Ozone pollution in the North China Plain spreading into the late-winter haze season

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Surface ozone is a severe air pollution problem in the North China Plain, which is home to 300 million people. Ozone concentrations are highest in summer, driven by fast photochemical production of hydrogen oxide radicals (HOx) that can overcome the radical titration caused by high emissions of nitrogen oxides (NOx) from fuel combustion. Ozone has been very low during winter haze episodes to levels approaching the air quality standard. This fast ozone production was driven by formaldehyde originating from high emissions of volatile organic compounds (VOCs). The North China Plain experiences severe summer ozone pollution, but ozone during winter haze (particulate) pollution events has been very low. Here, we show that the abrupt decrease in nitrogen oxide (NOx) emissions following the COVID-19 lockdown in January 2020 revealed a switch to VOC-relevant conditions. The ozone pollution season at northern mid-latitudes is typically limited to May–September months when UV fluxes are high (5, 14). However, high ozone events in winter have been observed in an oil/gas production basin in the United States and attributed to photolysis of carbonyls with HCHO as the leading species (12, 15). Ozone production in winter is generally in the VOC-limited regime (equivalently called the NOx-saturated regime) because of radical scavenging from oxidation of NOx to nitric acid (16). Thus, ozone is expected to increase as VOC emissions increase and as NOx emissions decrease.

Photolysis of HONO is a large source of HOx radicals in the North China Plain in winter (13, 17, 18). However, winter ozone production takes place in polluted regions by photochemical oxidation of volatile organic compounds (VOCs) in the presence of nitrogen oxides (NOx ≡ NO + NO2), catalyzed by hydrogen oxide radicals (HOx ≡ OH + peroxy radicals) as oxidants. NOx in urban air is mainly from fossil fuel combustion, while VOCs have a range of anthropogenic sources. The major sources of HOx radicals in urban air are the ultraviolet (UV) photolysis of ozone, nitrous acid (HONO), formaldehyde (HCHO), and other carbonyl compounds (11–13). The ozone pollution season at northern mid-latitudes is typically limited to May–September months when UV fluxes are high (5, 14). However, high ozone events in winter have been observed in an oil/gas production basin in the United States and attributed to photolysis of carbonyls with HCHO as the leading species (12, 15). Ozone production in winter is generally in the VOC-limited regime (equivalently called the NOx-saturated regime) because of radical scavenging from oxidation of NOx to nitric acid (16). Thus, ozone is expected to increase as VOC emissions increase and as NOx emissions decrease.

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concentrations are generally low (19), dropping down to a few parts per billion (ppb) during winter haze pollution episodes (20, 21) when weather conditions are stagnant (22, 23). Observations in the North China Plain show a strong negative correlation between PM$_2.5$ and ozone in winter (4, 24). The low ozone is attributed to titration by high NO emissions (19) and strong suppression of ozone formation under NO$_X$-saturated conditions (20). The abrupt lockdown of the North China Plain on January 24, 2020, in response to COVID-19 led to a sharp decrease in NO$_X$ emissions as indicated by ground-based and satellite data (25–27). Winter haze events with high PM$_{2.5}$ were still observed during the lockdown but were associated with increased ozone levels (28). The oxidizing environment resulting from the increased ozone was reported to be the driving reason for the high secondary PM$_{2.5}$ concentrations during the lockdown (29, 30). The increase of ozone during the lockdown has been tentatively attributed to weakened titration (29, 31), ozone production (32), and decreased PM$_{2.5}$ (27, 31). Understanding the factors controlling the high ozone observed in winter haze events during the lockdown provides insight into how continued decrease of emissions could result in growing ozone pollution in the North China Plain in winter–spring.

Here, we present air quality observations in the North China Plain during the COVID-19 lockdown and show that fast photochemical ozone production took place in haze events. We show that the COVID-19 lockdown underscores what has been a general trend since 2013 of increasing ozone pollution in winter–spring, including exceedances of the air quality standard of 160 µg m$^{-3}$ (75 ppb at Standard Temperature and Pressure [STP]) for the maximum daily 8-hour average (MDA8) ozone concentration. We explain this increase as driven by the 30% decrease of NO$_X$ emissions combined with flat VOC emissions for the 2013 to 2019 period under the Clean Air Action. Unlike in summer, we find that the ozone increase in winter–spring is not driven by decreasing PM$_{2.5}$ (6) but rather by decreasing NO$_X$ emissions. The decrease of NO$_X$ emissions has caused a major broadening of the ozone pollution season to now extend over much of the year. The COVID-19 lockdown experience shows that high ozone events may become more frequent and severe in winter–spring in the absence of action to reduce VOC emissions.

