see 256 Figure S1). The two true latitudes are set to 25°N and 40°N and the origin of the domain 257 is set at 34°N, 110°E. The left-bottom coordinates of the outmost domain are positioned 258 at x = -3114 km, y = -2448 km. The BASE case (heterogeneous reactions of Cl turned 259 260 off) and HET case (all heterogeneous reactions enabled) are compared to evaluate the impact of heterogeneous chlorine chemistry on nitrate formation. 261

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263 **3. Results**

264 3.1 Model performance evaluation

Predicted O₃, NO₂ and PM_{2.5} concentrations from the BASE case simulation are 265 evaluated against monitoring data at 12 sites in Beijing (Table S2) in 11 to 15, June 266 267 2017. The average NMB/NME values for O₃, NO₂ and PM_{2.5} across the 12 sites are -8%/29%, -7%/59% and -8%/53%, respectively. Predicted hourly Cl₂, ClNO₂ and N₂O₅ 268 concentrations were compared with observations measured at the Institute of 269 270 Atmospheric Physics (IAP), Chinese Academy of Sciences (39.98°N, 116.37°E) using a high-resolution time-of-flight chemical ionization mass spectrometer (CIMS) from 11 271 to 15 June 2017 (for site description, instrument introduction, and analytical method, 272 please refer to the study by Zhou et al. (2018)). Figure 1 shows that the concentrations 273 of Cl₂ and ClNO₂ in BASE case are rather low (close to 0), proving that the gas-phase 274 chemistry is not the major pathway to produce Cl₂ and ClNO₂. By contrast, the 275 simulated Cl₂ and ClNO₂ concentrations in HET case increase significantly, 276 correspondingly the NMB and NME changes from -100% to -54% and 100% to 61% 277 for Cl₂, and from -100% to -58% and 100% to 62% for ClNO₂, respectively (the particle 278 surface area concentrations is scaled up by a factor of 5 in daytime and 10 in nighttime 279 280 because this parameter is underestimated compared to the measured concentrations reported by Zhou et al. (2018)). The simulations of Cl₂ and ClNO₂ are improved as the 281 additional heterogeneous reactions prompt the production of gas phase molecular 282 chlorine. Overall, however, the Cl₂ and ClNO₂ concentrations are still underestimated. 283 284 Both BASE and HET simulations generally capture the hourly N₂O₅ concentrations as

well as the peak values (Figure 1(c)) with similar overall NMB and NME values.

- The uptake coefficient of O_3 could be an important factor affecting the predicted Cl₂ concentrations as it is found that the heterogeneous reaction of O_3 is the major source of Cl₂ during this period (see discussion in Section 3.2). The influence of different parametrizations of the uptake coefficient of N₂O₅ on ClNO₂ and nitrate concentrations are also discussed in Section 3.2.
- 291 Predicted NO₃⁻ and PCl concentrations are compared with observations measured 292 at an adjacent monitoring site located at the rooftop of School of Environment building 293 in Tsinghua University (THU, 40.00°N, 116.34°E, about 5 km from IAP) using an Online Analyser of Monitoring of Aerosol and Gases (MARGA) from 11 to 15 June 294 2017. According to Figure 1(d), the simulated nitrate concentration is slightly lower 295 than the observations most of the time. From the evening hours of 12 June to morning 296 hours of 13 June, observed and simulated nitrate concentration both increase 297 significantly. The NMB and NME values of hourly nitrate for the HET case (-5% and 298 39%, respectively) are slightly lower than those for the BASE case -10% and 46%) 299 300 during this high concentration period. The HET case also generally captures the dayto-day variation of PCl concentration and perform better than the BASE case, 301 302 correspondingly the NMB and NME are reduced from -48% and 72% to -37% and 67%. The substantial underestimation of PCl in the daytime on 15 June is likely caused by 303 304 missing local emissions during this period.
- 305

306 3.2 Impact of uptake coefficients of O₃ and N₂O₅ on chlorine species and nitrate

