

Comprehensive analyses of source sensitivities and apportionments of PM_{2.5} and ozone over Japan via multiple numerical techniques

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Abstract. Source sensitivity and source apportionment are two major indicators representing source–receptor relationships, which serve as essential information when considering effective strategies to accomplish improved air quality. This study evaluated source sensitivities and apportionments of ambient ozone and PM_{2.5} concentrations over Japan with multiple numerical techniques embedded in regional chemical transport models, including a brute forth method (BFM), a high-order decoupled direct method (HDDM), and an integrated source apportionment method (ISAM), to update the source–receptor relationships considering stringent emission controls recently implemented in Japan and surrounding countries. We also attempted to understand the differences among source sensitivities and source apportionments calculated by multiple techniques. While a part of ozone concentrations was apportioned to domestic sources, their sensitivities were small or even negative; ozone concentrations were exclusively sensitive to transport from outside Japan. Although the simulated PM_{2.5} concentrations were significantly lower than those reported by previous studies, their sensitivity to transport from outside Japan were still relatively large, implying that there has been a reduction in Japanese emissions, similar to surrounding countries including China, due to implementation of stringent emission controls. HDDM allowed us to understand the importance of the nonlinear responses of PM_{2.5} concentrations to precursor emissions. Apportionments derived by ISAM were useful in distinguishing various direct and indirect influences on ozone and PM_{2.5} concentrations by combining with sensitivities. The results indicate that ozone transported from outside Japan plays a key role in exerting various indirect influences on the formation of ozone and secondary PM_{2.5} components. While the sensitivities become closer to the apportionments when perturbations in emissions are larger in highly nonlinear relationships – including those between NH₃ emissions and NH₄⁺ concentrations, NO_x emissions and NO₃[−] concentrations, and NO_x emissions and ozone concentrations – the sensitivities did not reach the apportionments because there were various indirect influences including other sectors, complex photochemical reactions, and gas-aerosol partitioning. It is essential to consider nonlinear influences to derive strategies for effectively suppressing concentrations of secondary pollutants.

The air quality of Japan has gradually improved. However, ambient concentrations of fine particulate matter smaller than 2.5 micrometres ($PM_{2.5}$) and photochemical oxidants (predominantly ozone) exceed the Environmental Quality Standards (EQS). Therefore, we must develop effective strategies to suppress ambient $PM_{2.5}$ and ozone concentrations. Quantitative source–receptor relationships serve as essential information when considering effective strategies. There are two major indicators representing source–receptor relationships (Clappier et al., 2017). One is source sensitivity, which corresponds to a change in ambient pollutant concentrations caused by a certain perturbation in precursor emissions. The second is source apportionment, which corresponds to the contribution of precursor emissions to ambient pollutant concentrations. Receptor modelling, including Chemical Mass Balance (CMB) and Positive Matrix Factorization (PMF) methods, have been widely applied to evaluate source apportionments (Hopke, 2016). However, they have limitations when attempting to treat secondary pollutants, which form in the atmosphere via complex photochemical reactions. Moreover, receptor modelling cannot evaluate source sensitivities. Forward modelling using a regional chemical transport model is a powerful tool for evaluating both the source sensitivities and apportionments of primary and secondary pollutants.

Several numerical techniques have been developed for regional transport models to evaluate source sensitivities and apportionments (Dunker et al., 2002; Cohan and Napelenok, 2011). A simple technique for evaluating source sensitivities is the brute force method (BFM). Differences in the simulated pollutant concentrations between two simulation cases with and without perturbations in the input precursor emissions is considered as the sensitivity to a given emission source based on the BFM. This technique can require significant computational resources when evaluating the sensitivities to many emission sources. A decoupled direct method (DDM) is a numerical technique that simultaneously tracks the evolution of sensitivity coefficients, in addition to pollutant concentrations when solving model equations (Yang et al., 1997). This method has been extended to a high-order DDM (HDDM) to track high-order sensitivity coefficients (Hakami et al., 2003). The ozone source apportionment technology (OSAT) (Dunker et al., 2002) and particulate matter source apportionment technology (PSAT) (Wagstrom et al., 2008) are numerical techniques that evaluate the source apportionments of ozone and particulate matter concentrations, respectively, by tagging contributions of precursor emissions to simulated concentrations. An integrated source apportionment method (ISAM) is a similar numerical technique that evaluates source apportionments (Kwok et al., 2013). Each method has its strengths and weaknesses, such that it is important to appropriately interpret results that will be used to develop effective strategies.

Source sensitivities and apportionments of ambient pollutant concentrations over Japan have been evaluated using regional chemical transport models. Chatani et al. (2011) evaluated the sensitivities of simulated $PM_{2.5}$ concentrations over three metropolitan areas in Japan to domestic sources and transboundary transport in the 2005 fiscal year. Ikeda et al. (2015) evaluated the sensitivities of simulated $PM_{2.5}$ concentrations over the nine receptor regions in Japan to source regions in Japan, Korea, and China in 2010. These two studies only employed the BFM to derive source sensitivities of $PM_{2.5}$ concentrations. Itahashi et al. (2015) evaluated the sensitivities and apportionments of simulated ozone concentrations over East Asia to

sources in Japan, Korea, and China. That study presented a unique exercise discussing the differences in source sensitivities and apportionments derived by multiple techniques, including the BFM, HDDM, and OSAT, in Asia; these differences have
65 only been discussed in limited studies targeting the United States and Europe (Koo et al., 2009; Burr and Zhang, 2011; Thunis et al., 2019). Expanding targets is key to obtaining a more comprehensive and appropriate understanding of the source sensitivities and apportionments of pollutant concentrations, including ozone and PM_{2.5}, across Asia, including Japan, derived by multiple techniques.

In addition, recent studies (Ronald et al., 2017; Wang et al., 2017; Zheng et al., 2018) suggest that stringent emission
70 controls implemented in China have achieved improved air quality. These improvements should affect air quality not only in China but also across downwind regions including Japan. We must, therefore, update source sensitivities and apportionments when considering additional effective strategies aimed at further air quality improvement in Japan.

Mutual inter-comparisons of the source sensitivities and apportionments derived by multiple models and numerical techniques is one of the objectives of Japan's Study for Reference Air Quality Modelling (J-STREAM) (Chatani et al., 2018b).
75 Model inter-comparisons conducted in earlier phases of J-STREAM have contributed to the derivation of model configurations and development of emission inventories, both of which have contributed to improved model performance (Chatani et al., 2020; Yamaji et al., 2020). As one of the subsequent activities of J-STREAM, this study evaluates the sources sensitivities of ozone and PM_{2.5} concentrations simulated over regions in Japan for a recent year using the outcomes obtained in earlier phases of J-STREAM. Comprehensive analyses from various perspectives were performed to evaluate the sensitivities to eight
80 domestic and two natural emission source groups, as well as foreign anthropogenic emission sources and transboundary transport throughout the entire 2016 fiscal year. In addition, we perform mutual comparisons of the source sensitivities and apportionments of simulated ozone and PM_{2.5} concentrations. Although the target periods were limited to two weeks in four seasons, we discuss notable characteristics with respect to the differences in the source sensitivities and apportionments derived by the BFM, HDDM, and ISAM.

85 There are well-known nonlinear relationships between ambient concentrations of secondary pollutants including ozone and secondary components involved in PM_{2.5} (Seinfeld and Pandis, 1998). They are likely to cause deviations between source sensitivities and apportionments due to complex photochemical reactions and gas-aerosol partitioning. Nevertheless, it is important to investigate magnitudes of deviations and major causes of nonlinear relationships for considering effective strategies to suppress concentrations of secondary pollutants. Processes causing nonlinear relationships are universal
90 phenomena and not limited to Japan. The findings of this study contribute not only to solving remaining issues involving ozone and PM_{2.5} in Japan, but also to understanding on possible influences of nonlinear relationships in other countries and regions.

2. Methodology

2.1. Model configuration

The Community Multiscale Air Quality (CMAQ) modelling system (Byun and Schere, 2006) version 5.0.2, in which both the HDDM and ISAM are embedded, was selected to calculate the source sensitivities and apportionments, in addition to ambient pollutant concentrations. The carbon bond chemical mechanism with the updated toluene chemistry (CB05-TU) (Whitten et al., 2010) and aero6 aerosol module were employed. Input meteorological fields were simulated by the Weather Research and Forecasting (WRF) - Advanced Research WRF (ARW) version 3.7.1 (Skamarock et al., 2008).

Horizontal locations and resolutions of the four target domains, named as d01, d02, d03, and d04, remain unchanged since the first phase of J-STREAM (Chatani et al., 2018b), as shown in Fig. 1. Horizontal resolutions of d01, d02, d03, and d04 are 45×45 km, 15×15 km, 5×5 km, and 5×5 km, respectively. The top height of the model was lifted from 10,000 to 5,000 Pa to explicitly treat transport in the lower stratosphere (Itahashi et al., 2019a). The vertical layer heights were adjusted to be consistent with those of Chemical Atmospheric Global Climate Model for Studies of Atmospheric Environment and Radiative Forcing (CHASER) (Sudo et al., 2002), which was used to provide boundary concentrations, to avoid numerical diffusions to adjacent layers. Each vertical layer of CHASER from the ground to 80,000 Pa was further divided into two to simulate vertical variations in the lower atmosphere in more detail. The bottom layer height was approximately 28 m.

Several changes were applied to the original WRF configuration employed in the first phase of J-STREAM described in Chatani et al. (2018b) based on the outcomes of the model inter-comparisons. The input land use dataset was replaced with one created from geographic information system (GIS) data based on the 6th and 7th Vegetation Surveys released by the Biodiversity Centre of Japan, Ministry of Environment, which yielded improved performance for multiple meteorological parameters over urban areas (Chatani et al., 2018a). Lakes were added to the dataset based on the National Land Numerical Information Lakes Data. The shortwave and longwave radiation schemes were replaced with the Rapid Radiative Transfer Model for General circulations models (RRTMG) schemes (Iacono et al., 2008) to use the climatological ozone and aerosol profiles with spatial, temporal, and compositional variations (Tegen et al., 1997). Microphysics and cumulus schemes had significant influences on the simulated pollutant concentrations in the model inter-comparisons. A Morrison double-moment microphysics scheme (Morrison et al., 2009) and Grell–Devenyi ensemble cumulus scheme (Grell and Devenyi, 2002) were newly selected because they were characterized by better performance during the sensitivity experiments. Analysis datasets were replaced with the finer ones, i.e., the NCEP GDAS/FNL 0.25 Degree Global Tropospheric Analyses and Forecast Grids (ds083.3) (National Centers for Environmental Prediction/National Weather Service/NOAA/U.S. Department of Commerce, 2015) and Group for High Resolution Sea Surface Temperature (GHRSSST) (Martin et al., 2012), for the initial and boundary conditions, as well as grid nudging. Nudging coefficients are critical parameters for model performance (Spero et al., 2018), but forcing terms in the model equations may disturb physical consistencies. While nudging coefficients for winds were set to $1.0 \times 10^{-4} \text{ sec}^{-1}$ for all domains and vertical layers, those for temperature and water vapour were reduced to 5.0×10^{-5} , 3.0×10^{-5} , 1.0×10^{-5} , and $1.0 \times 10^{-5} \text{ sec}^{-1}$ for d01, d02, d03, and d04, respectively. In addition, nudging for the temperature and

125 water vapour within the planetary boundary layer in d03 and d04 was turned off to avoid excessive nudging to finer spatial
and temporal scales than the input analysis datasets, as well as to allow the simulated values to be in accordance with the
physical equations.