**Results and Discussion**

**High Ozone in Winter Haze during the COVID-19 Lockdown.** The national lockdown policy starting on January 24, 2020, drove a sharp decrease in transportation and industrial emissions, amplified by the Chinese Spring Festival holiday. We define here the lockdown period as January 24 to February 15 when lockdown measures were particularly strict (25). Fig. 1 and SI Appendix, Fig. S1 show 60 to 70% observed decreases in surface NO$_X$ concentrations and tropospheric NO$_X$ columns from the TROPOspheric Monitoring Instrument (TROPOMI), satellite, implying sharp reductions of NO$_X$ emissions. TROPOMI HCHO observations suggest a weaker reduction of VOC emissions. The surface NO$_X$ concentrations are from the Ministry of Ecology and the Environment (MEE) monitoring site network. For the North China Plain as defined in Fig. 1A, average decreases during the lockdown were 57% for surface NO$_X$, 67% for TROPOMI NO$_X$, and 13% for TROPOMI HCHO. By contrast, ozone concentrations measured at MEE sites increased substantially during the COVID-19 lockdown. Maximum MDA8 ozone at individual sites in the North China Plain increased from 41 ± 5 ppb before lockdown (Fig. 1D) to 59 ± 5 ppb during lockdown (Fig. 1H).

Fig. 2 shows the time series of MDA8 ozone, PM$_{2.5}$, and peroxyacetyl nitrate (PAN) in Beijing before and during the lockdown. Stagnant weather during the lockdown (29, 32) caused two severe haze episodes, as seen in the PM$_{2.5}$ peaks on January 28 and February 12. Ozone during these two events rose to 56 and 57 ppb, respectively, which positively correlated with PM$_{2.5}$. The correlation between ozone and PM$_{2.5}$ shifted from negative ($r = -0.53, P < 0.05$) before the lockdown to positive ($r = 0.52, P < 0.05$) during the lockdown. Data from previous years show persistent negative correlations between PM$_{2.5}$ and ozone in the wintertime in the North China Plain (4, 24). This is in contrast to summer when PM$_{2.5}$ and ozone are positively correlated except under very high PM$_{2.5}$ conditions (33). Odd oxygen (O$_X$ ≡ O$_3$ + NO$_X$) increases during the two haze episodes (SI Appendix, Fig. S2), indicating that the increase of ozone is not simply attributable to decreased titration by NO emissions and must be driven instead by photochemical production.

Further evidence for photochemical production of ozone comes from the increase in PAN during the haze pollution events of the lockdown period, reaching up to 4 ppb and synchronous with ozone and PM$_{2.5}$ (Fig. 2C). By contrast, PAN levels were very low before lockdown. PAN is produced by the same photochemistry as ozone, and under polluted conditions, its production rate is determined by the HO$_X$ source and the NO$_X$/NO ratio, the latter increasing with increasing ozone (34). Both ozone and PAN show diurnal cycles of depletion at night and increases in the daytime, with particularly large daytime increases during the haze episodes indicative of fast photochemical production (SI Appendix, Fig. S3).

We used the Goddard Earth Observing System Chemical Transport Model (GEOS-Chem) with 25-km resolution (Materials and Methods) to explain the high ozone in haze events during the COVID-19 lockdown. The model has been used previously to simulate PM$_{2.5}$ (35) and summertime ozone (33) in the North China Plain. Anthropogenic emissions are from the Multiresolution Emission Inventory for China (MEIC) (36) and updated to 2020 by applying sectoral scaling factors (Materials and Methods). The effect of the lockdown is simulated by decreasing NO$_X$ and VOCs emissions by 60% and 30%, respectively, consistent with independent estimates (25, 29) and with the NO$_X$ and HCHO observations of Fig. 1 and SI Appendix, Fig. S1. This decreases mean NO$_X$ concentrations at MEE sites over the North China Plain by 69% in the model compared to 57% in the observations. Tropospheric NO$_X$ columns decrease by 71% in the model compared to 67% in the TROPOMI observations. Correcting the TROPOMI retrieval for the change in NO$_X$ vertical profile during the lockdown (SI Appendix, Fig. S4) implies a 68% decrease rather than 67%. Tropospheric HCHO columns decrease by 29% in the model compared to 13% in the observations and 17% in the observations corrected for the change in vertical profile (SI Appendix, Fig. S4). HCHO columns in the Beijing area actually show a mean increase during the lockdown, both in the model and the observations (SI Appendix, Fig. S1), reflecting the two aforementioned stagnation events with amplified oxidant chemistry as discussed below. A sensitivity model simulation indicates that NO$_X$ and HCHO columns over the North China Plain would have changed by –17% and +3%, respectively, over the 6-wk period if emissions had stayed constant.