The uptake coefficients of O₃ and N₂O₅ may be important factors affecting the accuracy 307 of simulated nitrate concentrations. Some studies have confirmed that the reaction of 308 O₃ on CPS can indirectly affect the nitrate formation by increasing the atmospheric Cl₂ 309 and OH level (Li et al., 2016; Liu et al., 2018). According to Figure 1(a), the improved 310 model still substantially underestimates the concentration of Cl₂, which may be 311 associated with the underestimation of the uptake coefficients of O₃, which are 312 empirical and have not been confirmed by laboratory studies. The uptake coefficients 313 were increased by a factor of 10 (0.01 for daytime and 10^{-4} for nighttime) to evaluate 314 the sensitivity of Cl₂ production and nitrate formation to this parameter. Figure 2 shows 315 that the simulated Cl_2 and nitrate concentrations in daytime increase significantly 316 (especially for Cl₂) and sometimes can capture the peak value (such as the daytime peak 317 on 14 June). However, although the NMB and NME of Cl₂ and nitrate improve from -318

319 18% and 39% to 1% and 28% when the new uptake coefficients are used, the simulated 320 Cl_2 concentrations are still quite different from the observations (such as during the 321 daytime in 11 and 12 June, see Figure 2). A non-constant parameterization of the uptake 322 coefficients of O₃ that considers the influence of PCl concentrations, meteorology 323 conditions, etc., similar to those of OH and N₂O₅, might be needed. Further laboratory 324 studies should be conducted to provide a better estimation of this important parameter.

Several parameterizations for the uptake coefficient of N₂O₅ have been developed 325 326 for regional and global models and have been evaluated in several previous studies 327 (Tham et al., 20192018, McDuffie et al., 2018a, 2018b). In addition to the parameterization of Bertram and Thornton (2009) used in the HET case, two additional 328 simulations were performed to assess the impact of the uptake coefficient of N₂O₅ on 329 nitrate formation. The first simulation uses the original CMAQ parameterization of 330 Davis et al.(2008) and second simulation uses a constant value of 0.09, which is the 331 upper limit of the N₂O₅ uptake coefficient derived by Zhou et al. (2018) based on 332 observations. The results from the simulations with the parameterization of Bertram 333 and Thornton (2009) generally agree with the results using those based on Davis et al. 334 (2008) and the. The application of larger constant and fixed N₂O₅ uptake coefficient 335 336 leads to slightly better results, which might reflect the fact that the N₂O₅ concentrations are underestimated. Using the uptake coefficient of 0.09 can generally increase the 337 concentration of nitrate in some periods, but it also leads to significant increase of the 338 nitrate level (such as nighttime on 12-13 June and 13-14 June), which is 4-6 times 339 higher than those based on Bertram and Thornton (2009). Overall, predicted nitrate 340 concentrations are sensitive to changes in the changes in $\gamma_{N_2O_5}$, with approximately 50% 341 increase in the nitrate when a constant of $\gamma_{N_2O_5}$ of 0.09 is used. 342

343

344 3.3 Spatial distributions of nitrate and chlorine species concentrations

The regional distributions of averaged Cl_2 , $ClNO_2$, N_2O_5 and NO_3^- concentration from 11 to 15 June for the HET case are shown in Figure 3. Compared to the BASE case, the episode average concentrations of Cl_2 and $ClNO_2$ from the HET case increase significantly in the eastern region of Beijing, reaching up to 23 ppt and 71 ppt from near zero (Figure 3a and 3b). High concentrations are not found in the southern region with intensive emissions of chlorine species (Figure S2). The production of $ClNO_2$ requires the presence of chloride, NO_2 , and O_3 . In the areas close to the fresh emissions, O_3 is generally low (Figure S3), and the production of NO₃ (hence N₂O₅ and ClNO₂) is limited. Therefore, the production rate of ClNO₂ is generally low in areas affected by fresh emissions. Since the contribution of direct emissions to Cl_2 is low and it is predominantly produced secondarily in the atmosphere, high levels of Cl_2 are also found away from the fresh emissions.

