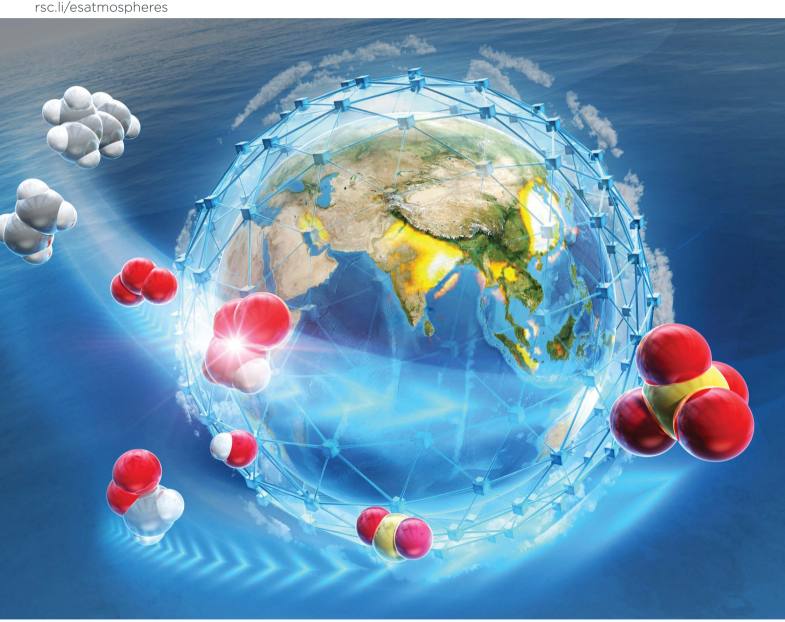
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Global-scale analysis of the effect of gas-phase Criegee intermediates (CIs) on sulphate aerosol formation: general trend and the importance of hydroxy radicals decomposed from vibrationally excited CIs†

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Stabilised Criegee intermediates (sCls), which are formed in the atmosphere through the ozonolysis of alkenes, are known precursors of sulphate aerosols $(SO_4^{2-}(p))$. Several previous studies have focused on the kinetics of sCI-related chemistry using both experimental and theoretical methods. Nonetheless, detailed evaluations of how the sCl affects global-scale $SO_4^{2-}(p)$ formation using chemical transport models (CTMs) have rarely been conducted. In this study, the impact of sCIs on $SO_4^{2-}(p)$ and other particulate matter was estimated using a global CTM by implementing approximately 100 chemical reactions associated with CI chemistry. The results suggest that sCIs contribute maximally less than 0.5% in remote areas, such as Amazon rainforests, Central Africa, and Australia. This value is lower than the previously estimated value, despite certain kinetic parameters related to CI chemistry being provisional due to insufficient data. Future work should focus on obtaining these kinetic parameters through experimental studies or theoretical calculations. The sCI that contributed the most to $SO_4^{2-}(p)$ formation was E-methyl glyoxal-1-oxide, which was generated by the ozonolysis of methyl vinyl ketone owing to its low-rate coefficient for the loss reaction of unimolecular decomposition and water vapour. The change in $SO_4^{2-}(p)$ enhanced the formation of secondary organic aerosols, whereas the reactions of the sCls with NO₂ decreased the formation of nitrate radicals. The results of the sensitivity analyses showed that in highly industrialised sites in China and India, OH radicals formed by the unimolecular decomposition of vibrationally excited CIs (vCIs) contributed to $SO_4^{2-}(p)$ formation, which maximally accounted for nearly ten times more than that of sCls, whereas the contribution of vCls and sCls to $SO_4^{2-}(p)$ formation was estimated to be almost equal in rural and remote sites. The estimated sCI loss by HNO₃ and organic acids was comparable to that of the unimolecular decomposition of sCIs and scavenging by water. This study provides full insight into the impact of gas-phase CI chemistry on a global scale.

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Environmental significance

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Atmospheric aerosols pose health risks to humans and animals and are also recognised as climate forcers. Therefore, understanding the fundamental chemistry of aerosol formation is crucial for the scientific community to address these issues. Criegee intermediates (CIs) are known precursors of sulphate and other atmospheric aerosols. While numerous studies have clarified the role of stabilised Criegee intermediates (sCIs) in aerosol formation, the impact of vibrationally excited Criegee intermediates (vCIs), which produce OH radicals through unimolecular decomposition, has been less explored. This study quantifies the significance of vCIs in aerosol formation and provides insights into Criegee chemistry using a global chemical transport model.

Introduction

Sulphate aerosols (SO₄²⁻(p)) are known as the nuclei of fine particulate matter (PM2.5), which has a negative impact on human health¹⁻³ and contributes to the positive and negative radiative forcing.4-6 As described in Fig. 1, the ozonolysis of alkenes forms primary molozonide (POZ) and decomposes to

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Fig. 1 Reactions of alkene ozonolysis and formation of CI.

vibrationally excited Criegee intermediates (vCIs) and carbonyl species.7-11 The vCIs then decompose to form hydroxy radicals (OH) or release energy through quenching in ambient air to form stabilised Criegee intermediates (sCIs).7-13 The sCIs are known to be strong oxidisers in the atmosphere, reacting with several air pollutants such as sulphur dioxide (SO2), nitrogen oxides (NO_x), and organic acids.⁷⁻¹¹ In particular, since the rate coefficient of the reaction between sCIs and SO2, which finally transforms into SO₄²⁻(p), was estimated to be high (approximately 10⁻¹¹ cm³ molecule⁻¹ s⁻¹), 7-11 sCIs have been considered as one of the main contributors of atmospheric $SO_4^{2-}(p)$ formation, and several studies analysed the impact of sCIs on SO₄²⁻(p) formation on both regional and global scales using chemical transport models (CTMs). Sarwer et al. included sCIrelated chemical reactions to the CTM and estimated the contribution of sCIs to SO₄²⁻(p) formation, concluding that sCIs contributed 20% to 75% of total $SO_4^{2-}(p)$ formation in the U.S.14 Percival et al. conducted regional and global analyses of the effect of sCIs on SO₄²⁻(p) formation, and they concluded that the contribution of sCIs is as high as that of gas-phase oxidation of OH, especially in urban areas, although the global CTM could not resolve the impact of sCIs on SO₄²⁻(p) formation in the urban area due to the averaged emissions in each calculated grid.15 On the other hand, Nakamura et al. reported negligible contributions of sCIs to $SO_4^{2-}(p)$ formation in the Greater Tokyo Area (GTA), the metropolitan area of Japan.¹⁶ The estimated amount of sCIs in GTA was consistent with a previous study that estimated the amount of sCIs in the urban area of the U.K.17 Our previous study suggested that isoprenederived sCIs enhance SO₄²⁻(p) formation in the forestry area of the Asian region, which was comparable to the rate of aqueous-S(IV) chemistries,18 although Newland et al. suggested the less importance of isoprene-derived sCIs to SO₂ oxidation on a global scale.19 Vereecken et al. estimated several rate coefficients related to sCIs using quantum chemical calculations combined with transition state theory and showed the importance of unimolecular decompositions of sCIs to the total evaluation of sCI-related pollution by the global CTM.20 Other several studies also evaluated the impact of sCIs on atmospheric pollutants, such as SO₂, organic acids, and peroxy radicals, by the CTM, sometimes combined with reaction rate coefficients derived from experiments and calculations using transition state theory.21-25