2.2. Emission inputs

Various improvements were applied to the original emission inputs used in the first phase of J-STREAM described
130 in Chatani et al. (2018b) based on the outcomes of the model inter-comparisons. Hemispheric Transport of Air Pollution
(HTAP) emissions version 2.2 (Janssens-Maenhout et al., 2015) was used for anthropogenic sources and international shipping
for Asian countries except for Japan. While the target year of HTAP v2.2 is 2010, the ratios of sectoral annual emissions
reported by Zheng et al. (2018) were multiplied for China, and those reported by the Clean Air Policy Support System (CAPSS)
(Lee et al., 2011) were multiplied for South Korea, to represent the changes in the precursor emissions of recent years. Itahashi
135 et al. (2018) suggested the importance of heterogeneous reactions involving Fe and Mn in sulphate formation. The speciation
profiles of Fu et al. (2013) were applied to consider other components, including Fe and Mn, in addition to originally available
black and organic carbon in PM_{2.5} emissions. The PM_{2.5} emission inventory developed by the Ministry of Environment for the
2015 fiscal year was used for on-road and other transportation sectors in Japan. Emissions from stationary sources in Japan
developed in J-STREAM (Chatani et al., 2018b) were fully updated to the 2015 fiscal year with the following improvements.
140 The emission database of large point sources discretized into sectors, facilities, and fuel types were newly developed by Chatani
et al. (2019) based on the Research of Air Pollutant Emissions from Stationary Sources to represent emissions characteristics
and speciation profiles including Fe and Mn. Missing fugitive volatile organic compound (VOC) emission sources, including
the use of repellents, air fresheners, aerosols inhalers, cosmetic products, and products for car washing and repair, were added
to be consistent with the Greenhouse Gas Inventory Office of Japan (2018). NH₃ emissions from fertilizer use and manure
145 management were replaced by the values reported by the Greenhouse Gas Inventory Office of Japan (2018). Fugitive VOC
and PM emissions from manure management were newly estimated based on the European Environment Agency (2016).
Emission factors of other NH₃ sources, including human sweat, human breath, dogs, and cats, were replaced by those reported
in Sutton et al. (2000). PM emissions from the abrasion of railways wires and rails were newly estimated as one of the major
sources of Fe and Mn. The method to estimate emissions from open agricultural residue burning were replaced by that used
150 by the Greenhouse Gas Inventory Office of Japan (2018). We applied the emission factors reported in Fushimi et al. (2017)
and Hayashi et al. (2014), as well as the temporal variations from Tomiyama et al. (2017). Biogenic VOC emissions were
estimated by Chatani et al. (2018a) using a detailed database of vegetation and emission factors specific to Japan. The surf
zone, defined as zones adjacent to beaches in the National Land Numerical Information Land Use Fragmented Mesh Data,
was newly added to estimate higher sea salt emissions from these areas (Gantt et al., 2015) in the CMAQ.

155 **2.3. Simulation setup**

Ambient pollutant concentrations in d01, d02, d03, and d04 were simulated for the entire 2016 fiscal year (from April 2016 to March 2017). Simulations for the preceding month (March 2016) were treated as spin-up. Sensitivities to the emission source groups, classified as listed in Table 1, were evaluated by the BFM, in which the emissions of each source group were reduced by 20% for the entire fiscal year in d02 and two selected weeks in spring (from May 6 to 20), summer (from July 21 to August 4), autumn (from October 20 to November 3), and winter (from January 19 to February 2 of 2017) in d03 and d04. These two weeks in the four seasons were the periods in which the monitoring campaigns for the ambient concentrations of the PM_{2.5} components were conducted throughout Japan. The reason for choosing 20% reduction as a perturbation range in BFM is that it is a typical range of emission reduction by potential emission controls. For s11 (transport through the boundaries of d02), the boundary concentrations of all species for d02 were reduced by 20%. Differences in the concentrations scaled by five between the simulations with and without 20% perturbations were treated as sensitivities in this study. In addition, source sensitivities and apportionments to all the emission source groups listed in Table 1 were evaluated by the HDDM and ISAM, respectively, using consistent inputs for the two coincident weeks in the four seasons in d02. The first- and second-order sensitivity coefficients to gaseous precursors of a single emission source group were calculated using HDDM. We note that the HDDM results were missing for the seasons other than winter because the simulations were not successfully completed because of numerical convergence problems. Table S1 in the Supplementary Material lists the annual total emission amounts for each source group in d02.

3. Results and discussion

3.1. Model performance on ozone and PM_{2.5}

We evaluated the model performance for the ozone and PM_{2.5} concentrations in d02 for the entire 2016 fiscal year. Table S2 in the Supplementary Material lists the statistics for the model performance of the maximum daily 8-h average ozone (MDA8O3) and daily mean PM_{2.5} concentrations. Table S2 includes the normalized mean bias (NMB), normalized mean error (NME), and correlation coefficient (R) (Emery et al., 2017) for entire Japan (JP), six regions (Kyushu-Okinawa, KO; Chugoku-Shikoku, CS; Kansai, KS; Tokai-Hokuriku, TH; Kanto-Koshinetsu, KK; Hokkaido-Tohoku, HT), and three areas designated by the Automobile NO_x-PM law as polluted urban areas (Osaka-Hyogo, OH; Aichi-Mie, AM; Shuto, ST). Figure 1 denotes the locations and abbreviations of the six regions and three designated areas. Automatic continuous monitoring data obtained at the ambient air pollution monitoring stations (APMSs) were used. Figure S1 in the Supplementary Material compares the observed and simulated monthly mean MDA8O3 and PM_{2.5} concentrations averaged at all stations in the regions.

The MDA8O3 were slightly overestimated in all regions. The observed MDA8O3 was the highest in May and lowest in December. There was another peak in August in western Japan. The model consistently reproduced these monthly variations. Overestimation occurred from the peak in May to the low in December. Values from December to March were slightly

underestimated. The overestimation in summer in this study is less evident than that reported in Chatani et al. (2020), who summarized the performance of the models that participated in the model inter-comparisons conducted in the first phase of J-STREAM. The improved performance obtained in this study may be due to the various improvements in the configurations described in section 2, as well as differences in the meteorological conditions. Kitayama et al. (2019) shows that CB05-TU, which was employed in this study, tends to yield lower ozone concentrations among major chemical mechanisms. All the criteria proposed by Emery et al. (2017) were attained in all regions.

The PM_{2.5} concentrations were underestimated in all regions. The statistics tended to be worse in eastern Japan as opposed to western Japan. The observed PM_{2.5} concentrations fluctuated with a peak in May and valley near September. Although the simulations reproduced these monthly variations, the absolute values were consistently underestimated. A possible reason is discussed in section 3.2. The criteria proposed by Emery et al. (2017) were attained for NME and R, but not for NMB due to persistent underestimation.

As mentioned in section 2, monitoring campaigns for the ambient concentrations of the PM_{2.5} components were conducted throughout Japan for the two target weeks in spring, summer, autumn, and winter. The components of the particulates collected on filters for 24 hours were analysed. These data are useful for the further validation of model performance for the PM_{2.5} components. Figure S2 in the Supplementary Material shows scatter plots of the observed and simulated daily concentrations of the PM_{2.5} components (SO₄²⁻, NO₃⁻, NH₄⁺, elemental carbon (EC), and organic carbon (OC)) at all locations throughout Japan during the monitoring campaigns in all four seasons. Table S1 summarizes their statistics for entire Japan and the four seasons. The simulated average concentrations of SO₄²⁻ and NH₄⁺ are similar to the observed values. Their observed and simulated values have significant correlations with R, i.e., approximately 0.7. The NO₃⁻ concentrations were overestimated with NME of over 100%. The R between the observed and simulated values is 0.441, which is significantly lower than SO₄²⁻ and NH₄⁺. A number of biased dots for NO₃⁻ occur in the scatter plot. While excessively higher simulated values appeared in summer, the model underestimated several of the higher values mainly observed in winter. Although previous studies have discussed issues of poor model performance associated with reproducing the NO₃⁻ concentrations in Japan (Shimadera et al., 2014; Shimadera et al., 2018), they have not yet been solved even after the application of various improvements. Both the EC and OC concentrations were underestimated. As OC is the second major component of PM_{2.5} following SO₄²⁻, its underestimation is one of the major causes of PM_{2.5} underestimation. Shimadera et al. (2018) also discussed the issues of poor model performance associated with reproducing OC concentrations in Japan, suggesting condensable organic matter as a key factor for this poor performance. Although studies on this issue have been conducted by Morino et al. (2018), they remain unsolved.

We note that it is important to recognize that source sensitivities and apportionments introduced in the subsequent sections may be affected by the model performance described in this section.

3.2. Source sensitivities of the annual mean ozone and PM_{2.5}

Figure 2 shows the source sensitivities of the annual mean ozone and PM_{2.5} concentrations derived by the BFM in all regions. Ozone is overwhelmingly sensitive to s11 (transport through the boundaries of d02). The sensitivities of ozone to domestic sources, including s01 (on-road vehicles) and s04 (stationary combustion), are negative in the three designated areas, which is caused by the titration of ozone because of the higher NO_x emissions in urban areas. While the sensitivity of PM_{2.5} to s11 is the highest, PM_{2.5} is also somewhat sensitive to domestic anthropogenic sources, including s01, s02 (ships), s04, and s08 (agriculture and fugitive ammonia). The sensitivities to domestic anthropogenic sources are higher in the three designated areas with higher precursor emissions. The sensitivity of PM_{2.5} to s12 (sea salt) is negative. The sums of the sensitivities of ozone to all the source groups are lower than their simulated concentrations, and the sums of the sensitivities of PM_{2.5} are higher than their simulated concentrations, due to the nonlinear relationships between their concentrations and precursor emissions.

