Radiative Forcing and Health Impact of Aerosols and Ozone in China as the Consequence of Clean Air Actions over 2012–2017

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Abstract We applied the chemical transport model GEOS-Chem to examine the changes in aerosols and tropospheric O3 in China from 2012–2017 and the associated radiative forcing and health impact. Simulated surface layer concentrations and column burdens of aerosols and O3 were evaluated by comparing with ground-based and satellite-retrieved measurements. Between 2012 and 2017, simulated annual mean concentrations of PM2.5 decreased by 21.0%, while O3 increased by 11.9% over eastern China (20–45°N, 105–122.5°E). Changes in aerosols and O3 over 2012–2017 jointly exerted a positive radiative forcing of 1.26 W/m² over eastern China, which was dominated by the less cooling from PM2.5 reductions (1.18 W/m²). The Global Exposure Mortality Model predicted 268.3 (247.3–291.6) thousand (9.6%) fewer deaths in eastern China in 2017 relative to 2012. These results suggest an appreciable health benefit and a potential warming as a consequence of clean air actions.

Plain Language Summary Stringent clean air actions have been implemented in China since 2013, aiming to improve air quality. As a result, observed concentrations of aerosols decreased while those of O3 increased in eastern China. In this work we assess the radiative forcing and health impact resulted from the changes in aerosols and O3 in China for the period of 2012–2017 by using a chemical transport model. The model reproduces the observed changes in PM2.5 and O3 in recent years and estimates a net positive radiative forcing of 1.26 W/m² in 2017 relative to 2012 over eastern China. Further estimates from the mortality model show that approximately 270,000 deaths are avoided. Results from our study suggest an appreciable health benefit as well as a potential warming as a consequence of clean air actions over 2012–2017.

1. Introduction

Air pollution has been a serious issue in China and has drawn the intense concern of the government. Starting in 2013, stringent clean air actions were taken in China, aiming to significantly improve air quality by 2017 (Action Plan on Air Pollution Prevention and Control, 2013). Consequently, Chinese anthropogenic emissions of SO2, NOx, CO, BC, and OC decreased by 21–59% from 2013 to 2017 (Zheng et al., 2018), and the average PM2.5 concentrations in 74 cities across China fell by 33.3% (16.3% to 50.3%；Huang et al., 2018). Meanwhile, O3 pollution has not improved and is seemingly getting worse (Huang et al., 2018; K. Li et al., 2019; X. Lu et al., 2018; Ma et al., 2016; Sun et al., 2016). Observed annual mean O3 concentrations during 2013–2017 were reported to change by 20.4% (~30.1% to 71.0%) in 74 key cities in China (Huang et al., 2018), and the summertime maximum daily average 8-hr (MDA8) O3 levels averaged over these cities increased by approximately 20% when concentrations in 2016–2017 were compared to those in 2013–2014 (X. Lu et al., 2018). Since PM2.5 and O3 changed so drastically and differently over the period of 2012–2017, estimates of the associated net radiative forcing and health impact are important for reexamining the clean air actions and for future planning.

Exposure to PM2.5 and O3 was estimated to cause 4.2 million and 254,000 deaths worldwide in 2015, respectively, and was reported to be a leading contributor to the global disease burden (Cohen et al., 2017). Several studies estimated the health benefits from PM2.5 reductions alone in China in recent years (Bi et al., 2019; X.
Both aerosols and tropospheric O3 play important roles in regional/global climate as short-lived forcers. Aerosols influence climate directly through scattering and absorbing infrared and solar radiation, and indirectly by altering cloud properties. Since preindustrial times, aerosols are estimated to have a global mean radiative forcing of $-0.9 (-1.5$ to $-0.4)$ W/m$^2$ (including both aerosol-radiation and aerosol-cloud interactions), partially mitigated the positive radiative forcing of $2.83 (2.26$ to $3.40)$ W/m$^2$ by long-lived greenhouse gases (IPCC, 2013). Tropospheric O3 is a greenhouse gas with anthropogenic radiative forcing of $0.40 (0.2$ to $0.6)$ W/m$^2$ (IPCC, 2013). Variations in ozone concentrations contribute to the climate change in the Northern and Southern Hemispheres and even influence precipitation in China (Son et al., 2009; Xie et al., 2016; Xie et al., 2017; Xie et al., 2018). Given these factors, efforts to improve air quality always come with climate consequences, as demonstrated by developed countries in North America and Europe over the past decades. Leibensperger et al. (2012a) reconstructed the historical aerosol trends in the United States and simulated a positive direct radiative forcing (DRF) of 0.8 W/m$^2$ and a positive indirect radiative forcing of 1.0 W/m$^2$ over the eastern United States between 1980 and 2010 when strict emission reduction regulations were enforced. As a result, a strong regional warming of 0.35 °C was identified in the eastern United States between 1980 and 2010 by the transient climate simulations with Goddard Institute for Space Studies general circulation model 3 (Leibensperger et al., 2012b). Similarly, in Europe during its air quality control period, the decrease in aerosols led to a positive European mean radiative forcing (direct plus indirect) of $1.26$ W/m$^2$ between 1981–1985 and 2001–2005, while the increase in O3 also exerted a positive radiative forcing with a smaller magnitude of $0.05$ W/m$^2$ (Pozzoli et al., 2011). In this regard, the radiative forcing resulted from the changes in air quality in China in recent years is of particular interest because of the very high concentrations of aerosols and O3 in China and the increasingly stringent air quality control policies.