The impact of the CI chemistry on the atmosphere has not been analysed in detail. Most previous studies have investigated the atmospheric implications of the newly estimated rate coefficients or the impact of specific kinetics by applying the global CTM, especially focusing on the impact of sCIs on ${\rm SO_4}^{2-}(p)$ formation. However, their impact on other aerosols has rarely

been investigated. In addition, vCIs may have a more significant impact on $SO_4^{2-}(p)$ formation than sCIs. Khan *et al.* suggested that the OH formation through the unimolecular decomposition of vCIs contributes maximally 13% of total OH formation in the terrestrial rainforest and boreal forest regions. ¹⁰ Also, Nakamura *et al.* ¹⁶ implied that the formation of $SO_4^{2-}(p)$ due to incorporation of sCIs was mainly due to the increase in OH concentration caused by the unimolecular decomposition of vCIs, not the direct oxidation of SO_2 by sCIs. Further studies should be conducted to investigate the role of vCIs in $SO_4^{2-}(p)$ formation through a comparative analysis of the contributions of sCIs and OH derived from vCIs.

This study aimed to reveal the impact of CI chemistry on SO₄²⁻(p) and other aerosol formation on a global scale and to compare the impact of sCIs and vCIs on SO₄²⁻(p) formation. This study comprised three steps: first, we applied the detailed sCI-related chemical mechanisms proposed by Nakamura et al., 16 most of which were originally based on the reactions of Vereecken et al. 20 to the global CTM to evaluate the effect of sCIs on $SO_4^{2-}(p)$ formation on a global scale. Second, we investigated sCI impact on ionic and SOA formation in addition to SO₄²⁻(p) to show the cross-effect of sCIs on other particulate species. Third, we conducted a sensitivity analysis to compare the contribution of $SO_4^{2-}(p)$ formation between sCIs and OH derived from the unimolecular decomposition of vCIs and the contribution of other sCI loss reactions. Although there have been many previous studies related to the impact of gas-phase reactions of sCIs on the atmosphere using the CTM, this study contributes to providing new insights into the atmospheric contribution of sCIs and vCIs and the fundamental information of the whole CI chemistry.

Methodology

Global chemical transport modelling

The Goddard Earth Observing System Chemistry model (GEOS-Chem v14.2.3)26 was used as the global CTM. The modern era retrospective analysis for research and applications version 2 (MERRA2) was used as the meteorological input with a grid resolution of $2^{\circ} \times 2.5^{\circ}$ (latitude × longitude) holding 72 vertical layers from the troposphere to the stratosphere.27 Emission inventories provided by the harmonized emission component (HEMCO) system were applied in this study.28 Briefly, the emission database for the community emission data system (CEDSv2) was chosen for anthropogenic emissions,29 hemispheric transport of air pollution (HTAPv3) was chosen for ship emissions,30 monthly mean data of the Aviation Emissions Inventory (AEIC2019) was chosen for aircraft emissions, 31 global fire emission database (GFED) was chosen for biomass burning,32 Aerosol Comparisons between Observations and Models (AeroCom) was selected for the SO₂ emissions from volcanoes,33 and the off-line model of emissions of gases and aerosols from nature (MEGAN) was chosen for BVOC emissions.34 The targeted period calculated in this study spanned from 1 July 2018 to 31 December 2019. The first six months (1 July 2018 to 31 December 2018) were treated as the spin-up period, and the entire year of 2019 was treated as the analysis

term. The reactions of the Criegee intermediates (CIs) were added to the full chemistry module of the kinetic pre-processor (KPP). The added CI reactions were based on Nakamura et al., who analysed the impact of sCIs on $SO_4^{2-}(p)$ formation in the GTA of Japan. 16 Nakamura et al. cited several reaction kinetics from the findings of Vereecken et al., who determined the rate coefficients of sCI atmospheric reactions using quantum chemical calculations coupled with transition state theory.20 The default KPP has already implemented the chemical reactions of CIs generated by the ozonolysis of ethylene and propene. In this study, those CI chemistries were excluded to unify the reactions applied in Nakamura et al.16 The details of the added reactions are described in a later section.

Validation of the modelling results using satellite data

Validation of modelling results on a regional scale was typically conducted through comparison with ground observations. However, considering the coarse resolution of our analysis (2° × 2.5°), a direct comparison with ground observations is challenging. The results were validated by comparison with satellite observations from the tropospheric monitoring instrument (TROPOMI).35-37 Alkenes, O3, and SO2 are the main species forming sCIs and SO₄²⁻(p), and comparing these species would be ideal for model validation. Owing to the lack of satellite observations of alkenes, HCHO, which has been widely used as an indicator of VOC emissions (including alkenes), 38 was used, along with tropospheric O₃ and SO₂. Level 2 data for HCHO, tropospheric O₃, and SO₂ were retrieved from COPERNICUS (O₃; spatial resolution: $30 \times 30 \text{ km}^2$)³⁹ and NASA Earth Observation Data (HCHO and SO_2 ; special resolution: $7 \times 3.5 \text{ km}^2$).40 Datasets were screened based on the recommended quality flags (<0.5) to produce Level 3 data, solar zenith angle < 85°, and cloud fraction < 30%.39 The TROPOMI data were re-gridded to equalize the spatial resolution to that of GEOS-Chem $(2^{\circ} \times 2.5^{\circ})$ via linear interpolation of the targeted grids. TROPOMI provides daytime data on the column densities of pollutants, which were compared with the daily average values calculated by GEOS-Chem. The determination factor (R^2) , root mean square error (RMSE), normalised mean bias (NMB), and normalised mean error (NME) were chosen as statistical indicators to verify the correlation between the observed and calculated results. Globally, five sites were selected for comparison: China (CHI), India (IND), South Africa (ZAF), Australia (AUS), and Bolivia (BOL: corresponding to the Amazon rainforests), where the impacts of sCIs were estimated to be high according to this study, the details of which are described in a later section. Detailed information on the five sites is provided in Table S1 of the ESI.†