The sensitivities of PM_{2.5} reflect the characteristics of the sensitivities of individual PM_{2.5} components. Figure S3 in the Supplementary Material shows the source sensitivities of the annual mean concentrations of the PM_{2.5} components derived by the BFM in all regions. The EC and primary organic aerosol (POA) are primary components. Sums of the sensitivities of these primary components to all the source groups are consistent with the simulated concentrations. In the three designated areas (OH, AM and ST), EC are specifically sensitive to s03 (non-road transport), and POA is specifically sensitive to s05 (biomass combustion). Sums of the sensitivities of SO₄²⁻, which is mainly a secondary component but almost non-volatile, to all the source groups are also equivalent to the simulated concentrations. SO₄²⁻ is highly sensitive to s09 (natural) in western Japan, i.e., the location of several active volcanoes. Significant nonlinearities exist in the sensitivities of NO₃⁻ and NH₄⁺, which are mainly secondary components. Specifically, although s08 mainly emits NH₃ and no NO_x, NO₃⁻ concentrations are highly sensitive to it because of the indirect influences. Details of these nonlinearities are discussed in section 3.6, which compares the source sensitivities and apportionments. The sensitivities of NO₃⁻ and NH₄⁺ to s12 (sea salt) are negative. Cl⁻ originated from sea salts and mostly involved in coarse particles tend to be replaced by NO₃⁻ because of the so-called chlorine loss caused by gas-aerosol partitioning (Pio and Lopes, 1998; Chen et al., 2016). Therefore, if sea salts are present, more HNO₃ gases are partitioned to coarse particles. That provides capacities for NO₃⁻ and associated NH₄⁺ involved in PM_{2.5} to evaporate to the gas phase, resulting in negative sensitivities of PM_{2.5} including NO₃⁻ and NH₄⁺ to sea salts. Nonlinearities are also significant to secondary organic aerosol (SOA). SOA are specifically sensitive to biogenic VOC emissions included in s09.

Table S3 in Supplementary Material lists the ratios of the source sensitivities of the annual mean ozone and PM_{2.5} concentrations simulated in the regions, which were compared with previous studies. While sums of the ratios of the sensitivities to all the source groups are not 100% because of the nonlinearities, they were often normalized to 100% in previous studies. Therefore, the ratios normalized to make their sums equal to 100% are also shown in Table S3. The annual mean PM_{2.5} concentrations simulated in this study for the three designated areas are 6–9 µg/m³, which is significantly lower than approximately 16 µg/m³ simulated by Chatani et al. (2011) for the corresponding areas in the 2005 fiscal year. However, their

250 ratios of the sensitivities to foreign anthropogenic sources were 48% in OH, 41% in AM, and 31% in ST; these are lower than the approximately 65% calculated in this study as the sums of the sensitivities to s10 (anthropogenic sources in other countries in d02) and s11. The normalized ratios for the sensitivities to the sources in Korea and China for 2010 were 71% in Kyushu, 57% in Kinki, and 39% in Kanto, reported in Ikeda et al. (2015), whereas in this study the sensitivities to s10 and s11 are 68% in KO (equivalent to Kyushu), 65% in KS (equivalent to Kinki), and 59% in KK (equivalent to Kanto). Relative contributions
255 of foreign sources evaluated in this study are even higher than previous studies for most areas of Japan despite the stringent emission controls implemented in China.

One of possible reasons for these elevated contributions is reduction of emissions in Japan. Zheng et al. (2018) showed that the emissions of PM_{2.5}, SO₂, and NO_x in China decreased by 31, 52, and 15%, respectively, from the 2010 to 2016 as a result of the stringent emission controls. If we compare the emissions reported in Chatani et al. (2011) with those used in this
260 study, which reflected changes in energy consumption and emission controls implemented since 2005, the emissions of PM_{2.5}, SO₂, and NO_x in Japan decreased by 29, 48, and 33%, respectively, from fiscal years 2005 to 2015. Therefore, the relative emission reductions in Japan may be larger than those in China if we assume certain increases in the emissions from 2005 to 2010. In particular, stringent emission controls implemented on diesel vehicles by the central and local governments were quite effective in suppressing PM_{2.5} emissions and ambient concentrations in urban areas (Kondo et al., 2012). A reduction in
265 the activity of the Miyakejima volcano in recent years has also resulted in lower SO₂ emissions. However, we can also state that the underestimations of the PM_{2.5} concentrations are larger in eastern than western Japan as described in section 3.1. Influences of domestic sources should be accumulated more in eastern than western Japan because the prevalent air flow over Japan is westerly. Therefore, worse model performance in eastern Japan imply underestimation of domestic emissions. Reductions of domestic emissions from fiscal years 2005 to 2015 may be overestimated.

270 Besides the changes in Chinese emissions, there are other reasons for the higher contributions from sources outside Japan. s11 includes all the components that pass through the boundaries of d02, such that it is affected not only by anthropogenic sources in China, but also anthropogenic sources in other countries, natural sources, and background concentrations.

Ozone concentrations have been relatively stable in Japan in recent years, while the NO_x and VOC concentrations
275 have been suppressed (Wakamatsu et al., 2013). Sensitivities derived in this study suggest that a continuous reduction in the NO_x emissions, because of the stringent emission controls implemented in Japan, has resulted in increases in the annual mean ozone concentrations caused by less titration of the ozone in urban areas. Suppressing the annual mean ozone concentrations further is difficult because they are practically insensitive to domestic sources. Trends in the transboundary transport of ozone likely have a significant effect on the mean annual ozone concentrations (Kurokawa et al., 2009; Chatani and Sudo, 2011). In
280 contrast, the annual mean PM_{2.5} concentrations are sensitive to domestic sources as well as transport from outside Japan. The stringent emission controls implemented in Japan and surrounding countries appear to have contributed to their decreasing trends in Japan. Additional efforts to reduce emissions may produce further improvements in the annual mean PM_{2.5} concentrations, whereas further validations of the emissions in Japan are necessary.

3.3. Monthly variations in source sensitivities of ozone and PM_{2.5}

285 Figure 3 shows the source sensitivities of the monthly mean ozone and PM_{2.5} concentrations derived by BFM simulated for the whole of Japan (JP) and ST, which is one of the three designated areas, including the Tokyo metropolitan area. Figure S4 in the Supplementary material shows the sensitivities of the PM_{2.5} components. Ozone is negatively sensitive to the domestic sources, including s01 (on-road vehicles) and s04 (stationary combustion), in winter because of the titration of ozone by higher NO_x emissions and inactive photochemical reactions in urban areas. The sensitivity of ozone to s11 (transport
290 through the boundaries of d02) is higher than the simulated concentrations, indicating that more ozone is transported from outside Japan and titrated by NO_x emissions in Japan. In contrast, negative sensitivities to domestic sources are less evident in summer even in the ST. Reductions in the ozone by titration are compensated by ozone formation from precursor emissions originating from domestic sources because of the more active photochemical reactions. Differences can be observed in the major source groups, which have positive sensitivities in summer in JP and ST. While the sensitivities of s02 (ships) and s04,
295 which mainly emit NO_x, are higher in JP, those of s07 (fugitive VOC) and s09 (natural), which mainly emit VOC, are higher in ST.

The sensitivity of PM_{2.5} to s11 is the highest in May due to transport by dominant westerly winds in this season. The sensitivity of POA is predominantly high, suggesting that it is affected by variable sources, such as open biomass burning. PM_{2.5} in summer is highly sensitive to s02, s04, and s09, which are mainly located in the southern sides of Japan, because of
300 dominant southerly winds, as well as active secondary formation, which are clearly reflected in the sensitivities of SO₄²⁻ to these sources. PM_{2.5} in winter is highly sensitive to s01 and s08 (agriculture and fugitive ammonia). A colder and more stable atmosphere in winter favours the accumulation of emissions from local sources and the partitioning of NO₃⁻ and NH₄⁺ to the aerosol phase, as reflected in their sensitivities.

As discussed for the annual mean concentrations, suppressing the monthly mean ozone concentrations is difficult
305 because the sensitivities to s11 are dominant in all months. In particular, the sensitivities to s01 and s04 are largely negative in urban areas in autumn and winter. Further reductions in their NO_x emissions may result in additional increases in the monthly mean ozone concentrations in these seasons. In contrast, the negative sensitivities are less evident in spring and summer. Reductions in the precursor emissions for domestic sources have the possibility to suppress, to a certain extent, the monthly mean ozone concentrations. Effective sources may be different in urban and other areas because of differences in ozone
310 formation regimes (Inoue et al., 2019). The effects that strategies have on various sources of precursor emissions for PM_{2.5} may vary seasonally because of differences in meteorological and photochemical conditions.

3.4. Source sensitivities per unit precursor emissions

Air quality standards are defined in terms of ambient concentrations, whereas targets for emission controls are defined in terms of emission amounts. Therefore, it is important to understand whether the sensitivities of ambient concentrations per
315 equal emission amounts of different sources are consistent or not. Figure 4 shows the sensitivities of the annual mean ambient

concentrations of PM_{2.5} components per annual total amount of corresponding precursor emissions of domestic anthropogenic sources (s01–s08) in all of Japan. All the values shown in Fig. 4 were normalized by the EC value for s01, which is inert and emitted only in the bottom layer. The horizontal and vertical locations of the emissions have an effect on the differences in the values for the primary components (EC and POA). Here, s02 includes ship emissions in surrounding oceans in d02, whose values suggest that approximately 40% of the ship emissions in d02 affect the concentrations of primary PM_{2.5} components over Japan. The values of s03 (non-road transport) and s04 (stationary combustion) are slightly lower because they include elevated sources, such as airplanes and large point sources. Slight differences among s01 (on-road vehicles), s05 (biomass combustion), and s06 (residential combustion), whose emissions were ingested only in the bottom layer, may be caused by differences in their horizontal distributions. Sources located in coastal areas may have lower influences as their emissions are transported beyond the land. Additional differences caused by photochemical reactions were observed for secondary components. The value to s05 include agricultural residue burning, which has large spatial and temporal variations, such that its emissions may be high where secondary formation is relatively active. The value of NH₄⁺ to s01 is significantly higher than that to s08 (agriculture and fugitive ammonia) because s01 co-emits NO_x and NH₃, which have a mutual correlation.

The effectiveness of equal reduction amounts of the precursor emissions may be different among sources because of photochemical reactions, as well as the locations of emissions. These factors may need to be considered when exploring effective strategies.