We find that the model can reproduce the elevated PM$_{2.5}$, ozone, and PAN during the winter haze episodes in the lockdown period (Fig. 2). The sensitivity simulation with no decreases of NO$_X$ and VOC emissions during the lockdown period underestimates ozone by 10 to 20 ppb (SI Appendix, Fig. S5). Fig. 2D shows the model sources of HO$_X$ radicals driving the photochemistry. HONO is the dominant HO$_X$ source before lockdown, consistent with previous observational (13, 17, 18) and model studies (37, 38). The source of HONO in GEOS-Chem is mainly from heterogeneous NO$_X$ chemistry (39–41) plus a minor direct emission from transportation (42) and is therefore greatly reduced during the lockdown, consistent with observations (43). We find, on average, a 54% decrease of HONO concentrations during the lockdown in the surface to 500-m column. The HONO photolysis rate increases by 19% during the lockdown, mainly due to changes in meteorology. Consequently, OH radical from the HONO photolysis decreases by 41% (Fig. 2D).
We find that the dominant HOx source during the lockdown is the photolysis of HCHO, which is much larger than prelockdown period despite the decrease in VOC emissions. The HOx source from HCHO photolysis is particularly large during the two haze events of January 28 and February 12, reflecting the stagnant conditions (SI Appendix, Fig. S6). The model includes primary HCHO emissions from the transportation and residential sectors (Materials and Methods), but we find that 95% of HCHO during lockdown is secondary from oxidation of VOCs. That fraction is 88% before lockdown, consistent with observational studies that find most of the HCHO in wintertime Beijing to be secondary (44, 45). Reactive alkenes and aromatics dominate the HCHO source and drive ozone production (SI Appendix, Fig. S7). We find in the model that HCHO acts as a photochemical amplifier whereby increased production of HCHO under stagnant weather conditions leads to higher OH concentrations and hence accelerates the oxidation of VOCs by OH. This radical amplification process combined with decreased NOx emissions allows for fast ozone production. The faster oxidation of VOCs also drives large increases of PAN in the model.

Growing Ozone Pollution during the Winter Haze Season, 2013 to 2019. The experience of the COVID-19 lockdown dramatically illustrates how the continuing decrease of NOx emissions in China under the Clean Air Action may lead to high-ozone episodes during the winter haze season. National NOx emissions have decreased by 30% over the 2013 to 2019 period (9, 36, 46, 47). Fig. 3 shows an increasing trend of maximum MDA8 ozone in March from 2013 to 2019, demonstrating the growing northward spread of ozone pollution into the winter haze season, with extensive exceedances of the air quality standard. PM2.5 decreases over the same period, but the daily correlation between PM2.5 and ozone is increasingly positive (Fig. 3), implying fast ozone production with simultaneously elevated PM2.5 during stagnant conditions. The same trend toward rapidly increasing ozone production from 2013 to 2019 is also observed in February (SI Appendix, Fig. S8), with ozone exceeding 75 ppb in the North China Plain by 2019 and the ozone–PM2.5 correlation switching from negative to positive. Odd oxygen over the North China Plain in 2019 suggests an onset of ozone production in mid-February (SI Appendix, Fig. S2).

Fig. 4 shows the seasonality of observed 2013 to 2019 ozone trends in the North China Plain after filtering of interannual meteorological variability with a stepwise multiple regression model (Materials and Methods) to better resolve the effect of anthropogenic emission changes (6, 8, 9). Trends in the original unfiltered data are shown in SI Appendix, Fig. S9. We find that
the ozone increase is largest in March and that the increase in February is comparable to that in summer. Fig. 4B shows the change in the observed frequency of ozone exceedances of the 160 μg m\(^{-3}\) air quality standard (days month\(^{-1}\)) for individual months between 2014 to 2016 and 2017 to 2019 for the 22 cities in the North China Plain with full records over the 6-y period. Exceedances are not yet observed in February, but exceedances in March rose to 2 days month\(^{-1}\) in the 2017 to 2019 period.