Incorporated chemical reactions

The chemical mechanisms of the ozonolysis reactions of 8 alkenes (ethylene (C₂H₄), propene (PRPE), methyl vinyl ketone (MVK), methacrolein (MACR), isoprene (ISOP), pinene (MTPA), other monoterpenes (MTPO), and limonene (LIMO)), unimolecular reactions sCIs, and eight types of bimolecular reactions between sCIs and air pollutants (water monomer (H2O), water

dimer ((H2O)2), nitrogen dioxide (NO2), sulphur dioxide (SO2), nitric acid (HNO₃), formic acid (HCOOH), acetic acid (CH₃-COOH), and organic acids with more than C3-chains (RCOOH)) were incorporated into KPP. There were eight types of sCIs generated from alkene ozonolysis: formaldehyde oxide (SCI1), acetaldehyde oxide (SCI2), methyl vinyl ketone oxide (SCI3), methacrolein oxide (SCI4), two types of pinene oxides (SCI5 and SCI6), methyl glyoxal-1-oxide (SCI7), and methyl glyoxal-2-oxide (SCI8), of which all the sCIs aside from SCI1 are E- and Zisomers. Detailed structures of the sCIs are shown in Fig. S1 of the ESI.† Most of the chemical mechanisms and rate coefficients are cited from Nakamura et al. 16 Additionally, they analysed the effect of sCIs on $SO_4^{2-}(p)$ formation in the GTA using the community multiscale air quality modelling system (CMAQ)41 with the SAPRC-07 chemical mechanism,42 and several expressions of the chemical species were different from those of KPP. The corresponding species were chosen to substitute the SAPRC-07 components with KPP. The added reactions are summarised in Tables S2 to S10 of the ESI.†

Alkene ozonolysis

Table S11 of the ESI† lists the incorporated alkene ozonolysis reactions and rate coefficients whose information is originally cited from Master Chemical Mechanism (MCM).43 The default reactions of alkenes and O3 defined in KPP were substituted by the reactions listed in Table S11† because MCM has more detailed chemistries of sCI generation. The 15 sCI structures shown in Fig. S1 of the ESI,† including E- and Z-isomers of several sCIs, were generated by ozonolysis of the alkenes. The branching ratio of E- and Z-isomers was assumed to be 0.5.16,18 The alkene ozonolysis reactions leading to the generation of sCIs and OH are shown in Fig. S1.† For example, ethylene ozonolysis is defined as follows:

$$C_2H_4 + O_3 \rightarrow 0.37SCI1 + 0.13OH + ...$$
 (R1)

The formation of OH by (R1) can be considered the end product of the unimolecular decomposition of vCIs.

Unimolecular decomposition of sCIs

Unimolecular decomposition of sCIs is one of the major pathways of sCI loss,²⁰ and Table S2† lists the chemical mechanisms of the unimolecular reaction of sCIs and their kinetic parameters. Three types of unimolecular decompositions, 1,3-ring closure, 1,4-H-migration, and 1,5-ring closure, were considered, and the details of these reactions are described by Vereecken et al.20

Reactions of sCIs with H₂O and (H₂O)₂

The reactions of sCIs with water monomers (H2O) and dimers $((H_2O)_2)$ are also one of the major paths for the loss of sCIs; Tables S3 and S4 of the ESI† list the chemical mechanisms and kinetic parameters of the reaction. Note that the kinetic parameters of the reactions of sCIs and (H2O)2 depend on the ratio of H₂O to (H₂O)₂ in the atmosphere, which is described by the equilibrium constant K_w , expressed by eqn (1) according to a previous study.⁴⁴

$$K_{\rm w} = \frac{\left[({\rm H_2O})_2 \right]}{\left[{\rm H_2O} \right]^2}$$
 (1)

Further details of the derivation of the rate coefficients listed in Table S4† have been described in a previous study. 16

Reactions of sCIs with SO2, NO2, HNO3, and organic acids

Tables S5–S11† show the incorporated reactions and kinetic parameters of the reactions between sCIs and SO₂, NO₂, HNO₃, and organic acids. The main difference from Nakamura et~al. is the definition of the reactions between sCIs and NO₂. In a previous study, ¹⁶ the products of sCIs + NO₂ reactions were determined as NO₃ and aldehydes (or ketones). Caravan et~al. ⁴⁵ found out that the main product of the reaction was the sCIs–NO₂ adduct, and only approximately 30% of sCIs–NO₂ adduct decomposed to NO₃ and aldehydes (or ketones), indicating that NO₃ formation through the sCIs + NO₂ reaction was overestimated by Nakamura et~al. ¹⁶ For this reason, in this study, we multiplied 0.3 with NO₃ and aldehyde (or ketone) as the stoichiometric factor to incorporate the findings of Caravan et~al. ⁴⁵

Sensitivity analysis to compare the contributions of sCIs and vCIs in $SO_4^{2-}(p)$ formation

Sensitivity analysis was conducted to reveal the important CIrelated reaction scheme for $SO_4^{2-}(p)$ formation. Seven scenarios were used to evaluate sensitivity: unimolecular decomposition of vCIs (S1), reactions of sCIs with SO₂ (S2), unimolecular decomposition of sCIs (S3), reactions of sCIs with water (S4), reactions of sCIs with NO₂ (S5), reactions of sCIs with HNO₃ (S6), and reactions of sCIs with organic acids (S7). The reactions evaluated in scenarios S1 and S2 contributed to $SO_4^{2-}(p)$ formation through SO_2 oxidation by OH or sCIs. The other scenarios (S3-S7) were loss reactions of sCIs, which indirectly contributed to $SO_4^{2-}(p)$ formation. Note that although scenario S1 focuses on the contribution of vCIs to SO₄²⁻(p) formation, the exact species of vCIs are not defined by the chemical mechanism of this study. This is because vCIs are intermediate species in the ozonolysis reaction of alkenes and are either immediately decomposed into OH or stabilised through collisions with bath gas to form sCIs. Thus, scenario S1 only calculates the total sensitivity to ozonolysis minus the sensitivity to the reactions of sCIs. To evaluate the contribution of each process to $SO_4^{2-}(p)$ formation, the sensitivity, S_i , of reaction scheme i was formulated as follows:

$$S_i = \frac{k_i}{\left[SO_4^{2-}(p)\right]} \frac{\delta\left[SO_4^{2-}(p)\right]}{\delta k_i},$$
 (2)

where k_i is the rate coefficient of the targeted reaction scheme i and δk_i is the functional derivative of k_i . [SO₄²⁻(p)] refers to the column density of SO₄²⁻(p) in the scenario with Criegee chemistry and δ [SO₄²⁻(p)] refers to the change in [SO₄²⁻(p)] after the reaction rates were multiplied by $(1 + \delta k_i/k_i)$. In this study, the value of $\delta k_i/k_i$ was set to 0.1. For example, the

sensitivity of sCIs and SO₂ reactions to SO₄²⁻(p) formation was calculated by multiplying the rate coefficients of all sCIs + SO₂ reactions by 1.1 and investigating the change in the column density of SO₄²⁻(p). In terms of sensitivity scenario S1, the rate coefficients of the ozonolysis reactions of alkenes were multiplied by 1.1; however, this operation also led to the enhancement of later sCI-related reactions, which resulted in including both the effects of vCIs and sCIs on the sensitivities of $SO_4^{2-}(p)$ and other aerosol formation. To account for this problem, another scenario, S8, was calculated, which contained the operation of multiplying 1.1 by the rate coefficients of all sCIrelated chemical reactions. The results of scenario S1 were then subtracted by the results of scenario S8 to ensure that S1 held only the sensitivities of the vCIs to aerosol formation. In the Results and discussion section, the sensitivity of S1 contains the subtracted result for scenario S8.

Results and discussion

Evaluation of the modelling performance in comparison with the observed data of TROPOMI

Fig. 2 shows the relationships between the observed and modelled column densities of O3, HCHO, and SO2. According to Fig. 2(a) and (b), O₃ and HCHO showed high determination coefficients with $R^2 = 0.83$ and 0.57, respectively, while the value of R^2 for SO₂ was 0.37, indicating a moderate linear correlation between the observed and modelled results. The slopes of all three chemicals were less than 1.0, with SO₂ showing the lowest value, followed by O3 and HCHO. This suggests that the modelled results underestimated the observed results, especially in SO₂. One of the emission sources of SO₂ is anthropogenic activities, and thus the emission of SO2 is highly dependent on time. Fig. 2 shows a comparison between the 1 h observed data by TROPOMI during the daytime and the daily averaged modelled results by GEOS-Chem, and this is the reason why the modelled results underestimated the daytime observational results. Similarly, O3 exhibited higher concentrations during daytime compared to the nighttime, as it is generated by the photochemical reaction of NO2 and the related HO_X cycle; thus, the modelled results underestimated the observed data due to the large concentration difference between daytime and nighttime. In contrast, the sources of HCHO are mostly secondary products of atmospheric oxidation of VOCs including nocturnal reactions. Therefore, HCHO is produced at all times of the day, leading to a relatively small underestimation of the modelled HCHO compared to the observed results. In addition, the results obtained using TROPOMI were linearly interpolated to fit the grid resolution of GEOS-Chem, which may have caused errors between the observed and calculated results. Comparisons between TROPOMI and GEOS-Chem were conducted using the column density rather than the surface mixing ratio, which may have contributed to error accumulation across vertical layers. These complex analyses resulted in relatively large discrepancies between TROPOMI and GEOS-Chem. Additionally, the accuracy of emission inventories for air pollutants is not perfect, as these inventories are estimated using statistical methods. Collectively, these factors influenced

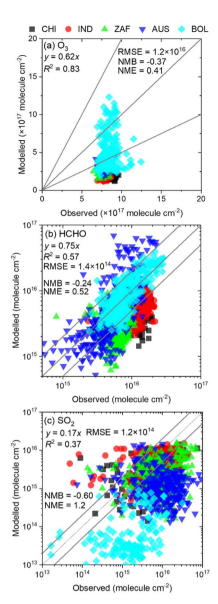
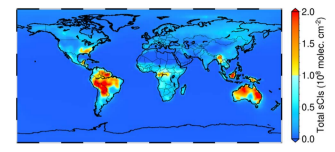


Fig. 2 Comparison of observed and modelled results of the column density of (a) tropospheric O₃, (b) HCHO, and (c) SO₂. HCHO and SO₂ are shown in the scale of common logarithm for readability

the observed differences between TROPOMI and GEOS-Chem. Overall, although discrepancies were observed between TRO-POMI and GEOS-Chem for SO₂, the calculated column densities of O₃, HCHO, and SO₂ correlated with the observed data in terms of determination coefficients, indicating an acceptable agreement between the datasets.

Distribution of sCIs formed by alkene ozonolysis in 2019

Fig. 3 shows the annual average column density of the total sCIs in 2019 calculated using GEOS-Chem. The column density was high in the equatorial regions of the Amazon rainforests, Australia, Central Africa, Southeast Asia, and some countries of Central America and North America, including the U.S. and Mexico. The column densities of the precursors of the sCIs, O_3 , and total alkenes are shown in Fig. S2 of the ESI.† In



Annual mean total column density of sCIs in 2019.