3.5. Differences in source sensitivities among domains

Nesting is a technique in air quality simulations aimed at obtaining improved model performance using finer meshes over target regions, as well as representing large-scale transport in coarser meshes in a computationally effective manner. This study employed d03 and d04 with finer 5 × 5 km meshes over OH, AM, and ST, which include all the major target urban areas. We emphasize the importance of observing how much the sensitivities evaluated in d03 and d04 are different from those in d02 using coarser 15 × 15 km meshes. Figure 5 shows the sensitivities to all the source groups over OH, AM, and ST evaluated in d02, d03, and d04 averaged for the two target weeks during the four seasons. The ozone concentrations simulated for the summer in d02 and d03 or d04 are slightly different. Negative sensitivities to s01 (on-road vehicles) and s04 (stationary combustion) are correspondingly higher. Finer meshes tend to result in slightly larger influences of ozone titration. Although the simulated PM_{2.5} concentrations are slightly different in different domains, the relative contributions of the source groups to the sensitivities are consistent. These results suggest that differences in horizontal resolutions between d02 and d03 or d04 do not cause critical differences in the sensitivities when they are spatially and temporally averaged over the target areas and two weeks. They also support the validity of the discussions in this study, which are mostly based on the results obtained in d02.

3.6. Mutual comparisons of source sensitivities and apportionments derived by BFM, HDDM, and ISAM

3.6.1. Overall differences among techniques

Figure 6 shows the apportionments derived by ISAM and sensitivities derived by BFM and HDDM of the simulated ozone and PM_{2.5} concentrations to all the source groups for the whole of Japan (JP) and ST averaged for the two target weeks during the four seasons. We used the following treatments in Fig. 6. Only the sensitivities to the gaseous precursor emissions were calculated by HDDM. The sensitivities to emissions and boundary concentrations of primary aerosol components (EC, POA, and other primary components) calculated by BFM were also used for HDDM. The simulated SOA concentrations were characterized as apportionments of “OTHR” in ISAM in this study because apportionments of SOA concentrations were not calculated by ISAM embedded in CMAQ version 5.0.2. The HDDM sensitivities were evaluated using first- and second-order sensitivity coefficients ($S^{(1)}$ and $S^{(2)}$) based on the following Taylor expansion (Eq. (1)):

$$C(+\Delta\varepsilon) = C(0) + \Delta\varepsilon S^{(1)}(0) + \frac{\Delta\varepsilon^2}{2} S^{(2)}(0), \quad (1)$$

where $C(+\Delta\varepsilon)$ and $C(0)$ are the simulated concentrations with and without the perturbations, respectively; $\Delta\varepsilon$ is a perturbation ratio; and $S^{(1)}(0)$ and $S^{(2)}(0)$ are the first- and second-order sensitivity coefficients, respectively. The HDDM-20 corresponds to the value calculated by applying $\Delta\varepsilon = -0.2$ and multiplication by 5. If a sensitivity is represented by a second-order polynomial function, HDDM-20 is equivalent to the value obtained by BFM. However, the influence of the second-order term for a perturbation beyond 20% is not reflected in HDDM-20 because the value at a 20% perturbation is just linearly extrapolated. They are reflected in the HDDM-100, which corresponds to the value calculated by applying $\Delta\varepsilon = -1.0$. Differences between BFM and HDDM-20 correspond to the deviations of sensitivities from second-order functions, and differences between HDDM-20 and HDDM-100 correspond to the influences of the second-order term for a perturbation beyond 20%. Sums of the apportionments of all the source groups derived by ISAM represent, in principle, the simulated concentrations.

Not only the sensitivities described in previous sections, but also the apportionments of ozone to s11 (transport through the boundaries of d02) are dominant, suggesting that ozone over Japan is predominantly transported from outside Japan. There are certain positive apportionments of ozone to domestic sources, including s01 (on-road vehicles), s02 (ships), and s04 (stationary combustion), in the spring and summer, indicating that a certain amount of ozone originates from precursors emitted from these sources. Nevertheless, sensitivities of ozone to domestic sources are small or even negative. Let us consider a simple example. Ozone transported from outside Japan reacts with NO emitted in Japan and forms NO₂ (step 1). Next, NO₂ is photochemically decomposed to NO and O, followed by ozone regeneration via a rapid reaction between O and O₂ (step 2). Potential ozone (ozone + NO₂) is preserved in these two steps (Itahashi et al., 2015). Regenerated ozone is apportioned to NO sources in Japan by ISAM in this case. However, if ozone transported from outside Japan increases and enough NO is available, there is a subsequent equivalent increase in NO₂ formation and ozone regeneration. This indicates that regenerated ozone is sensitive to transport from outside Japan. In contrast, if NO emissions in Japan increase, ozone concentrations decrease after

step 1 or remain unchanged after step 2. This suggests that the sensitivities to NO sources in Japan are negative after step 1 or zero after step 2. Their sensitivities cannot become positive in this example. In reality, a certain amount of the NO is oxidized by other species, including RO₂ that originates from VOCs emitted in Japan. They result in net ozone formation and positive sensitivities, which compensates negative sensitivities to a certain extent. The apportionment of ozone concentrations to s11 is smaller than their sensitivities in autumn and winter in ST. The apportionments to domestic sources are negligible in these seasons. Ozone is titrated by high NO emissions in urban areas in step 1, whereas step 2 is not fully reached because of the inactive photochemical reactions.

There are differences in the source apportionments and sensitivities of PM_{2.5}, which reflect those of the PM_{2.5} components, shown in Fig. S5 in the Supplementary Material. Sensitivities of gaseous HNO₃ and NH₃, which are counterparts of NO₃⁻ and NH₄⁺ in the gas phase, are also shown in Fig. S5. The source apportionments and sensitivities of primary components (EC and POA) are consistent. While the sums of the source apportionments and sensitivities of SO₄²⁻ to all the sources are also consistent, there are differences in the relative contributions of the source groups. The apportionment to s11 corresponds to the concentrations of SO₄²⁻ transported from outside Japan. The higher sensitivities are affected by additional indirect influences, i.e., SO₂ is oxidized to H₂SO₄ via gaseous and aqueous reactions, and is then predominantly partitioned to SO₄²⁻. Gaseous SO₂ is oxidized by OH, a part of which originates in ozone. Therefore, s11, which has an overwhelmingly high sensitivity to ozone, also has higher sensitivities of SO₄²⁻ oxidized from SO₂. In contrast, if SO₂ emissions are reduced under fixed OH, other SO₂ remaining in the atmosphere has the opportunity to be oxidized to SO₄²⁻. Therefore, the sensitivities to downwind domestic sources are smaller than their apportionments. Similar discussions are applicable to NO₃⁻. The apportionments of NO₃⁻ and HNO₃ to s11 are lower than their sensitivities, indicating that a certain amount of the NO₃⁻ and HNO₃ is not directly transported from outside Japan. Ozone overwhelmingly affected by s11 enhances the oxidation of NO_x to HNO₃ through OH, followed by a smaller amount that is further partitioned to NO₃⁻. This causes indirect influences on the sensitivities to s11. Such influences are apparent in the horizontal distributions of the apportionments and sensitivities of concentrations of related species to s11 for the two target weeks of spring shown in Fig. S6 in the Supplementary Material. The sensitivities of SO₄²⁻ and NO₃⁻ are higher than their apportionments over Japan. The sensitivities of SO₂ and NO₂ over Japan are correspondingly negative, suggesting that they are oxidized by OH that originated in ozone transported from outside Japan. The isolated higher sensitivities over Japan, particularly visible for those of NO₃⁻, clearly suggest that they are not directly transported from outside Japan.

Section 3.2 discussed higher relative contributions than previous studies and less contrasts between western and eastern Japan for the sensitivities of PM_{2.5} to s11 obtained in this study. Oxidation of SO₂ and NO_x emitted from domestic sources by OH that originated in ozone transported from outside Japan is another factor that causes higher sensitivities of s11. The entirety of Japan is equally affected by ozone transported from outside Japan, as shown in Fig. 2(a), because of its long lifetime in the atmosphere, resulting in less contrast in the sensitivities of PM_{2.5} to s11 between western and eastern Japan, whereas the sensitivities of domestic emissions are small. Ozone governs the oxidative capacity of the atmosphere (Prinn, 2003). If ozone transported from outside Japan is not as reduced in future, efforts to reduce SO₂ and NO_x emissions in Japan

will not effectively contribute to the reduction in the concentrations of SO_4^{2-} and NO_3^- because OH that originated in ozone transported from outside Japan affects their formation.

There is no apportionment of NO_3^- to s08 (agriculture and fugitive ammonia), which emits NH_3 and no NO_x , in accordance to the principle. Nevertheless, NO_3^- is highly sensitive to s08, and is affected by the relationships between NH_4^+ and NO_3^- . Here, NH_4^+ and NO_3^- are mutual counter ions in NH_4NO_3 , whose formation is enhanced when both are available. More NH_3 emissions can induce the partitioning of HNO_3 to NH_4NO_3 . These influences can be observed in the correspondingly negative sensitivities of gaseous HNO_3 to s08. While the apportionments of NH_4^+ are dominated by s08 in ST, its sensitivities are significantly smaller than the apportionments. Both $(\text{NH}_4)_2\text{SO}_4$ and NH_4NO_3 are major forms of NH_4^+ , where, as discussed above, NH_4NO_3 formation is sensitive to NH_3 emissions. In contrast, the sensitivities of SO_4^{2-} to s08 are negligible (Fig. S5(c2)), suggesting that $(\text{NH}_4)_2\text{SO}_4$ formation is predominantly limited by SO_2 sources, including s02 and s04. Their influences are reflected in the sensitivities of NH_4^+ to s02 and s04 (Fig. S5(e2)). These results are consistent with Clappier et al. (2017), who discuss the differences between apportionments and sensitivities in different regimes involving SO_2 , NO_x , and NH_3 using idealized example cases.

There is the certain degree of sensitivity of $\text{PM}_{2.5}$ to s08, as shown in Fig. 2(b), which are indirectly caused by the interactions between NH_4^+ and NO_3^- . There have been several studies that have highlighted the importance of NH_3 emission controls to reduce $\text{PM}_{2.5}$ concentrations (Pinder et al., 2007; Wu et al., 2016; Guo et al., 2018). Such discussions are applicable to Japan. However, Liu et al. (2019) suggested that NH_3 emission control could worsen acid rain because nitric acid is not neutralized and remains in the atmosphere. When seeking strategies to achieve sustainable developments, it is necessary to consider other environmental aspects, including acid rain and nitrogen cycles, as well as the reduction of $\text{PM}_{2.5}$ concentrations.