In this study, we use the global 3-D chemical transport model GEOS-Chem to simulate the changes in PM$_{2.5}$ and tropospheric O3 over China from 2012 to 2017. Associated changes in radiative forcing are calculated by the coupled radiative transfer module Rapid Radiative Transfer Model (RRTMG), while attributable mortalities are estimated based on health impact functions with simulated PM$_{2.5}$ and O3 concentrations as the inputs. Integrated or respective roles of PM$_{2.5}$ and O3 are all examined.

2. Methods

We simulate concentrations of aerosols and O3 using the nested grid version (0.5° latitude by 0.625° longitude over Asia 11°S to 55°N, 60–150°E) of the GEOS-Chem model (version 11-01; http://acmg.seas.harvard.edu/geos/) driven by Modern-Era Retrospective Analysis for Research and Applications Version (MERRA-2) assimilated meteorological data (Gelaro et al., 2017). The model includes fully coupled O3-NO$_x$-hydrocarbon chemistry (Bey et al., 2001; Park et al., 2004) and online aerosol calculations including sulfate (Park et al., 2004), nitrate (Pye et al., 2009), ammonium, black carbon and organic carbon (Park et al., 2003), mineral dust (Fairlie et al., 2007), and sea salt (Alexander et al., 2005). Photolysis rates are computed by Fast-JX scheme (Bian & Prather, 2002). Stratospheric O3 is simulated based on linearized O3 parameterization (LINZ; McLinden et al., 2000). Aerosol thermodynamic equilibrium is calculated by the ISORROPRIA II package (Fountoukis & Nenes, 2007). Heterogeneous reactions of aerosols, such as irreversible absorption of NO$_x$ and NO$_2$ on wet aerosols (Jacob, 2000), hydrolysis of N$_2$O$_5$ (Evans & Jacob, 2005), and the uptake of HO$_2$ by aerosols (Thornton et al., 2008) are included. Dry deposition for gas and aerosols is computed based on the resistance-in-series model (Y. H. Wang et al., 1998; Wesely, 1989), and wet deposition follows the scheme of Liu et al. (2001). PM$_{2.5}$ concentration is calculated as the sum of sulfate, nitrate, ammonium, BC, and OC from the model; sea salt and mineral dust aerosols are not considered in PM$_{2.5}$ in this study because they are emitted naturally.
Global anthropogenic emissions in the GEOS-Chem model are from the Community Emissions Data System (Hoesly et al., 2018). For emissions in China, the emissions from Shared Socioeconomic Pathways (Gidden et al., 2019) for year 2015 are used and the annual scaling factors from Zheng et al. (2018) are applied to obtain emissions for other years during 2012–2017. Biomass burning emissions follow the Global Fire Emissions Database Version 4 (van der Werf et al., 2017), and biogenic volatile organic compounds emissions are calculated online by Model of Emissions of Gases and Aerosols from Nature Version 2 (Guenther et al., 2006). Other natural emissions such as NOx from lightning and soil are computed based on MERRA-2 meteorology. From 2012 to 2017, annual emissions (anthropogenic plus natural) of SO2, OC, BC, CO, and NOx decreased sharply over eastern China (105–122.5°E, 20–45°N) by 62.1%, 29.4%, 28.9%, 22.0%, and 21.1%, respectively (Figure S1 in the supporting information), and the largest reductions centered in densely populated city clusters including the North China Plain, the Yangtze River Delta, the Pearl River Delta, Fenwei Plain, central China (e.g., Hubei and Hunan provinces) and the Sichuan Basin (Figure S2). Emissions of NH3 experienced a small decrease of 4.6%, while nonmethane volatile organic compounds emissions increased by 3.3%.