a comparison between Fig. 3 and S2,† the abundance of sCIs was nearly correlated with the abundance of alkenes. Fig. S3-S8 of the ESI† show the column density of the detailed species of the sCIs, and Fig. S9[†] provides a detailed description of the alkene species. According to Fig. S3-S8,† SCI1 (formaldehyde oxide), ESCI3 (E-methyl vinyl ketone oxide), ESCI4 (E-methacrolein oxide), and ESCI7 (methyl glyoxal-1-oxide) were the main contributors of the total amount of sCIs. While SCI1 is generated from both anthropogenic and biogenic alkenes, ESCI3 and ESCI4 are generated from isoprene, a biogenic alkene, and ESCI7 is generated from methyl vinyl ketone, a byproduct of isoprene oxidation. The column density of ESCI7 was the highest among all sCIs. The high concentration of ESCI7 resulted from the low rates of unimolecular decay and ESCI7 and water reactions. Because of the geometrical structure of ESCI7, only 1,3-ring-closure was considered for the unimolecular isomerisation of ESCI7. For example, ESCI3, which is generated by isoprene ozonolysis, involves two unimolecular isomerisation processes: 1,3-ring closure and 1,4-H-migration. The rate coefficients of the 1,3-ring closure for ESCI7 and ESCI3 and of 1,4-H-migration for ESCI3 at 298 K were 0.058 s⁻¹, 0.0040 s⁻¹, and 51 s⁻¹, respectively, calculated using the values listed in Table S2 of the ESI.† Similarly, assuming the concentration of water at 298 K as 8.3×10^{17} molecule cm⁻³, 46 the reaction rates of SCI1, ESCI3, ESCI4, and ESCI7 with water were calculated to be 2866 s⁻¹ (716 s⁻¹ for H_2O and 2150 s⁻¹ for $(H_2O)_2$, 0.525 s⁻¹ (0.065 s⁻¹ for H_2O and 0.460 s⁻¹ for $(H_2O)_2$), $668 \text{ s}^{-1} \text{ (}158 \text{ s}^{-1} \text{ for H}_2\text{O and } 510 \text{ s}^{-1} \text{ for (H}_2\text{O})_2\text{), and } 22.0 \text{ s}^{-1}$ $(2.50 \text{ s}^{-1} \text{ for } H_2O \text{ and } 19.5 \text{ s}^{-1} \text{ for } (H_2O)_2)$, respectively, as calculated from the values listed in Tables S3 and S4 of the ESI.† Thus, because of the relatively high concentration of MVK in the atmosphere, which is shown in Fig. S9 of the ESI,† and because of the lower reaction rate in the main loss processes, which enabled a longer atmospheric lifetime, ESCI7 showed a high concentration in the atmosphere, as discussed by Vereecken et al.20 For similar reasons, E-isomers of sCIs are more dominant than Z-isomers because of the high reactivity of Z-isomers to unimolecular isomerization.20 Some of the previous studies suggested the importance of (E- and Z-)SCI3 and (E- and Z-)SCI4 as atmospheric oxidisers because SCI3 and SCI4 are directly generated from the ozonolysis of isoprene, which is the most relevant VOCs on a global scale.18,19 Nevertheless, this study showed the potential importance of ESCI7 as the most relevant atmospheric oxidiser among the sCIs owing to its long

atmospheric lifetime. Therefore, further studies on the kinetics of ESCI7 in the atmosphere should be conducted in the future. Fig. S9† suggests that ethylene, isoprene, and methyl vinyl ketone are the relevant alkenes in the atmosphere and that ethylene is emitted from both anthropogenic and biogenic sources. Combining these results, most sCIs are formed from biogenic sources, particularly isoprene, on a global scale. The seasonal trend of the column density of sCIs in 2019 is shown in Fig. S10 of the ESI,† where the year is divided into four seasons: January-March (JFM), April-June (AMJ), July-September (JAS), and October-December (OND). In the Northern Hemisphere, the seasons of AMJ and JAS exhibit relatively high concentrations, particularly in the southeastern regions of the United States and the eastern part of Russia (Siberia). This is because AMI and IAS correspond to warm to hot seasons in the Northern Hemisphere, during which high amounts of BVOCs are emitted. Conversely, except for the Amazon rainforest, the seasons of JFM and OND show relatively high concentrations of sCIs in the Southern Hemisphere, also due to high-temperature seasons in those regions. Although the Amazon rainforest is located in the equatorial region and the Southern Hemisphere, its rainy season during JFM, AMJ, and OND results in JAS having the highest concentration of sCIs.

Impact of Criegee chemistry on sulphate formation

Fig. 4 shows the calculated results of the column densities of $SO_4^{2-}(p)$ for (a) annual mean $SO_4^{2-}(p)$ (µg cm⁻²), (b) the change in SO₄²⁻(p) due to Criegee chemistry (μg cm⁻²), and (c) the ratio of the change in $SO_4^{2-}(p)$ (%). According to Fig. 4(a), $SO_4^{2-}(p)$ is distributed globally, but its concentration is high in the Northern Hemisphere, where countries such as China, India, and Middle East are located, which results in high SO₂ emissions from the burning of fossil fuels, as shown in Fig. S2(b).† According to Fig. 4(b), the change in $SO_4^{2-}(p)$ was significant in China and India, both of which emit substantial amounts of SO2 due to the fossil fuel combustion, and in the Amazon rainforests, Central Africa to South Africa, and Australia, which are dominated by forests, causing high BVOC emissions and high sCI concentrations, as shown in Fig. 3. Thus, the effect of Criegee chemistry on $SO_4^{\ 2-}(p)$ formation was high in high-SO₂ and BVOC emissions. Fig. 4(c) demonstrates that the contribution of Criegee chemistries to the rate of change of SO₄²⁻(p) is maximum less than 0.5% in the Amazon rainforests, Australia, and Central Africa, the value of which is not drastic in terms of the air pollution issue. The percentage contribution of Criegee chemistry to $SO_4^{2-}(p)$ formation is high in those regions because the background concentration of $SO_4^{2-}(p)$ is low in remote areas, so the incorporation of the Criegee chemistry showed higher responsiveness in those regions. As mentioned above, SCI1, ESCI3, ESCI4, and ESCI7 were the dominant sCIs on a global scale, and these four sCIs mainly contributed to the formation of $SO_4^{2-}(p)$. Seasonal trends in the column density of SO₄²⁻(p) are shown in Fig. S11-S14 of the ESI,† with the seasons divided in the same manner as in Fig. S10:† January-March (JFM), April-June (AMJ), July-September (JAS), and October-December (OND). The absolute changes in $SO_4^{2-}(p)$ in the eastern part of China during JFM and OND, which

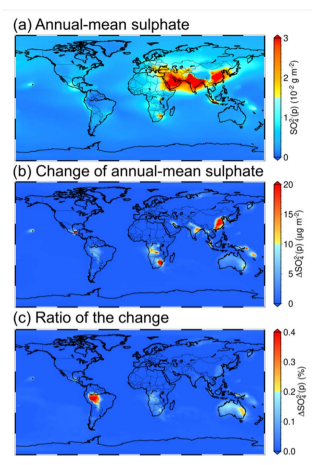


Fig. 4 (a) Annual mean column density of sulphate (g m $^{-2}$), (b) the change in the column density of sulphate before and after the introduction of Criegee chemistry (mg m $^{-2}$), and (c) the rate of change of sulphate before and after the introduction of Criegee chemistry (%).

correspond to the cool to cold seasons, show a significant increase due to high SO_2 emissions driven by increased fossil fuel demand. In the Amazon rainforest, the changes in $SO_4^{\,2-}(p)$ are the highest during JAS, as JFM, AMJ, and OND coincide with the rainy seasons. The concentration of sCIs is also the highest during JAS, as discussed in the previous section.