3.6.2. Nonlinear responses in sensitivities

Differences among the sensitivities derived by BFM, HDDM-20, and HDDM-100 are mostly small, suggesting that, in most cases, HDDM is able to calculate sensitivities consistent with BFM. Slight differences were found in the sensitivities of NO_3^- derived by them. Figure 7(a) shows the sensitivities of the daily NO_3^- concentrations to the source groups located within d02 (s01–s10) derived by BFM, HDDM-20, and HDDM-100 for the two target weeks in winter in ST. The sensitivities derived by BFM are slightly higher than those derived by HDDM-20. While the sensitivities derived by HDDM-20 are only affected by gaseous precursor emissions, those derived by BFM contain minor contributions of primary emitted NO_3^- . They are one of the factors that may result in higher sensitivities derived by BFM. However, differences were found even in the sensitivities to s08 (agriculture and fugitive ammonia), which mostly emits NH_3 . Differences should be recognized as difficulties in representing sensitivities only with first- and second-order sensitivity coefficients derived by HDDM.

Sums of the sensitivities derived by HDDM-100 are higher than those derived by HDDM-20 for all days, indicating nonlinear responses of NO_3^- concentrations against precursor emissions. Daily variations of two additional indicators are shown in Fig. 7(b). One is a nonlinear index (Cohan et al., 2005), which is calculated as follows:

$$\text{Nonlinear index} = \left| \frac{0.5S^{(2)}}{S^{(1)}} \right|. \quad (2)$$

445 This corresponds to an absolute ratio of the second- to first-order sensitivity terms when a perturbation is $\Delta\epsilon = \pm 1.0$, indicating the strength of the nonlinearities. Another indicator is an available NH_3 ratio, which corresponds to a ratio of $\text{NH}_3 + \text{NH}_4^+$ (those stoichiometrically equivalent to SO_4^{2-} are subtracted) to $\text{HNO}_3 + \text{NO}_3^-$, indicating an abundance of potential NH_4^+ that can be combined with NO_3^- . Here, s08 has the highest nonlinear indices that cause the overall nonlinearities, implying that the NO_3^- concentrations have nonlinear responses to NH_3 emissions. Daily variations in the nonlinear indices of
450 s08 and available NH_3 ratios are well correlated; nonlinearities are higher when available NH_3 ratios are lower. The formation of NH_4NO_3 tends to be more constrained by NH_3 with less available NH_3 , as shown by Xing et al. (2011). A typical situation occurred on 30 January. Negative sensitivities of s04 (stationary combustion) suggest that SO_2 emissions of s04 remove NH_3 to form $(\text{NH}_4)_2\text{SO}_4$ and prevent NH_4NO_3 formation. The HDDM can represent such complex nonlinear relationships involving multiple species.

455 In addition to BFM with 20% perturbation (denoted as BFM-20), additional simulations were conducted to derive sensitivities by BFM with 100% perturbation (denoted as BFM-100) for s04, which emits NO_x but not NH_3 , and s08, which emit NH_3 but not NO_x . Figure S7 in the Supplementary Material shows the sensitivities derived by BFM-20, BFM-100, HDDM-20, and HDDM-100, and apportionments derived by ISAM of the daily NO_3^- and NH_4^+ concentrations to s04 and s08 for the two target weeks in winter in ST. The sensitivities derived by BFM-100 are higher than those derived by BFM-20
460 because of the nonlinear responses. Similar features are evident in the sensitivities derived by HDDM-100 and HDDM-20, implying that HDDM is capable of representing directions of nonlinear responses beyond 20% perturbation. It is notable that the sensitivities derived by BFM with a larger perturbation become closer to the apportionments for NO_3^- to s04, and NH_4^+ to s08. However, there are still deviations among them caused by indirect influences of factors including other sectors, complex photochemical reactions, and gas-aerosol partitioning. Moreover, NO_3^- and NH_4^+ concentrations are never apportioned but
465 nonlinearly sensitive to s08 and s04, respectively.

3.6.3. Dependence of ozone formation on NO_x and VOC

ISAM has the capability to separately calculate apportionments of ozone to NO_x and VOC emissions of a given source based on ozone formation conditions (Kwok et al., 2013). It is important to understand relationships between apportionments and sensitivities of ozone to NO_x and VOC emissions. Additional simulations were conducted to separately
470 derive the sensitivities of ozone to NO_x and VOC emissions of s01 (on-road vehicles) by BFM with 20% (BFM-20) and 100% (BFM-100) perturbations.

Figure 8 shows the sensitivities derived by BFM-20 and BFM-100, and apportionments derived by ISAM of daily ozone concentrations to the NO_x and VOC emissions of s01 for the two target weeks in summer in ST. The apportionment to the NO_x emissions is higher than the apportionment to the VOC emissions. While there are differences in the magnitudes of
475 the apportionments and sensitivities to the VOC emissions, their daily variations are consistent. The sensitivity to the NO_x

emissions is mostly negative, but became positive on 25 July when the apportionments, as well as the ozone concentrations, were the highest. The dominant winds were northerly until 24 July and switched to southerly on 25 July. Precursors and the ozone formed from them were transported to the south and returned to ST. Therefore, the aged air mass passed over ST on 25 July 25th. Influences of ozone formation from NO_x emissions were higher than the immediate titration by them for this condition.

Figure S8 in the Supplementary Material shows the sensitivities derived by BFM-20 and BFM-100, and apportionments derived by ISAM of the hourly ozone concentrations to the NO_x and VOC emissions of s01 on 25 July in ST. Hourly variations in the apportionments and sensitivities to the VOC emissions are consistent. Whereas the sensitivities to the NO_x emissions during the night are slightly negative because of titration, their higher positive sensitivities during the daytime indicate the contribution of the NO_x emissions to the high ozone concentrations.

We note that the sensitivities to VOC emissions derived by BFM-20 and BFM-100 are almost identical. That means ozone formation from VOCs is linearly related to emissions. The sensitivities of NO_x emissions derived by BFM-20 and BFM-100 are also almost identical when they are negative. That means titration of ozone by NO_x is also linearly related to emissions. In contrast, the sensitivities to NO_x emissions derived by BFM-100 are higher than those derived by BFM-20 when they are positive. That means ozone formation from NO_x is nonlinearly related to emissions. Cohan et al. (2005) also reported that the sensitivities of ozone concentrations are lower when perturbations of precursor emissions are smaller because other remaining precursors are more likely to contribute to ozone formation instead. This may also be the reason why the sums of the sensitivities to all the sources are lower than the simulated ozone concentrations in spring and summer (Figs. 2, 3, and 5). While the sensitivities derived by BFM-100 become closer to the apportionments, the apportionments are still higher than the sensitivities as discussed for NO₃⁻ and NH₄⁺ in section 3.6.2. That implies effects on concentrations of ozone, NO₃⁻, and NH₄⁺ may be less than those inferred by BFM-100 and ISAM when reductions of emissions of NO_x and NH₃ are small.

Figure S9 in the Supplementary Material shows the horizontal distributions of the apportionments and sensitivities of the ozone concentrations to the s01 NO_x and VOC emissions averaged for the two target weeks in summer. The sensitivity to NO_x emissions is negative in urban and coastal areas where NO_x emissions from on-road vehicles are high. There are consistencies in the horizontal distributions of the positive sensitivities and apportionments to NO_x and VOC emissions. While there are quantitative differences in the magnitudes of the sensitivities and apportionments due to nonlinear influences, ISAM provides spatial and temporal variations in the apportionments to NO_x and VOC emissions consistent with the sensitivities derived by BFM.

4. Conclusions

Sensitivities and apportionments of ozone and PM_{2.5} concentrations over regions in Japan for the 2016 fiscal year to emissions from twelve source groups were evaluated by the BFM, HDDM, and ISAM using emissions data that take into account the latest stringent emission controls. Ozone was predominantly sensitive to transport from outside Japan. While PM_{2.5}

concentrations were lower than those simulated by previous studies for past years because of emission reductions, the relative contributions of transport from outside Japan to the total sensitivities were even larger, suggesting that emissions in Japan have been similarly reduced to surrounding countries, including China. Moreover, sensitivities of $\text{PM}_{2.5}$ included indirect influences of ozone predominantly transported from outside Japan via the oxidation of precursors by OH to secondary $\text{PM}_{2.5}$ components. There was a certain sensitivity of $\text{PM}_{2.5}$ to domestic sources, but the sensitivity of ozone to domestic sources was significantly smaller or even negative because of titration and nonlinear responses against precursor emissions.

Sensitivities and apportionments of primary species were consistent. Fundamental differences were found between them for secondary species. Whereas apportionments represent direct contributions, sensitivities include indirect influences. Clappier et al. (2017) and Thunis et al. (2019) have suggested that sensitivities can provide more useful information than apportionments when considering effective strategies. This study indicates that apportionments simultaneously evaluated with sensitivities can be useful in distinguishing direct and indirect influences, i.e. they cannot be distinguished only by sensitivities. For example, the sensitivities of SO_4^{2-} and NO_3^- to the transport from outside Japan encompassed at least two undistinguished influencing factors, including the direct transport of SO_4^{2-} and NO_3^- , which were evaluated by their corresponding apportionments, and oxidation of SO_2 and NO_x emitted from domestic sources by OH originating in ozone transported from outside Japan. In addition, the titration of ozone by NO_x emissions and inter-correlations between NH_4^+ and NO_3^- in their partitioning were also identified as key indirect influences on ozone and $\text{PM}_{2.5}$.

Sensitivities of $\text{PM}_{2.5}$ derived by BFM and HDDM were mostly consistent except for NO_3^- and NH_4^+ . There were differences between the sensitivities of NO_3^- and NH_4^+ calculated with the first- and second-order sensitivity coefficients derived by HDDM and those derived by BFM. HDDM revealed possibilities to indicate directions of nonlinear responses to larger perturbations in emissions. The sensitivities derived by BFM become closer to the apportionments derived by ISAM when perturbations in emissions are larger in highly nonlinear relationships, including those between NH_3 emissions and NH_4^+ concentrations, NO_x emissions and NO_3^- concentrations, and NO_x emissions and ozone concentrations. However, the sensitivities did not reach the apportionments because of the various indirect influences, including other sectors, complex photochemical reactions, and gas-aerosol partitioning. The dependence of ozone formation on the NO_x and VOC emissions derived by ISAM was spatially and temporally consistent with sensitivities derived by BFM.