357 The spatial distribution of N_2O_5 concentrations differs from that of other species (Figure 3c). While the concentrations of most of the species are higher in the southern 358 region, the N₂O₅ concentrations are lower in some parts of this region. This is because 359 360 the O₃ concentration in the core urban areas is low due to high NO_x emissions. The N₂O₅ concentrations from the HET case are approximately 16% lower on average 361 (Figure 3d) because the Bertram and Thornton (2009) parameterization used in the HET 362 case generally gives higher uptake coefficients than the parameterization of Davis et al. 363 (2008) used in the BASE case (Table 3). 364

365 Although the higher uptake coefficients of N₂O₅ in the HET case facilitate faster conversion of N₂O₅ to nitrate, the nitrate concentrations do not always increase. During 366 367 daytime hours, nitrate concentrations in the HET case increase due to higher OH (Figure 3e and Figure 3f), increased OH see Figure S4). At nighttime, in contrast, the nitrate 368 369 concentration decreases significantly in some regions by about 22%, mainly due to lower molar yield of nitrate from the N₂O₅ heterogeneous reaction in the HET case 370 (Figure 3g and Figure 3h). Although ClNO₂ produced in the N₂O₅ reaction also 371 372 produces nitrate through a heterogenous reaction when the particle pH is above 2, which 373 is true for most regions (see Figure \$4\$5), the uptake coefficient of ClNO₂ is significantly lower than that of N₂O₅ (0.01 \sim 0.09 for N₂O₅ and 6 \times 10⁻³ for ClNO₂), 374 375 leading to an overall decrease of nitrate production. As the ClNO₂ production from the heterogeneous reaction leads to less N₂O₅ conversion to non-relative nitrate, it may 376 change the overall lifetime of NOx and their transport distances. The magnitude of this 377 change and its implications on ozone and PM2.5 in local and downwind areas should be 378 379 further studied.

380

381 3.4 Relationship between nitrate formation and chlorine chemistry

382 Nitrate productions from the homogeneous and heterogeneous pathways in Beijing are

383 approximated by the difference in predicted nitrate concentrations between the BASE

384 or HET case and a sensitivity case without heterogenous reactions. Averaging over the

five-day period, approximately 58% of the nitrate originates from HNO₃ gas-to-particle

partitioning and 42% is from heterogeneous reactions (Figure 4). This conclusion generally agrees with measurements at Peking University (PKU) (52% from the heterogeneous process and 48% from HNO₃ partitioning) on four polluted days (average in September 2016 reported by Wang et al. (2017). Slightly higher contributions of the homogeneous pathway in this study is expected because of high OH concentrations during the day and lower particle surface areas at night.

392 The nitrate formation from different homogeneous and heterogeneous pathways in the BASE case and HET case are further studied. Contributions of different gas phase 393 394 pathways are determined using the process analysis tool in CMAQ. Contributions of different heterogeneous pathways are determined using a zero-out method that turns of 395 one heterogeneous pathway at a time in a series of sensitivity simulations. Figure 4 396 shows that the reaction of OH and NO₂ is always the major pathway for the formation 397 of nitrate through homogeneous formation of HNO₃ and gas-to-particle partitioning. 398 However, its nitrate production rate through this homogeneous pathway decreases 399 significantly from daytime to nighttime (from 1.81 μ g m⁻³ h⁻¹ to 0.33 μ g m⁻³ h⁻¹ on 400 average). The nitrate production from other HNO₃ partitioning pathways in the daytime 401 is negligible. At nighttime, homogeneous reaction of N₂O₅ with water vapor accounts 402 403 for approximately 5% of the overall homogeneous nitrate formation. For the heterogeneous pathways, daytime production rate is approximately 0.6 µg m⁻³ h⁻¹ with 404 1/3 of the contributions from NO₂ and 2/3 from N₂O₅. Nighttime production on nitrate 405 from the heterogeneous pathways is approximately 3.1 μ g m⁻³ h⁻¹, of which 85% is due 406 407 to N_2O_5 and 15% is due to NO_2 .

Comparing the BASE case and the HET case shows that, when the chlorine 408 chemistry is included, the gaseous HNO₃ produced by OH reacting with NO₂ increases 409 significantly in the HET case. Correspondingly, the nitrate production rate reaches up 410 to 2.04 μ g m⁻³ h⁻¹ in the daytime due to increased atmospheric OH concentrations 411 predicted by the chlorine reactions. Similar conclusions are also obtained by Li et al. 412 (2016) and Liu et al. (2017) based on observations and model simulations. The 413 heterogeneous production of nitrate from the reaction of N₂O₅ uptake decreases by 414 approximately 27% in the HET case due to the production of gas phase ClNO₂. 415 According to the study by Sarwar et al. (2012; 2014), including the heterogeneous 416 reaction of N₂O₅ with PCl decreased the nocturnal nitrate concentration by 11-21% in 417 the United States, which was slightly less than the current study for Beijing. It is likely 418 because PCl concentrations in the United States are significantly lower than those in 419