The all-sky DRF of each aerosol species and O3 is calculated via the online coupled radiative transfer module RRTMG (for general circulation models). The RRTMG resolves the spectrum to 14 shortwave bands and 16 longwave bands covering wavelengths from 0.23 to 56 μm. Microphysical properties of each species follow Heald et al. (2014). Instantaneous shortwave and longwave radiative fluxes are computed every 3 hr. The DRF of each species is calculated as the difference of parallel simulations with and without anthropogenic emissions.

The baseline simulations include changes in both emissions and meteorology from 2012 to 2017, and there is a 6-month spin-up before January 2012. Associated premature mortalities are estimated based on simulated concentrations and the specific steps are described in section 3.4.

3. Results and Discussions
3.1. Model Evaluation

Here we conduct comparisons with ground-based and satellite-retrieved measurements (described in Text S1) to see whether the nested grid version of the GEOS-Chem model with the emissions described in section 2 can reproduce the magnitudes and spatiotemporal patterns of surface layer PM2.5 and O3 concentrations, aerosol optical depth (AOD), and tropospheric column ozone (TCO).

Simulated surface layer PM2.5 and O3 concentrations are evaluated using measurements from the CNEMC data set over the period of 2014–2017. Nationwide 841 sites are selected from the CNEMC data set and averaged on the MERRA-2 grid (corresponding to 243 model grids) for comparison. Figure S3 displays the spatial distributions of simulated and observed seasonal mean PM2.5 concentrations in four seasons for 2014–2017. The model reproduces the observed spatial distributions of seasonal mean PM2.5, with high correlation coefficients ($r$) ranging from 0.75 to 0.83 and normalized mean biases (NMB) of 2.0–34.1% for different seasons. In June–August (JJA), March–May, and September–November (SON), both simulated and observed seasonal mean PM2.5 levels were highest over the North China Plain. In DJF, severely polluted zones extended to a larger area of China (including eastern China, northeastern China, and the Sichuan Basin) as shown by a number of grids with both observed and simulated PM2.5 exceeding 100 μg/m³. The model also reproduces the features of PM2.5 seasonality, with a high $r$ value of 0.79 between observed and simulated monthly mean PM2.5 for the ensemble of 243 grids (Figure S4). The monthly mean PM2.5 from both simulations and observations peaks in wintertime due to the strengthened anthropogenic emissions as well as low temperatures favoring nitrate ammonium formation and shows valleys during summertime because of the enhanced wet removal from the East Asian summer monsoon.

Regarding surface O3, the model generally captures the observed seasonal mean spatial patterns in DJF, JJA and SON with $r$ values of 0.60, 0.30, and 0.41, respectively (Figure S5). Observed O3 hot spots such as the Pearl River Delta in DJF, northern China in JJA, and the coastal areas (including Shandong province, the Yangtze River Delta, and the Pearl River Delta) in SON are well reproduced by the model. In March–May, compared with measurements, the model has low biases over the North China Plain and the Yangtze River Delta (with a NMB of −25.7%), whereas high biases outside these two regions (with a NMB
of +29.3%), which lead to a low coefficient correlation over the whole of China when taking all grids into account. Over four seasons, the model generally overestimates O3 concentrations with NMBs in the range of 1.0–35.8%. Monthly variations of surface layer O3 are also captured by the model with a high r value of 0.65 between observed and simulated monthly mean O3 concentrations for the ensemble of 243 grids (Figure S4). Both simulated and observed monthly mean O3 concentrations are high during warm months (April to September), which are driven by enhanced photochemical O3 production owing to high temperatures and strengthened natural emissions of O3 precursors.