We conducted GEOS-Chem simulations with 2013 and 2019 emissions to understand the effect of changes in anthropogenic emissions on the seasonality of ozone increase in the North China Plain (Materials and Methods). NO\(_x\) emissions in the MEIC inventory for the North China Plain decreased by 30% from 2013 to 2019, VOC emissions stayed flat, and observed PM\(_{2.5}\) concentrations decreased by about 50% in all seasons (8, 9, 36). Model results shown in Fig. 4A indicate that these changes led to ozone increases in all months of the year, peaking in spring and fall. We previously found in GEOS-Chem that the PM\(_{2.5}\) decrease was the principal driver for the ozone increase in summer due to decreased scavenging of HO\(_2\) radicals assuming a reactive uptake coefficient of 0.2 (6, 33). Here, we reproduce this result and find that the ozone trend in July would have been negative if not for the PM\(_{2.5}\) decrease. However, we also find that the PM\(_{2.5}\) effect is largely limited to summer. In other seasons, oxidation of NO\(_x\) to nitric acid is the dominant HO\(_2\) sink (SI Appendix, Fig. S10). This explains the maximum ozone increases in the model in spring and fall when photochemistry is active but strongly VOC-limited (16).

The model underestimates the observed ozone increase in all seasons except in the fall. The MEE sites are generally located in urban centers, where VOC-limited conditions are strongest and may not be captured at the 25-km resolution of GEOS-Chem (6, 48). Similarly, wintertime increases in ozone due to decreased local titration by NO emissions would not be well represented at the 25-km model scale. There is additional uncertainty in the HO\(_2\) reactive uptake coefficient and how it may vary with PM\(_{2.5}\) composition (49). The model underestimate of the springtime trend may reflect an increase in background ozone, which is particularly important in that season (50). Ozoneonde measurements over the 2013 to 2018 period in Beijing (51) show an increase in free tropospheric ozone in spring but not in other seasons (SI Appendix, Fig. S11), which may be due to rising anthropogenic emissions in South and Southeast Asia (52).

Implications for Air Pollution Control. The extension of the ozone pollution season into winter–spring is observed not only for the North China Plain but across China (SI Appendix, Fig. S12). Regulatory attention has so far focused on summertime ozone pollution (53), but we find that the rate of ozone increase is larger in February to May months and that exceedances of the air quality standard can already occur as early as March in the North China Plain. Ozone is rapidly becoming a year-round air pollution problem in China, and one may expect more frequent occurrences of joint ozone–PM\(_{2.5}\) pollution episodes. For example, a pollution event with MDA8 ozone of 140 ppb and 24-h PM\(_{2.5}\) of 100 μg m\(^{-3}\) averaged over all Beijing MEE sites was observed on April 30 to May 1, 2020. Beyond public health, this broadening of the ozone pollution season threatens crop production (54), particularly for the spring growth of winter wheat in North China, which accounts for two-thirds of China’s total wheat yield (55).

The fast increase of ozone and HO\(_x\) radicals outside of summer would stimulate the formation of secondary PM\(_{2.5}\), including nitrate, sulfate, and organic components. Long-term PM\(_{2.5}\) composition measurements in Beijing show an increase in the contributions of secondary species to total PM\(_{2.5}\) (30, 56). Increased production of secondary PM\(_{2.5}\) has been blamed for the occurrence of haze pollution episodes during the COVID-19 lockdown (29).

This rapid broadening of the ozone pollution season in China can be explained by the fast decrease of NO\(_x\) emissions combined with very high and flat VOC emissions. VOC oxidation produces carbonyls, in particular HCHO, whose photolysis produces HO\(_x\) radicals that drive fast ozone production and accelerate further VOC oxidation. This process has been previously reported to occur in oil/gas fields in the United States in winter (12), and here we see that it operates in urban China as well. Increasing background ozone in spring (51, 52, 57) could also contribute to the increase in surface ozone pollution.

NO\(_x\) emission controls in China have been motivated mainly by the goal of decreasing nitrate PM\(_{2.5}\), and further controls are expected in the future (http://env.people.com.cn/n1/2020/0515/c1010-31710781.html). The Chinese government has recently announced controls on VOC emissions in June to September months in order to decrease ozone pollution (53). Our results suggest that extending these VOC emission controls year round would avoid further spread of ozone pollution outside the summer season.