420 Beijing (the monthly PCl concentration is 0.06 μ g m⁻³ in the United State against ~1 μ g

- 421 m^{-3} in Beijing) so that PCl is depleted quickly. The contributions of NO₂ uptake to
- 422 nitrate also decrease by 22% because of the lower rate constant of the reaction of NO₂
- 423 with PCl. In contrast, the contribution of ClNO₂ reacts with particle surface to nitrate
- 424 production increases by 6% in the HET case. The overall nitrate concentration in the
- 425 HET case is about 22% higher than that in the BASE case during this study period.
- 426

427 **4. Conclusions**

428 In this work, a modified CMAQ model incorporated with heterogeneous reactions for the production of molecular chlorine and other reactive chlorine species is developed 429 and its impact on of the nitrate formation predictions are evaluated. The contributions 430 from different homogenous and heterogeneous pathways to nitrate formation are also 431 quantified. High concentration of Cl₂ and ClNO₂ do not occur in the southern part of 432 the Beijing-Tianjin-Hebei region with intensive emissions of chlorine species as higher 433 concentrations of O₃ and N₂O₅ associated with the heterogeneous formation of these 434 species generally occurred in the downwind areas. CTMs without a complete treatment 435 of the chlorine chemistry can underestimate daytime nitrate formation from the 436 437 homogeneous pathways, particularly from HNO₃ gas-to-particle partitioning due to underestimation of OH concentrations and overestimate the nighttime nitrate formation 438 from the heterogeneous pathways due to missing chlorine heterogeneous chemistry. 439

440

441 *Data availability*. The data in this study are available from the authors upon request 442 (shxwang@tsinghua.edu.cn)

443

Author contributions. XQ, QY, SW and JH designed the study; YS, BL, AS, XY
provided observation data; XQ, QY, SW, JZ, QX, DD, LD and JX analyzed data. XQ,
QY and SW wrote the paper.

- 447
- 448 *Competing interests*. The authors declare that they have no conflict of interest.
- 449

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Figure 1 Comparison of observed hourly Cl₂, ClNO₂, N₂O₅ (at the Institute of
Atmospheric Physics, Chinese Academy of Sciences), NO₃⁻ and PCl (at Tsinghua
University) in urban Beijing with predictions from the BASE and the HET cases

593 University) in urban Beijing wit594 during 11-15 June 2017.



Figure 2 Comparison of observed and predicted Cl₂ and NO₃⁻ concentrations under different uptake coefficient of O₃ (HET: daytime $\gamma_{O_3} = 1 \times 10^{-3}$, nighttime $\gamma_{O_3} = 1 \times 10^{-5}$; HET-Co: daytime $\gamma_{O_3} = 1 \times 10^{-2}$, nighttime $\gamma_{O_3} = 1 \times 10^{-4}$).



600

Figure 3 Spatial distributions of episode-average (a) Cl_2 , (b) $ClNO_2$, (c) N_2O_5 , (e) daytime nitrate (ANO₃) and (g) nighttime nitrate concentrations from 11-15 June 2017, and the differences in the episode-average (d) N_2O_5 (HET case – BASE case), (f) daytime nitrate and (g) nighttime nitrate. Units are $\mu g m^{-3}$.



606 607 Figure 4 Contribution 608 formation.

Saatan	Emissions						
Sector	HC1	Cl ₂	PC1				
Power plant	22.8	1.2	6.75				
Industry	587.3	20.1	89.2				
Residential	202.4	8.1	34.7				
Biomass burning	0.182	0	0.14				
Municipal solid waste	1080.2	0	8.47				
Cooking	0	0	426.8				
Total	1892.9	29.4	566.1				

Table 1 The sectoral emissions of HCl, Cl₂ and PCl in Beijing in 2017. Unit: Mg year⁻¹

- Table 2 Major gas-phase and heterogeneous pathway of producing nitrate in original CMAQ and newly added or revised heterogeneous reactions
- 614 in improved CMAQ.