Understanding the influences that various factors have on sensitivities can contribute to the establishment of effective strategies. However, accurate sensitivities and apportionments depend on model performance. Uncertainties remain in model performance, as discussed in section 3.1. If specific emission sources affect overall model performance, source sensitivities and apportionments derived by models may be skewed. Figure s10 in the Supplementary Material shows source sensitivities of the annual mean $\text{PM}_{2.5}$ concentrations derived by BFM in the regions. The values shown in (b) were uniformly scaled by the ratios of observed and simulated concentrations of $\text{PM}_{2.5}$ components shown in Table S2. The scaled sensitivities of $\text{PM}_{2.5}$ to the transport from outside Japan are higher by 1.0–2.2 $\mu\text{g}/\text{m}^3$ (15–40%) because of their high contributions to underestimated POA and SOA. The scaled sensitivities of $\text{PM}_{2.5}$ to other sources are different by 0–0.5 $\mu\text{g}/\text{m}^3$. This case assumes that deviations between observed and simulated $\text{PM}_{2.5}$ concentrations can be proportionally explained by the source sensitivities. Uncertainties

could be higher if specific sources cause poor model performance. In particular, this study revealed NH_4^+ and NO_3^- concentrations are nonlinearly sensitive to NH_3 and NO_x emissions. Uncertainties in NH_3 and NO_x emission sources could largely influence source sensitivities as well as model performance of NH_4^+ and NO_3^- concentrations. More studies are
545 necessary to increase the confidence in source sensitivities and apportionments as well as model performance. In addition, sensitivities obtained by the BFM with a single perturbation may be inappropriate for applications to different perturbation ranges when nonlinearities are higher. High-order sensitivity coefficients calculated by the HDDM could help evaluate the importance of nonlinear responses.

This study demonstrated that a combination of sensitivities and apportionments derived by the BFM, HDDM, and
550 ISAM can provide critical information to identify key emission sources and processes in the atmosphere. The sensitivities and apportionments were derived with the consistent model configurations and inputs in this study. However, model configurations and inputs may not necessarily be consistent. Itahashi et al. (2019b) reported that source sensitivities can be changed by the regional chemical transport model with improved treatments for aqueous reactions. Uncertainties in the sensitivities and apportionments caused by different model configurations and inputs should be explored as the next step of J-STREAM.

555 **Data availability**

The input datasets are available upon request at <http://www.nies.go.jp/chiiki/jstream.html>. The output datasets are available upon request to the authors.

Author contribution

SC designed this study, conducted BFM simulations, and prepared the manuscript. HS conducted ISAM simulations, and SI
560 conducted HDDM simulations. KY prepared the meteorology and additional inputs.

Competing interests

There are no conflicts of interest to declare.

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Table 1. Emission source groups to which sensitivities and apportionments were evaluated in this study.

Group	Included emission sources
s01	On-road vehicles
s02	Ships
s03	Non-road transport (machineries, railways, and airplanes)
s04	Stationary combustion (power plants, industries, and commercial)
s05	Biomass combustion (smoking, cooking, and agricultural residue burning)
s06	Residential combustion
s07	Fugitive volatile organic compounds
s08	Agriculture (except for agricultural residue burning) and fugitive ammonia
s09	Natural (volcanoes, biogenic, and soil)
s10	Anthropogenic sources in other countries in d02
s11	Transport through boundaries of d02
s12	Sea salt

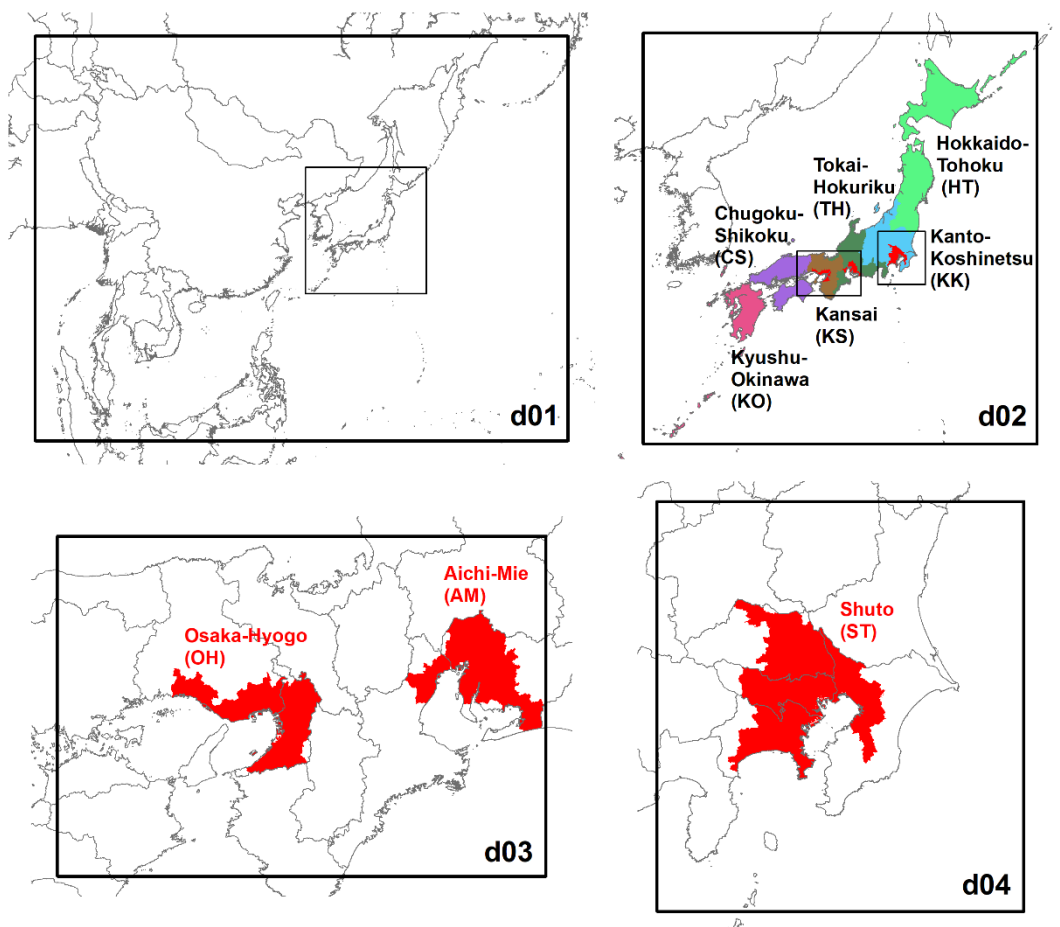
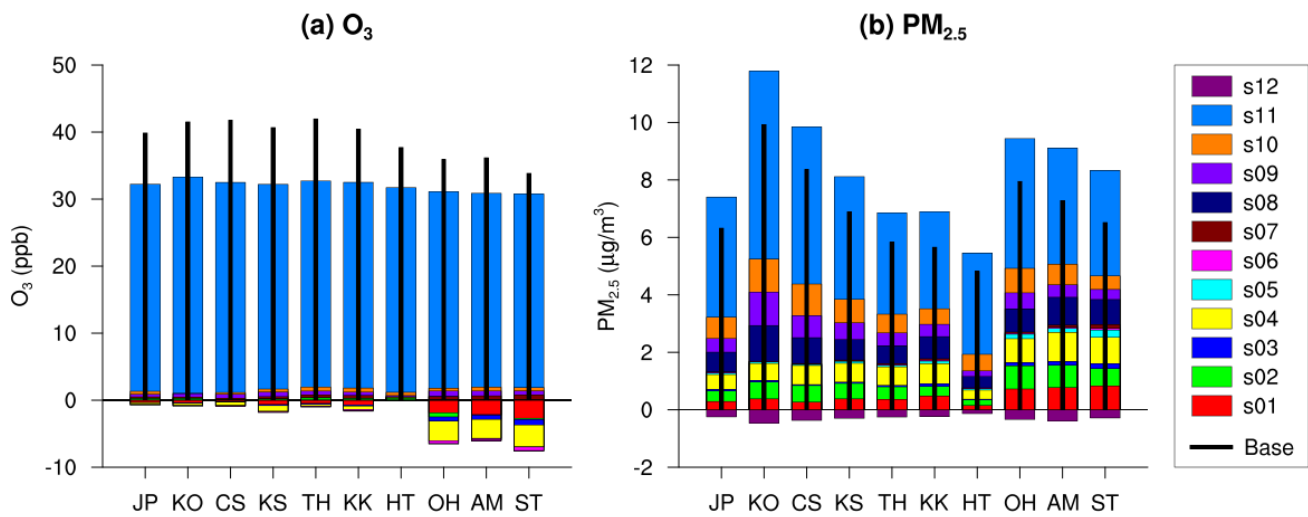


Figure 1. Target domains for the simulations in this study. Results are summarized for six colour coded regions in d02 and three designated areas shown in red in d02, d03, and d04. Their abbreviations are shown in parentheses.



770 **Figure 2. Source sensitivities of the annual mean ozone and PM_{2.5} concentrations derived by BFM in the regions. Thick black lines represent the simulated concentrations.**