Estimation of radiative forcing of air pollutants relies not only on surface layer concentrations but also on column burdens. Simulated AOD and TCO are thus compared with the satellite retrievals from Moderate Resolution Imaging Spectroradiometer (MODIS) and Ozone Monitoring Instrument/Microwave Limb Sounder (OMI/MLS), respectively (Figures S6 and S7). For 2012–2017, observed annual mean MODIS AOD exhibited high values of 0.8–1.0 over eastern China with hot spots occurring in the North China Plain, the Yangtze River Delta, the Sichuan Basin, and central China. This spatial distribution of AOD is well captured by the model with low biases over the Yangtze River Delta and Xinjiang province and high biases over the Sichuan Basin and Northeast China. Simulated AOD averaged over China is 0.30, very close to the value from MODIS observation (0.27). Good agreement is also found between the simulated and observed TCO. Both simulated and OMI/MLS retrieved annual mean TCO show high values of 35–45 DU over eastern China and low values of 20–30 DU over the Tibet plateau. For 2012–2017, simulated annual mean TCO is 33.3 DU over whole China, a little smaller than the value of 35.5 DU from the OMI/MLS data set.

We also examine the changes in annual mean surface layer concentrations of PM2.5 and O3, AOD, and TCO in China, to see whether the model can capture the observed changes in recent years. From 2014 to 2017 when CNEMC data are available, observed annual mean PM2.5 averaged over the 243 grids decreased rapidly by 15.6 μg/m³ in 2017 relative to 2014, and observed annual mean O3 increased by 4.6 ppbv in 2017 (Figure S8a). The model satisfactorily captures these tendencies of changes but with smaller magnitudes; between 2014 and 2017, simulated PM2.5 and O3 show changes of −10.6 μg/m³ and 3.1 ppbv, respectively, when simulated values are averaged over the 243 grids. Following the trend of the surface layer PM2.5 concentrations, MODIS AOD decreased by 0.07 over China from 2012–2017 (Figure S8b). The model reproduces the decrease in AOD, with a simulated decrease of 0.07 over China. While the observed surface layer O3 increased, TCO from OMI/MLS shows a decrease of 1.02 DU over 2012–2017 for China, which may be attributed to the relatively high bias in its 2012 levels (according to Figures S7 and Figure 8a in a previous work; Ziemke et al., 2019). Despite this, the model captures the interannual variation of TCO and simulates an increase of 1.81 DU over 2012–2017. Overall, the baseline simulation can reproduce the surface layer concentrations and column burdens of aerosols and tropospheric O3 in China reasonably well.

### 3.2. Simulated Changes in Air Quality in China From 2012 to 2017

Figure 1 shows the simulated changes in annual mean surface layer PM2.5, AOD, surface layer O3, and TCO over China from 2012 to 2017. Simulated PM2.5 concentrations were highest over eastern China, with a regional mean value of 50.0 μg/m³ in 2012 before the start of clean air actions. From then on, PM2.5 levels began to drop rapidly and by 2017 had decreased 10.5 μg/m³ (21.0%) over eastern China, with largest changes (−16 to −40 μg/m³) found in the North China Plain, the Sichuan Basin, and central China. Simulated AOD shows a similar spatiotemporal pattern as the simulated PM2.5. The AOD over eastern China decreased from 0.52 in 2012 to 0.39 in 2017 with a relative change of −25.0%.

Simulated annual mean surface layer O3 levels are highest over western China (>55 ppbv) because of the large downward transport from the stratosphere. Over eastern China, simulated surface-layer O3 had a mean value of 37.23 ppbv in eastern China in 2012 and increased gradually thereafter by 4.44 ppbv (11.9%) in 2017. The largest increases of O3 of 5–10 ppbv were simulated in eastern China (including the North China Plain and the Yangtze River Delta). Such increases in O3 concentration can be partly attributed to the contemporaneous PM2.5 reductions according to K. Li et al. (2019), because the decrease of PM2.5 can lead to a slowdown of the aerosol sink of HO2 radicals and thus accelerate the O3 production. Following the tendency of changes in surface O3, simulated TCO over eastern China increased by 2.95 DU (7.8%) over 2012–2017.
3.3. Radiative Forcing From Aerosols and Tropospheric O₃ in China Over 2012–2017