The contributions of sCIs to the change in $SO_4^{2-}(p)$ shown in Fig. 4(c) are lower than the results of Newland et al., who evaluated the impact of monoterpene-derived sCIs on $SO_4^{2-}(p)$ formation, which suggested that sCIs from monoterpenes account for 1.2% of annual SO₄²⁻(p) formation in the terrestrial tropics.24 The difference might come from the included sCI chemical reactions. Newland et al. considered the reactions of sCIs with SO2, H2O (monomers), and the unimolecular decomposition of sCIs, whereas this study included additional reactions involving (H₂O)₂, NO₂, HNO₃, and three organic acids. The inclusion of these additional reactions would lower the concentration of sCIs, thereby leading to slower oxidation of SO2. Caravan et al. also suggested a 1-10% contribution of $SO_4^{2-}(p)$ formation in the terrestrial tropics from isoprenederived Criegee intermediates (SCI3 in this study),23 but for the same reason as Newland et al.,24 the estimated contribution might be overestimated.

Effect of Criegee chemistry on NO₃⁻(p), NH₄⁺(p), and SOA formation

According to a previous study, the incorporation of Criegee chemistry led to not only an increase in $SO_4^{2-}(p)$ but also a change in other particulate species owing to the change in ionic balance and the enhancement of particle formation through nucleation around the surface of $SO_4^{\ 2-}(p)$.¹⁸ These particulate species include nitrate aerosol (NO₃⁻(p)), ammonium aerosol (NH₄⁺(p)), and secondary organic aerosol (SOA), which are defined in ISORROPA II implemented in GEOS-Chem.⁴⁷ Fig. 5 shows the change in the composition of particulate matter at five sites: CHI, IND, ZAF, AUS, and BOL (Table S1†), which showed high SO₄²⁻(p) formation caused by Criegee chemistry incorporation, as shown in Fig. 4. According to Fig. 5(a), all sites showed an increase in $SO_4^{2-}(p)$ and SOA, the results of which are consistent with a previous study by Hata et al. 18 Budisulistiorini et al. suggested that SOA formation is strongly correlated with the ambient concentration of $SO_4^{2-}(p)$, and a 25% decrease in SO₄²⁻(p) induces a 70% decrease in SOA.48 For this reason, the high increase in SOA shown in Fig. 5(a) is due to the increase in $SO_4^{2-}(p)$ induced by Criegee chemistry incorporation. On the other hand, a decrease in NO₃⁻(p) was estimated for all the analysed sites, which is

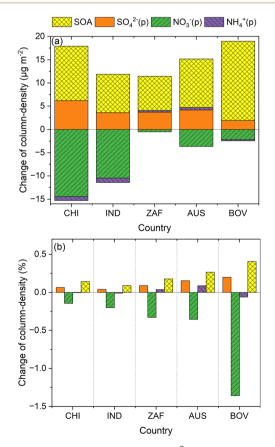


Fig. 5 Cross-effects of the change of $SO_4^{2-}(p)$ to $NO_3^{-}(p)$, $NH_4^{+}(p)$, and SOA by the introduction of Criegee chemistry for (a) absolute change and (b) ratio.

inconsistent with the results of a previous study by Hata et al.18 Hata et al. treated sCIs + NO₂ reactions to completely form NO₃ radicals. However, according to the findings of Caravan et al., only 30% of NO₃ was estimated to be formed by sCIs + NO₂ reactions, and the remaining NO2 was considered to form the adduct sCIs-NO2.45 This new finding was implemented in this study. Thus, the formation of sCIs-NO2 additives resulted in NO₂ removal from the atmosphere, leading to a decrease in NO₃⁻(p). As shown in Fig. 5(a), the decrease in NO₃⁻(p) was large in CHI and IND, where highly anthropogenic air pollutants were likely to be emitted, as implied by the column density of SO₂ (Fig. S2(b)†). These two sites emitted high amounts of NO, subsequently forming NO_2 via a reaction with O_3 , which caused a strong decline in NO₃⁻(p) due to NO₂ removal by sCIs. This suggests that in areas with high emission, the increases in $SO_4^{2-}(p)$ and SOA were cancelled by the reduction of $NO_3^{-}(p)$, all of which were caused by Criegee chemistry incorporation. In contrast, the decrease in NO₃⁻(p) was low at the remaining sites with relatively lower anthropogenic emissions. The behaviour of $NH_4^+(p)$ differed in each site. Because NH_4^+ is the counter ion of SO_4^{2-} , NO_3^- , and other anion species, NH_4^+ (p) increased when the total cations increased (ZAF and AUS) and decreased when the total cations decreased (CHI and IND). According to Fig. 5(b), the change in column densities of $SO_4^{2-}(p)$, $NO_3^{-}(p)$, $NH_4^+(p)$, and SOA at the five sites ranged from -1.3% to 0.4%depending on the sites and species. The ratios were higher for AUS and BOL than for CHI and IND. This trend may have been due to the lower background concentrations of particulate species in AUS and BOL. The lower anthropogenic emissions led to lower particulate species in AUS and BOL, which have resulted in high responsiveness to concentration fluctuations derived from Criegee chemistry incorporation. In contrast, ZAF showed moderate anthropogenic emissions according to the column density of SO₂ (Fig. S2(b)†), indicating an intermediate trend in the ratio (%) between the high- and low-emission sites.

Sensitivity analysis of the rate coefficients of Criegee-induced chemical reactions for the formation of sulphate aerosol

vCIs vs. sCIs: Which CIs are important to $SO_4^{2-}(p)$ formation?

Fig. 6 shows the results of the sensitivity analysis of the vCIs and sCIs (scenarios S1 and S2) for $SO_4^{2-}(p)$ formation at the five sites estimated by eqn (2). Fig. 6 suggests that vCIs contributed more to the formation of $SO_4^{2-}(p)$ than sCIs at all analysed sites. In areas with high emissions, CHI, IND, and ZAF, the contribution of vCIs was much higher than that of sCIs, whereas in sites with lower anthropogenic emissions, AUS and BOL, the contribution was equal. At sites with high to moderate anthropogenic emissions, such as CHI, IND, and ZAF, bimolecular reactions between sCIs and other pollutants, such as NO₂ and HNO3, occurred frequently, which decreased the oxidation of SO_2 by sCIs. Thus, a relatively high vCI contribution to $SO_4^{2-}(p)$ formation was estimated in these sites. In contrast, in AUS and BOL, low concentrations of air pollutants led to a relatively high

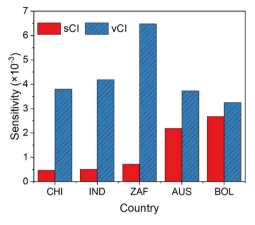


Fig. 6 Sensitivities of sCI and vCI chemical reactions to ${\rm SO_4}^{2-}(p)$ formation in the five analysed sites.

contribution of sCIs to ${\rm SO_4}^{2-}(p)$ formation in AUS and BOL, which was almost comparable to the contribution from vCIs. Previous studies have separately investigated the atmospheric roles of vCIs and sCIs. Khan *et al.* suggested an approximately 13% contribution of OH formation by the unimolecular reactions of vCIs in the terrestrial rainforests and boreal forest regions. Numerous studies have focused on the importance of sCIs in the formation of ${\rm SO_4}^{2-}(p)$ in both remote and urban areas. However, the contribution of ${\rm SO_4}^{2-}(p)$ formation in vCIs and sCIs has not been compared in previous studies. This study quantitatively evaluates their contributions through a detailed sensitivity analysis. The OH from the unimolecular decomposition of vCIs is a more dominant oxidiser of ${\rm SO_2}$ than the direct oxidation of ${\rm SO_2}$ by sCIs, especially in urban areas.