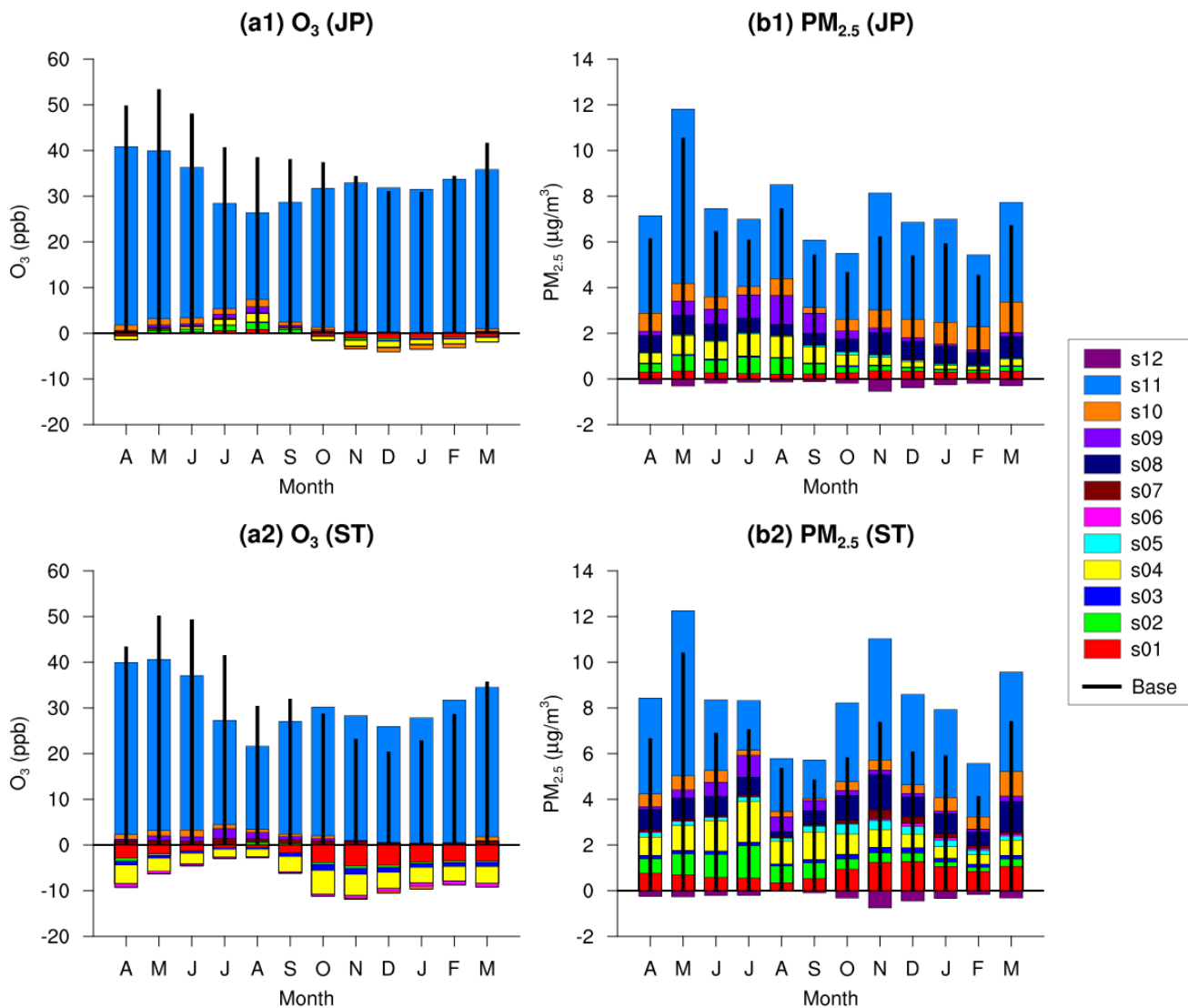


Figure 3. Source sensitivities of the monthly mean ozone and $PM_{2.5}$ concentrations derived by BFM in entire Japan (JP) and ST. Thick black lines represent the simulated concentrations.

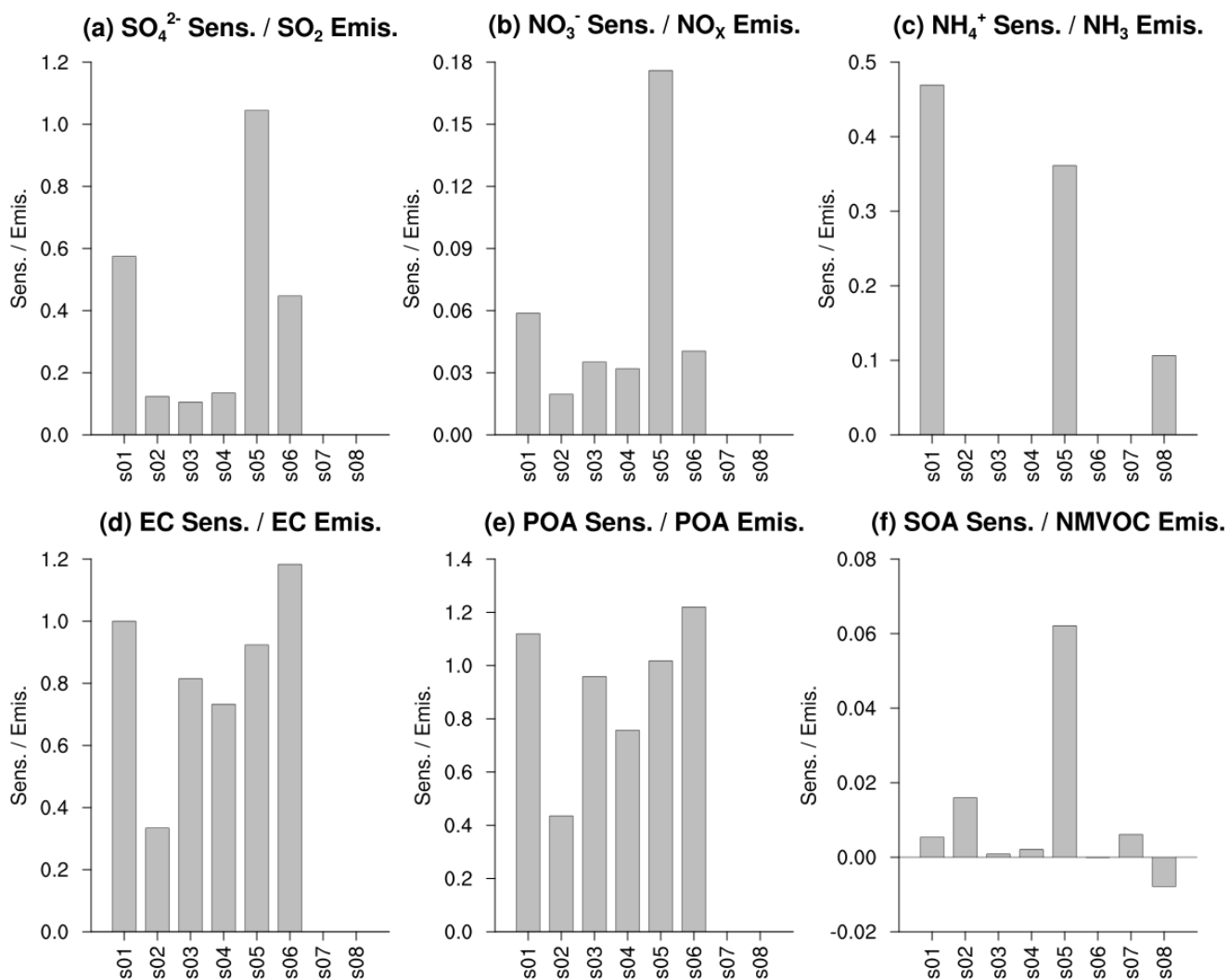


Figure 4. Sensitivities of the annual mean ambient concentrations of PM_{2.5} components in d02 per total annual amounts of corresponding precursor emissions from domestic anthropogenic sources (s01-s08) in entire Japan. All of them are normalized by the EC value for s01.

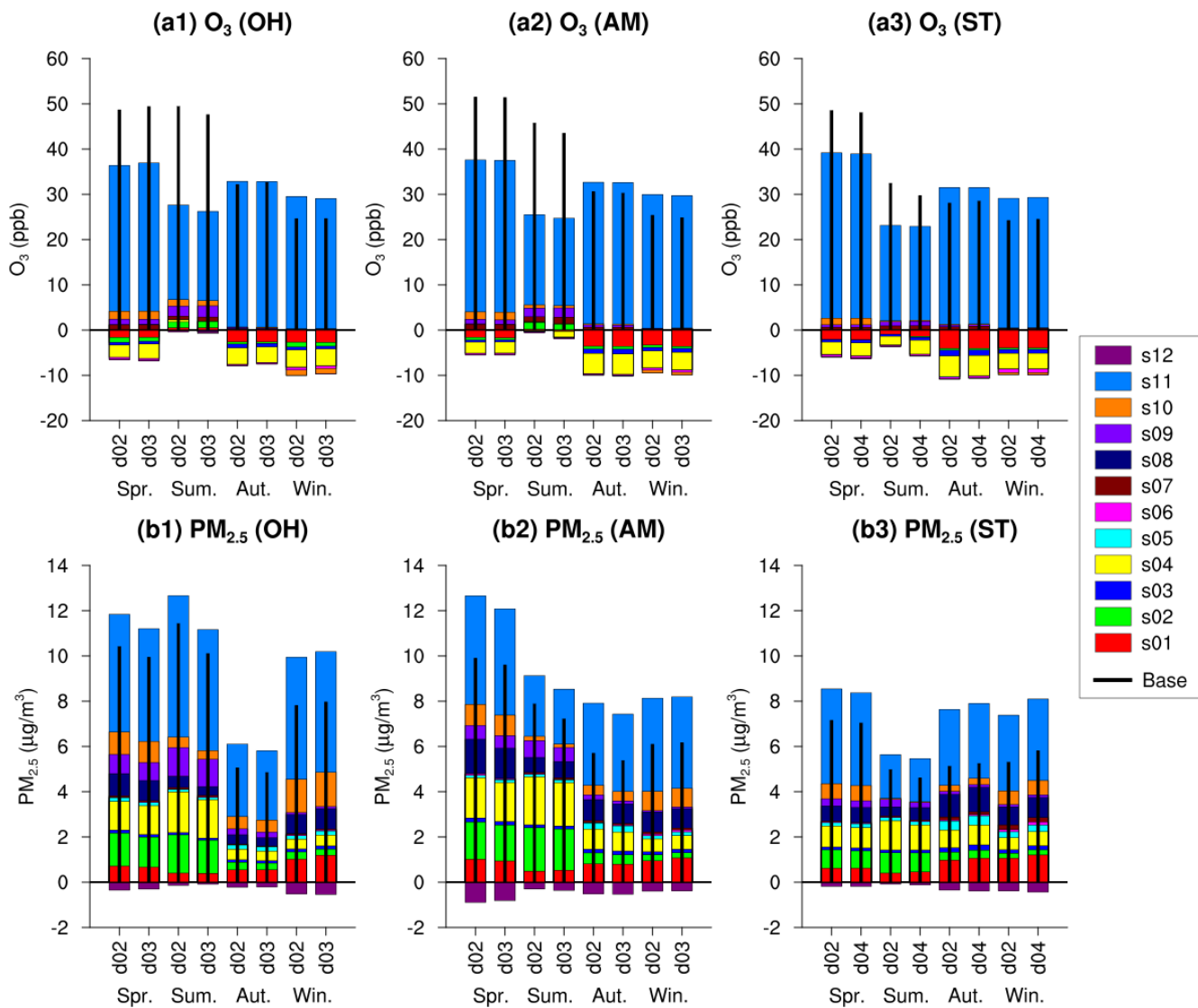


Figure 5. Sensitivities of the simulated ozone and $PM_{2.5}$ concentrations to all source groups over OH, AM, and ST evaluated in d02, d03, and d04 for the two target weeks in the four seasons.