Туре	Reactions	No.	Reference	Comment
Original CMAQ				
Gas-phase chemistry	$OH + NO_2 \rightarrow HNO_3$	R1		
	$N_2O_5 + H_2O \rightarrow 2HNO_3$	R7		
	$HO_2 + NO_3 \rightarrow 0.2HNO_3 + 0.8OH + 0.8NO_2$	R8		
	$NO_3 + VOCs^a \rightarrow HNO_3$	R9		
Heterogeneous	$N_2O_5(g) + H_2O(aq) \rightarrow 2H^+ + 2NO_3^-$	R5		
chemistry	$2NO_2(g) + H_2O(aq) \rightarrow HONO(g) + H^+ + NO_3^-$	R10		
Improved CMAQ				
Newly added or	$N_2O_5(g) + H_2O(aq) + Cl^{-}(aq) \rightarrow ClNO_2(g) + NO_3^{-}$	R6	Bertram and Thornton (2009)	Revise R5
revised heterogeneous	$2NO_2(g) + Cl^- \rightarrow ClNO(g) + NO_3^-$	R11	Abbatt et al. (1998)	Revise R10
reactions	$NO_3(g) + 2Cl^- \rightarrow Cl_2(g) + NO_3^-$	R12	Rudich et al. (1996)	Increase NO ₃ -
	$O_3(g) + 2Cl^- + H_2O(aq) \rightarrow Cl_2(g) + O_2(g) + 2OH^-$	R13	Abbatt et al. (1998)	Affect OH
	$2OH^{-}(g) + 2Cl^{-} \rightarrow Cl_{2}(g) + 2OH^{-}$	R14	George et al. (2010)	Affect OH
	$\text{ClONO}_2(g) + \text{Cl}^- \rightarrow \text{Cl}_2(g) + \text{NO}_3^-$	R15	Deiber et al. (2004)	Affect OH
	$HOCl(g) + Cl^- + H^+ \rightarrow Cl_2(g) + H_2O$	R16	Pratte et al. (2006)	Affect OH
	$\operatorname{ClNO}_2(g) + \operatorname{Cl}^- + \operatorname{H}^+ \rightarrow \operatorname{Cl}_2(g) + \operatorname{HONO}(g) (pH < 2.0)$	R17	Riedel et al. (2012)	Affect OH
	$CINO_2(g) + H_2O(aq) \rightarrow Cl^- + NO_3^- + 2H^+ (pH \ge 2.0)$	R18	Rossi (2003)	Increase NO ₃ -

^a: presents different VOCs species. In the SAPRC-11 mechanism, the VOCs species include CCHO (Acetaldehyde), RCHO (Lumped C3+
 Aldehydes), GLY (Glyoxal), MGLY (Methyl Glyoxal), PHEN (phenols), BALD (Aromatic aldehydes), MACR (Methacrolein), IPRD (Lumped
 isoprene product species).

619Table 3 Observed day (D) and night (N) NO3⁻ concentrations (Obs.) and predicted620uptake coefficient of N2O5 (γ_{N2O5}) and nitrate concentrations (Pred.) using the621parameterizations of γ_{N2O5} by Bertram and Thornton (2009) (Scenario 1), Davis et al.,622(2008) (Scenario 2) and the upper-limit value derived by Zhou et al. (2018) (Scenario6233)

	NO ₃ -	Scenario1		Scer	nario2	Scenario3	
	Obs.	γN2O5	Pred.	γn205	Pred.	γn205	Pred.
06/11-D	2.54	0.033	1.59	0.008	1.32	0.09	2.17
06/11-12-N	2.42	0.043	1.67	0.037	1.37	0.09	2.12
06/12-D	3.39	0.028	2.16	0.032	2.74	0.09	3.13
06/12-13-N	4.24	0.021	4.02	0.022	4.05	0.09	6.04
06/13-D	2.57	0.012	1.18	0.008	1.06	0.09	2.47
06/13-14-N	4.10	0.022	4.45	0.022	4.45	0.09	7.13
06/14-D	0.95	0.001	1.34	0.001	1.33	0.09	1.64
06/14-15-N	2.75	0.013	1.00	0.007	0.96	0.09	2.33
06/15-D	0.75	0.001	0.66	0.001	0.66	0.09	1.11