In summary, we have found that winter haze episodes in the North China Plain following the COVID-19 lockdown in January 2020 featured fast photochemical production of ozone with concentrations reaching 60 to 70 ppb. This can be explained by the sharp decrease in NO\(_x\) emissions combined with high VOC emissions, driving ozone production through HCHO photolysis. This rapid ozone production during the COVID-19 winter haze episodes in January to February illustrates a more general trend from 2013 to 2019 of increasing ozone pollution in winter–spring, driven by a 30% decrease of NO\(_x\) emissions while VOC emissions...
have stayed flat. Curbing ozone pollution in China will require VOC controls year round as NOx emissions continue to decrease. Such controls will benefit not just the North China Plain but also the country as a whole.

Materials and Methods

Observations. Surface hourly concentrations of ozone, PM2.5, and CO concentrations are reported by the MEE at a network that started from ~450 sites in 2013 and grew to ~1,500 sites by 2020 and included about 360 cities. We accessed the data through http://beijingair.sinaapp.com. We computed MD8 ozone concentrations and 24-h average concentrations for other air pollutants from the hourly data for the 2013 to 2020 period. The reported concentrations are in the unit of μg m⁻³ for gas species under standard conditions of temperature and pressure (STP; 273 K, 1,013 hPa) until August 31, 2018. This reference state was changed to (298 K, 1,013 hPa) on September 1, 2018. We converted concentrations to ppb following ref. 9. We also used PAN concentrations measured by researchers from the China Meteorological Administration at a site on the campus of Minzu University of China (39.95°N, 116.32°E) in urban Beijing. The measurements were made with an online gas chromatograph equipped with an electron capture detector (58).

Satellite observations of NO2 (59) and HCHO (60) columns from the TROPOMI instrument were accessed from https://s5phub.copernicus.eu/dhus/. TROPOMI provides daily global coverage with 5.5 × 3.5 km² pixel resolution. TROPOMI data have been applied in tracking anthropogenic emission changes during the lockdown (26, 61). We used the TROPOMI Level 2 observations with quality assurance values larger than 0.75 for NO2 (version 1.3.2) and larger than 0.5 for HCHO (version 1.1.8).

Chemical Transport Model Simulations.

GEOS-Chem model description. We simulated air quality over China by using the GEOS-Chem atmospheric chemistry model (version 12.7.1; http://www.geos-chem.org). The model includes detailed ozone–NOx–VOC–aerosol–halogen tropospheric chemistry (62, 63) and is driven by NASA GEOS-FP (Forward Processing) meteorological data (64) with 0.25° × 0.3125° horizontal resolution. We use that native resolution over a nested eastern China domain (30 to 45°N; 108 to 125°E). Chemical boundary conditions at the edges of the nested domain are updated every 3 h from a global simulation with 4° × 5° resolution.

The GEOS-Chem simulation of ozone pollution over China has been evaluated in a number of recent studies (6, 50, 65). Our model configuration largely follows that of ref. 33, but we use a higher spatial resolution. The standard GEOS-Chem model includes HONO and HCHO formed from NO2 heterogeneous chemistry and VOC oxidation, respectively (39, 66). Here, we added HONO emissions from transportation by applying a HONO/NOx emission ratio of 1.7% following ref. 42, and we multiplied transportation HCHO emissions in the standard model by a factor of 5 for the cold season (November to March) following refs. 67 and 48. These direct emissions made minor contribution to HONO and HCHO as compared to chemical production, 2013 to 2019 simulations. To quantify the effect of changing 2013 to 2019 emissions on ozone, we performed a set of simulations with fixed meteorology (September 2018 to August 2019) and perturbed emissions, using 6 mo of initialization. Anthropogenic emissions over China are from the MEIC, including seasonal (month-to-month) variability (36). MEIC emissions are available for the 2013 to 2017 period and were accessed from http://www.meicmodel.org. NOx and SO2 emissions for the 2017 to 2019 period were scaled on the basis of national emission trends reported by the MEE (46, 47). CO and PM2.5 primary emissions for the 2017 to 2019 period were scaled on the basis of the MEE network data. In our nested eastern China domain, the decreases of emissions from 2013 to 2017 and from 2017 to 2019 are, respectively, 23% and 8% for NOx, 62% and 11% for SO2, 27% and 13% for CO, and 34% and 15% for PM2.5. Anthropogenic emissions of ammonia (mainly from agriculture) and VOCs show no significant trend from 2013 to 2017 in the MEIC inventory. The lack of trend in VOC emissions is consistent with satellite HCHO data over eastern China showing no significant trend over the 2013 to 2019 period (9). Thus, we assume constant anthropogenic emissions of ammonia and VOCs from 2017 to 2019.