The all-sky DRF at the tropopause by aerosols and tropospheric O₃ is presented in Figure 2. In 2012, aerosol DRF was the strongest (−10 to −16 W/m²) over the North China Plain, the Sichuan Basin and central China, which is consistent with the spatial distribution of AOD in 2012 (Figure 1). The aerosol DRF averaged over eastern China was −7.80 W/m² in 2012, which was slightly larger than the values (−4.8 to −6.08 W/m²) estimated by previous studies (Chang et al., 2009; J. W. Li et al., 2013; K. Li et al., 2016; Qian et al., 2016).
2003) as listed in Table S1. These differences in estimated aerosol DRF can be caused by discrepancies in anthropogenic emissions, chemical mechanisms of the models as well as the treatment of aerosol optical properties. In 2017 as compared to 2012, a strong positive aerosol DRF of about 2.0 W/m² is found at the tropopause covering the Sichuan Basin, central China, and Guizhou-Guangxi provinces (Figure 2), which is consistent with the spatial distribution of the changes in AOD between 2012 and 2017 (Figure 1). The aerosols are estimated to have a positive DRF of 1.18 W/m² over eastern China in 2017 relative to 2012, which is mostly contributed by the less cooling of 1.55 W/m² from sulfate aerosols (Figure S9). The decrease in absorbing aerosol BC exerts a cooling of −0.44 W/m² over eastern China, partly counteracts the warming effect resulted from the reductions in scattering aerosols (Figure S9).

Simulated all-sky tropopause DRF of tropospheric O₃ is found to be highest over eastern China (0.6–1.0 W/m²) in 2012, resembling the spatial distribution of TCO (Figure 1). The averaged DRF of tropospheric O₃ is 0.72 W/m² over eastern China in 2012, comparable to the values (0.53 to 0.87 W/m²) obtained by other studies (Chang & Liao, 2009; Chang et al., 2009; W. Wang et al., 2005; Zhu & Liao, 2016) in Table S1. From 2012 to 2017, the increase in tropospheric O₃ enhanced the warming, with a positive DRF of 0.08 W/m² over eastern China (Figure 2). Over the studied period of 2012–2017, the positive DRF caused by O₃ increases is much smaller than the positive DRF brought by reductions in aerosols.

When considering effects of both aerosols and tropospheric O₃, the net DRF from the changes in aerosols plus tropospheric O₃ during 2012–2017 is 1.26 W/m² at the tropopause over eastern China, which is spatially dominated by the pattern of changes in aerosols. Figure S10 also shows the DRF at the surface. Over the 2012–2017 period, changes in aerosols and O₃ are estimated to exert a positive surface DRF of 3.66 and 0.08 W/m², respectively, for eastern China. The IPCC (2013) has reported a global warming of 0.85 °C (0.65–1.06 °C) in surface temperature between 1880 and 2012, and the related global mean radiative forcing values of well-mixed greenhouse gases, tropospheric O₃, and aerosols are +2.83, +0.4, and −0.9 W/m², respectively, in 2011 compared to 1,750 levels. Therefore, our predicted net DRF (aerosols plus tropospheric O₃) of 1.26 W/m² over eastern China in 2017 relative to 2012 is very large and has important implications for regional climate.

### 3.4. Health Impacts of PM$_{2.5}$ and O₃ in China From 2012 to 2017

The premature mortalities attributable to the ambient air pollutants exposures are calculated following the approach from Apte et al. (2015), which can be briefly described as

$$\Delta\text{Mortality} = y_0 \left( \frac{RR(C) - 1}{RR(C)} \right) \text{Pop}$$

where $\Delta\text{Mortality}$ is the premature mortality caused by PM$_{2.5}$ or O₃ exposures, $y_0$ is the baseline mortality rate for a specific disease, Pop is the exposed population, and $RR(C)$, referred as concentration-response functions (CRFs), relate changes in air pollutant concentrations $C$ to changes in relative risk $RR$ for each specific disease (see Text S2 for details). For O₃, we use the CRF from Jerrett et al. (2009) that associates long-term O₃ exposures to the relative risk of respiratory diseases. For PM$_{2.5}$, two kinds of CRFs are applied for further comparisons. One is the integrated exposure-response model (IER) from R. T. Burnett et al. (2014), which is widely used in the mortality estimations in the global burden of disease, World Health Organization, and many previous studies (Apte et al., 2015; Huang et al., 2018; X. C. Lu et al., 2019). The IER is a disease-specific model that is constructed for five causes of deaths by PM$_{2.5}$ including lower respiratory infections (LRI), chronic obstructive pulmonary disease, lung cancer, ischemic heart disease, and stroke. Another CRF we used is the recent Global Exposure Mortality Model (GEMM), which relaxed several strong assumptions required by IER and is assumed to provide better estimates for highly polluted environments such as China (R. Burnett et al., 2018).