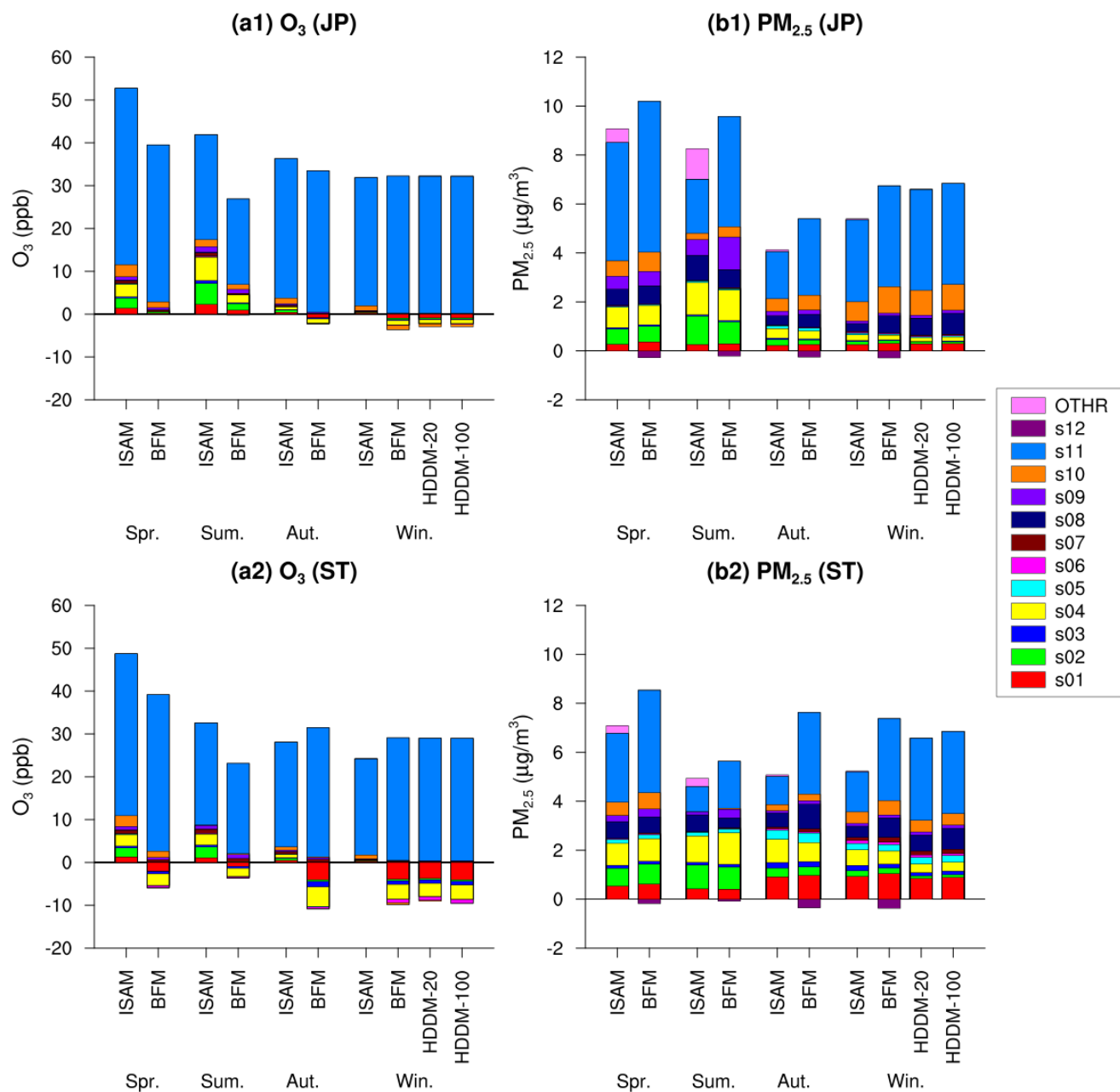


Figure 6. Apportionments derived by ISAM and sensitivities derived by BFM and HDDM of the simulated ozone and $PM_{2.5}$ concentrations to all source groups in JP and ST for the two target weeks in the four seasons.

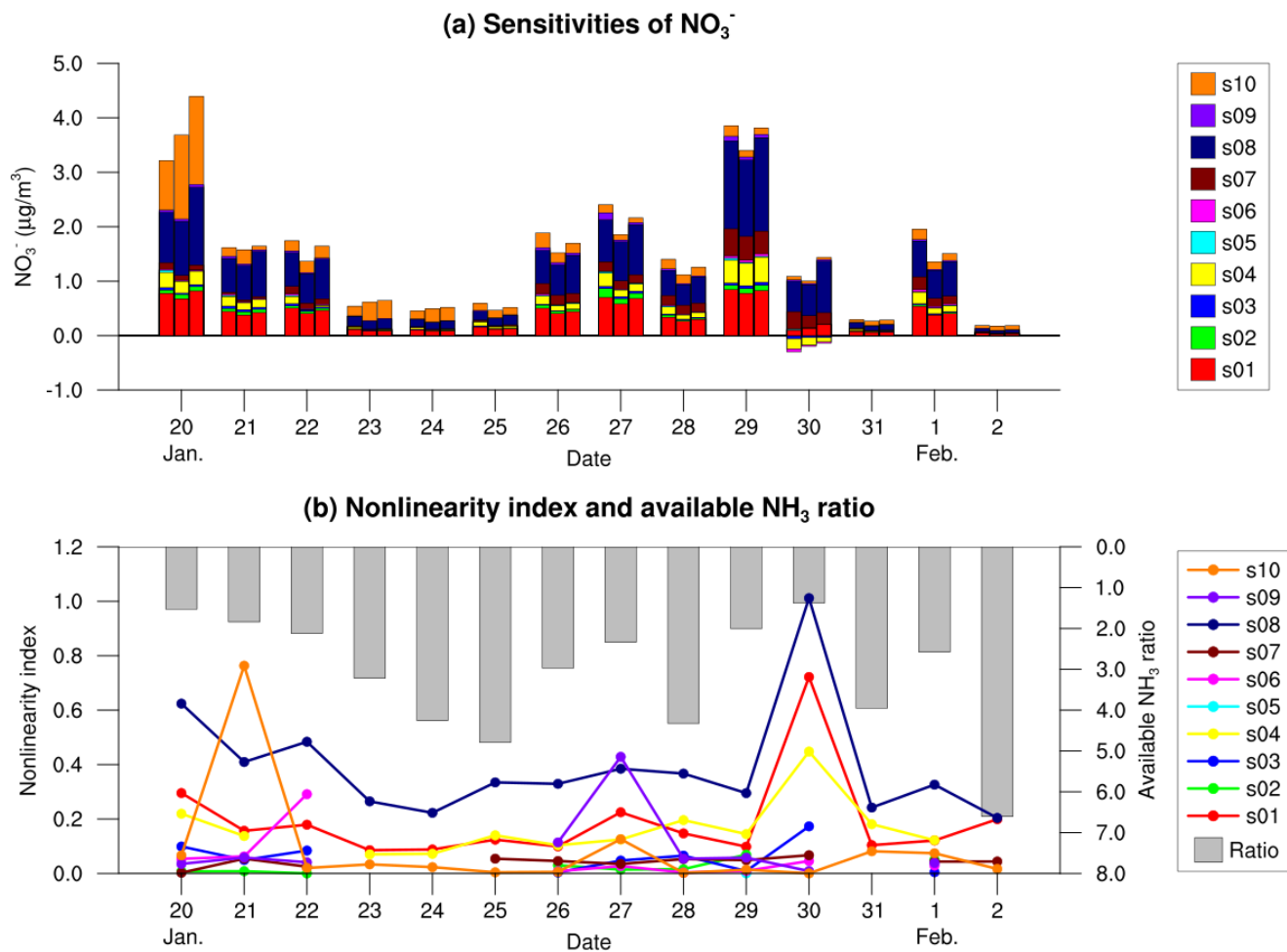
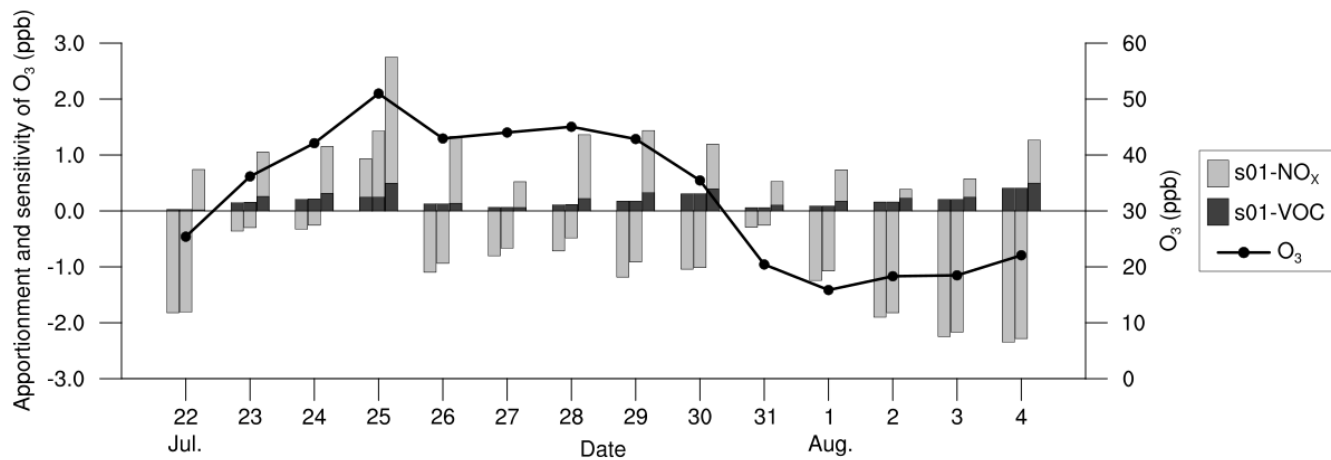


Figure 7: (a) Sensitivities of the daily NO_3^- concentrations to the source groups located within d02 (s01–s10) derived by BFM (left), HDDM-20 (middle), and HDDM-100 (right) and (b) daily nonlinearity index and available NH_3 ratios for the two target weeks in winter in ST. Nonlinearity indices for first-order sensitivity coefficients less than $0.001 \mu\text{g}/\text{m}^3$ are not shown as they are likely to be affected by numerical noise.



800 **Figure 8.** Sensitivities derived by BFM-20 (left) and BFM-100 (middle), and apportionments derived by ISAM (right) of daily ozone concentrations (shown by a line with markers) to the NO_x and VOC emissions of s01 for the two target weeks during the summer in ST.

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