Simulations with Chinese anthropogenic emissions for years 2013 and 2019 were conducted to compute the 2013 to 2019 ozone trend in different months due to anthropogenic emission changes. A sensitivity simulation changing only anthropogenic NOx and VOC emissions was also conducted to isolate the effect of PM2.5 changes (33) by difference.

Effect of COVID-19 lockdown. To simulate ozone changes caused by the COVID-19 lockdown starting on January 24, 2020, we conducted GEOS-Chem simulations from January 1, 2020, to February 15, 2020, after 3 mo of initialization,
The PM2.5 effect (33) is isolated in the model by removing the effect of emissions, both using the same September 2018 to August 2019 meteorological conditions and by removing the effect of interannual meteorological variability with a calculated by linear regression on the monthly mean values at MEE sites for more details. All observed ozone trends are statistically significant at the 95% confidence level, except for July and November, which have P values of 0.07 and 0.06, respectively. (B) The number of days per month with observed ozone levels above the air quality standard (160 μg m⁻³, corresponding to 75 ppb at STP), averaged over the 2014 to 2016 and 2017 to 2019 periods for the 22 MEE network cities in the North China Plain with full 2014 to 2019 records.

with the same model settings as the 2013 to 2019 simulations. Prelockdown anthropogenic emissions before January 24 were scaled from 2019 by extending the rate of 2017 to 2019 reductions to 2020. Relative to the pre-lockdown period, NOx and VOC emissions over the simulation domain were reduced by 60% and 30%, respectively. This was done by applying reduction ratios of 90% for transportation emissions, 20% for power plants, 60% for other industry (distilleries, chemical plants, and IPM agricultural processing), and no changes for the residential and agriculture sectors. These reduction ratios are estimated from economic data in January to February of 2020 (68) and are consistent with the surface and satellite data for NOx and HCHO shown in Fig. 1 and SI Appendix, Fig. S1. Other studies have estimated emission reductions of 50 to 60% for NOx and 30 to 40% for VOCs during the lockdown period (25, 29). We apply the emission reductions from January 24 to February 15 when the lockdown policy was most severe (25, 29, 32). The sector-based reductions also result in model emission decreases for SO2, CO, and PM2.5 of 40%, 39%, and 22%, respectively. Mean photoysis rates of NO2, HONO, and HCHO in the surface to 500-m model column increased, respectively, by 19%, 19%, and 24% between January 1 to 23 and January 24 to February 15, mainly reflecting changes in meteorology.

To quantify the role of meteorology in driving air quality change during the lockdown, a sensitivity simulation with no emission reductions was conducted.

Removing Meteorological Influence on Ozone Trends. To isolate the role of anthropogenic emission changes in driving 2013 to 2019 ozone trends, we removed the effect of interannual meteorological variability by using the stepwise multiple linear regression (MLR) model following refs. 6 and 9. Nine meteorological variables from the Modern-Era Retrospective analysis for Research and Applications, version 2 (MERRA-2) database (69), including temperature, 10-m zonal and meridional wind speed, 850-hPa meridional wind speed, relative humidity, boundary layer height, cloud cover, rainfall, and sea level pressure (6), were considered as candidate ozone covariates on the 0.5° × 0.625° MERRA-2 grid. We applied the MLR model to fit the day-to-day variability of MD8 ozone independently for each site and each season. To avoid overfitting, only the three locally dominant meteorological parameters were regressed onto the final fit for the meteorologically-driven 2013 to 2019 trend in monthly MD8 ozone. The residual trend was then taken to reflect the role of anthropogenic emissions following refs. 6 and 9.

Data Availability. The measurements, GEOS-Chem model code, and reanalysis data in this study are publicly available for download. Surface measurements of hourly air pollutants from China’s MEE sites can be downloaded from http://beijingair.sinaapp.com. The TROPOMI satellite data can be freely accessed from https://5phub.copernicus.eu/dhus/. The anthropogenic emission inventory is available from http://www.meicmodel.org. The GEOS-Chem model code is open source (https://doi.org/10.5281/zenodo.3676008). All other study data are included in the article and/or SI Appendix.

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