The GEMM comprises two forms of applications, which are GEMM 5-COD and GEMM noncommunicable disease (NCD) + LRI; the first one is used for estimating separate mortality by five causes of deaths aforementioned (5-COD: LRI, chronic obstructive pulmonary disease, LC, ischemic heart disease, and stroke), thus fitting for the comparison with assessments from the IER; and the second one aims for broader causes of deaths as the sum of NCDs and LRI.
Figure 3 shows the estimated annual deaths owing to ambient O₃ and PM₂.₅ (based on two CRFs) pollutions in eastern China from 2013 to 2017 relative to 2012 levels. For PM₂.₅, estimates based on two different models (a) IER and (b) GEMM are displayed. Bars indicate the estimates for each specific cause of death attributable to PM₂.₅ and O₃. Black lines indicate the total estimates of PM₂.₅ plus O₃. Error bars indicate the 95% confidence interval (CI). In (b), total estimates from GEMM noncommunicable disease + LRI are indicated in the dashed line. IER = integrated exposure-response model; GEMM = Global Exposure Mortality Model; LRI = lower respiratory infections; IHD = ischemic heart disease; COPD = chronic obstructive pulmonary disease; LC = lung cancer.

Figure 3 shows the estimated annual deaths owing to ambient O₃ and PM₂.₅ (based on two CRFs) pollutions in eastern China from 2013 to 2017 relative to 2012 levels. The increases in O₃ concentrations resulted in more deaths of respiratory diseases over the period, with increased numbers varied from 4.4 thousand in 2013 to 16.0 thousand in 2017 for eastern China. As for PM₂.₅, the attributable deaths show a continuous decline. By 2017, the improvement of PM₂.₅ pollution is estimated to led to 84.5, 198.3, and 284.3 thousand avoided deaths in eastern China based on IER, GEMM 5-COD, and GEMM NCD + LRI, respectively, far offsetting the excess deaths brought by O₃ increase. The estimates from GEMM are much larger than those from IER (a ratio of 0.3–0.4 between IER and GEMM), suggesting that more health benefits may come along with the improvement of PM₂.₅ pollution than what previous studies have expected. Overall, changes in PM₂.₅ and O₃ are estimated to ease the burden of disease from ambient air pollution in eastern China by −5.9%, −8.4%, and −9.6% with avoided deaths numbers of 68.55 (95% CI: 49.3–101.6) thousand, 182.35 (95% CI: 147.3–211.4) thousand and 268.32 (95% CI: 247.3–291.6) thousand based on IER, GEMM 5-COD, and GEMM NCD + LRI, respectively. For more information on estimates for full coverage of China, please see Table S2.

4. Conclusions

We calculate the radiative forcing and premature mortality from aerosols and O₃ in China for the air pollution control period of 2012–2017 by using the GEOS-Chem model. Simulated surface-layer concentrations and column burdens of aerosols and O₃ are evaluated using the ground-based and satellite-retrieved measurements. Between 2014 and 2017, the model satisfactorily reproduces the observed 71.3% decrease in surface layer PM₂.₅ concentration and 72.1% increase in surface layer O₃ concentration averaged over 243 grids across China.

Over the 2012–2017 period, simulated annual mean concentration of PM₂.₅ decreased by 21.0% over eastern China (105°–122.5°E, 20°–45°N) and that of O₃ increased by 11.9%. Reductions in aerosols led to a warming effect on the Earth system with a regional mean tropopause DRF of 1.18 W/m² over eastern China in 2017 relative to 2012. The increase in tropospheric O₃ also resulted in a positive DRF, with a smaller magnitude of 0.08 W/m² over eastern China. Further mortality assessment shows that PM₂.₅ reductions avoided 284.3 thousand deaths in eastern China in 2017 relative to 2012 based on GEMM NCD + LRI, far offsetting the excess deaths (16.0 thousand) caused by O₃ increases. In summary, changes in PM₂.₅ and O₃ from 2012–2017 are estimated to jointly exerted a positive DRF of 1.26 W/m² and led to 268.3 (95% CI: 247.3–291.6) thousand (9.6%) fewer deaths in eastern China.
The results in our study show an appreciable health benefit related to air pollution control from 2012 to 2017 in China, indicating a great effectiveness of clean air actions in terms of health impact. However, a potentially warming effect is also suggested on regional climate, which should be considered in future planning. It should be noted that the GEOS-Chem model is driven by assimilated meteorological fields MERRA-2, and the feedbacks between meteorology and air pollutants are not included. Fully coupled chemistry-climate model estimates are thus better to understand the climatic responses to the control of anthropogenic emissions.

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References


References From The Supporting Information