Sensitivities of the sCI-loss reactions *via* unimolecular decomposition and water

Fig. 7 shows the sensitivity of the unimolecular decomposition of the sCIs and the reaction of the sCIs with water for the formation of $SO_4^{2-}(p)$ (scenarios S3 and S4). Most of the results showed a negative impact on $SO_4^{2-}(p)$ formation because both

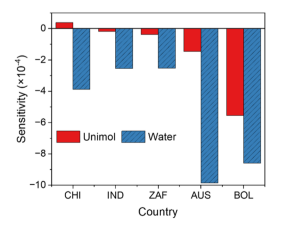


Fig. 7 Sensitivities of sCI-loss reactions by the unimolecular decomposition of sCIs (Unimol) and by the reaction of sCIs and water in the five analysed sites.

the unimolecular decomposition of sCIs and the reaction of sCIs with water are sCI-scavenging reactions, leading to a decrease in the total amount of $SO_4^{\ 2-}(p)$ formation. Only the sensitivity of the unimolecular decomposition of sCIs in CHI showed a positive value, possibly because the products of the unimolecular decomposition of sCIs included OH and the precursors of OH, such as aldehydes, which surpassed the negative effect of the consumption of sCIs. At all the analysed sites, the sensitivity of the reaction of sCIs and water was higher than that of the unimolecular decomposition of sCIs, indicating that the reaction of sCIs with water is more important for the consumption of sCIs and subsequent reduction of SO₄²⁻(p) formation. Only BOL showed high sensitivity to the unimolecular decomposition of sCIs, which was almost equal to the sensitivity of water. This was due to the abundant sCIs derived from BVOCs in BOL, which are in the terrestrial Amazon rainforests and have intense BVOC emissions (Fig. S9†). Vereecken et al. showed that sCIs from BVOCs tend to be lost faster by unimolecular decomposition than sCIs from AVOCs,20 leading to high sensitivity of the unimolecular reactions of sCIs.

Cox et al. 17 and Vereecken et al. 20 estimated the contributions of sCI-loss reactions in urban and suburban areas and found that the contributions of unimolecular decomposition and water scavenging of sCIs are equal or that unimolecular decomposition is higher, and the results are opposite to the results shown in Fig. 7. This study investigated the temperature dependence of the ratio of H₂O/(H₂O)₂, which differed from the results of previous studies. Furthermore, the contributions of the two sCI-loss processes were evaluated using the functional derivative of the sensitivity equation defined by eqn (2), while previous studies estimated the contributions by comparing the ratio of the rate coefficients between unimolecular decomposition and water weighed by the ambient concentrations.17,20 Therefore, it is difficult to completely compare the results between this study and previous studies due to the difference in the definition of the sensitivity and the sCI-chemical reactions included in this study.

Sensitivities of the sCI-loss reactions via NO₂, HNO₃, and organic acids

Fig. 8 shows the sensitivities of the sCI-loss reactions by NO₂, HNO₃, and organic acids to SO₄²⁻(p) formation (scenarios S5, S6, and S7). At all analysed sites, the sensitivity of NO2 showed a positive contribution to $SO_4^{\ 2-}(p)$ formation, whereas the sensitivities of HNO3 and organic acids showed negative contributions. We applied the reaction of sCIs and NO₂ to form 30% of NO₃, according to Caravan et al.45 NO₃ is known as a nocturnal oxidiser and enhances the HO_x cycle,49 and this scheme might contribute to the positive sensitivity of NO₃ to SO₄²⁻(p) formation. However, the sensitivities of the sCI-loss reactions of HNO₃ and organic acids were negative because these species consumed sCIs, which decreased the sCIs + SO₂ reactions. It is important to note that the negative sensitivities of the reactions of HNO₃ and organic acids with sCIs to $SO_4^{2-}(p)$ formation were of the same order of magnitude as those of unimolecular decomposition and water reactions, especially at remote sites, including AUS and BOL. Therefore, the presence of

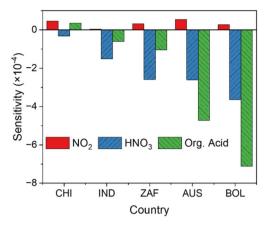


Fig. 8 Sensitivities of the reactions of sCls with NO₂, HNO₃, and organic acids (denoted as Org. Acid in the figure)

HNO₃ and organic acids in the atmosphere led to the decrease of $SO_4^{2-}(p)$ formation through oxidation by sCIs, which are as significant as the unimolecular decomposition of sCIs and the reaction of sCIs and water.

Overall conclusions of the sensitivity analyses

This study shows the sensitivity of the sCI chemical reactions to the formation of $SO_4^{\ 2-}(p)$ according to the detailed sensitivity analysis method defined by eqn (2). The results suggest that in all the analysed sites, the OH formed from vCIs more influenced the formation of $SO_4^{2-}(p)$ than sCIs, but in the remote sites, their difference became closer. Compared with the unimolecular reaction of sCIs and the reaction of sCIs with water, water was more relevant to the sCI-loss reactions at all analysed sites. The loss of sCIs by HNO₃ and organic acids is as important as the loss by unimolecular decomposition and water. In terms of the comparison of vCIs and sCIs with SO₄²⁻(p) formation, this is the first study to clarify the quantitative relationship between them in highly industrialised and rural/ remote sites, concluding that sCI chemistry should be considered in rural and remote areas.

Uncertainties of this work

Although several impacts of CIs on $SO_4^{\ 2-}(p)$ formation were updated in this study, the uncertainties of the analysis must be addressed. There are two types of uncertainties in the analyses: those associated with the CTM itself and those related to the chemical reactions incorporated in this study. Regarding the uncertainties of the CTM, several factors contribute, including emission inventories, meteorological parameters, and gas, heterogeneous-, and aqueous-phase chemistries incorporated in the default CTM. The combinations of these uncertainties ultimately resulted in the outcomes shown in Fig. 2, which indicate a correlation between the observed and modelled results for the column density of O₃, HCHO, and SO₂, although the replicability is not perfect. As discussed above, the discrepancies may stem from the differences in grid resolution between the observed and calculated results, as well as the

accumulation of errors in the vertical layer summation process calculated by the CTM. There is also a possibility of inaccuracies in the emission inventories. Nevertheless, we must accept the errors caused by the CTM itself, as all current CTMs have inherent uncertainties. It is important to note that the results obtained in this study should be treated as one of the conclusions with potential errors stemming from the CTM.

The uncertainties arising from the incorporated reactions of CIs should be treated more carefully, as the inclusion of CI chemistry is the main focus of this study. The results in Fig. 5 suggest that only 30% of NO₃ formation by sCIs + NO₂ reactions is considered to lead to NO2 removal,45 thus contributing to a reduction in tropospheric NO₃, a phenomenon opposite to that observed in previous work.18 Furthermore, the sensitivity analyses shown in Fig. 6-8 indicate that the sensitivities of other CI-related chemical reactions are higher than those of the sCIs + NO2 reactions. This suggests that the definition and accuracy of the chemical mechanisms applied in the CTM affect the atmospheric impacts induced by CI chemistry. For example, Sarwer et al. conducted CTM calculations by adding the chemical mechanisms of sCIs for both low- and high-rate coefficients of sCIs + water reactions to clarify the potential influence of uncertainties in these reactions.50 The results suggested that the contribution of sCIs to $SO_4^{2-}(p)$ was more than an order of magnitude higher when lower rate coefficients for sCIs + water reactions were applied. Lade et al. conducted experimental and theoretical approaches to determine the rate coefficient of SCI1 (formaldehyde oxide) and water reactions, as well as CTM calculations to assess the atmospheric impact of the water monomer and dimer.51 They concluded that more than 98% of SCI1 is consumed by the water dimer, but the uncertainties in the product yields of SCI1 + water reactions limit the understanding of SCI1 chemistry. In terms of the rate coefficients for the unimolecular decomposition of sCIs, this study used the values obtained by Vereecken et al., 20 which were estimated based on classical TST calculations coupled with the semi-empirical method and structure-activity relationships. However, the experimental and theoretical studies for the determination of the rate coefficient for the unimolecular decomposition of dimethyl-substituted CIs, conducted by Lester and Klippenstein, suggested the importance of applying highpressure limit calculations (master equation method) to determine the rate coefficient under ambient conditions and of adjusting tunnelling parameters.52 The rate coefficient estimated by Vereecken et al. was reported to be approximately 1.5 times higher than that of Lester and Klippenstein. The branching ratios of the reaction mechanisms of CI chemistry incorporated in this study also hold several uncertainties, as the products of nearly all reactions involve assumptions due to a lack of information (e.g. assuming 0.5 for the branching ratio of E- and Z-isomers of sCIs after alkene ozonolysis, assuming complete dissociation of unimolecular decay to form OH, assuming the same rate coefficients for all sCIs + HNO₃ reactions, etc.).

Despite these uncertainties, one of the main topics of this study is to clarify the importance of vCI-initiated OH formation in $SO_4^{2-}(p)$ formation. According to the results in Fig. 6-8, the sensitivities of the reaction kinetics of vCIs and sCIs on SO₄²⁻(p) formation are an order of magnitude higher than those of the sCI-

loss reactions. For these reasons, although several reaction mechanisms incorporated in this study involve uncertainties, the contribution of vCIs remains plausible. Further studies should be conducted to clarify the uncertainties in reaction kinetics, branching ratios, and other aspects of CI chemistry in future work.

Summary

The impact of chemical reactions related to CIs on $SO_4^{2-}(p)$ and other aerosol species was evaluated using a global CTM. The absolute amount of $SO_4^{2-}(p)$ formation was found to be high in China and India because of high SO2 pollution, while the percentage of contribution of Criegee chemistry was maximally approximately 0.5% in the Amazon rainforests. The most relevant sCIs on a global scale estimated in this study was ESCI7, the E-isomer of the sCIs formed by the ozonolysis of MVK, owing to the relatively low rate of atmospheric loss processes, which indicates the necessity for further kinetic studies related to ESCI7 chemistry. The results of the sensitivity analyses suggest that in all analysed sites, the OH formed by the unimolecular decomposition of vCIs can have more influence on $SO_4^{2-}(p)$ formation than on the direct oxidation of SO_2 by sCIs, especially in industrialised areas. This indicates that, in terms of aerosol formation via Criegee chemistry, the unimolecular decomposition of vCIs is more important in aerosol formation, and sCIs has increasing importance in the remote area. The sCI loss reactions involving HNO3 and organic acids are as important for $SO_4^{2-}(p)$ formation as those involving the unimolecular decomposition of sCIs and water.

The results of this study showcase the overall global impact of CIs on the formation of tropospheric $SO_4^{2-}(p)$ and related aerosol species, but there are several assumptions for the rate coefficients set in this study (e.g., temperature dependence was not considered for the reactions of sCIs with SO2, NO2, etc., and the branching ratio of E- and Z-isomers was set to 0.5) because of the limited information available on the gas-phase kinetics of sCIs. Further kinetic studies on sCIs should be conducted in future. Furthermore, several studies have revealed the heterogeneous kinetics between the gas/liquid interface of CIs reacting with H₂O, alcohol, etc., forming hydroxy hydroperoxide (HHP), alkoxy alkyl hydroperoxide (AAHP), liquid-phase OH, and RO2, which are important oxidisers in the liquid phase that contribute to the formation of particulate matter.53-55 Besides the detailed gas-phase chemistry of CI, the heterogeneous kinetics of CI should be evaluated by chemical transport modelling in future work to detail the effect of CI on the atmosphere.

Data availability

The data supporting this article are included as part of the main text and in the ESI.†

Author contributions

Hiroo Hata proposed the research concept, conducted the global chemical transport modelling, and wrote the manuscript

draft. Yuya Nakamura determined the kinetic parameters of the Criegee intermediates and checked the manuscript for accuracy. Jairo Vazquez Santiago analysed satellite data obtained from TROPOMI and checked the manuscript for accuracy. Kenichi Tonokura supervised and checked the work for accuracy.

Conflicts of interest

There are no conflicts to declare.

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