



1 **Assessment of meteorology vs control measures in China fine**
2 **particular matter trend from 2013-2019 by an environmental**
3 **meteorology index**

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12 **Abstract**

13 A framework was developed to quantitatively assess the contribution of meteorology
14 variations in the trend of particular matter (PM) concentrations and to separate the
15 impacts of meteorology from the control measures in the trend, based upon an
16 Environmental Meteorology Index (EMI). The model-based index EMI realistically reflects
17 the role of meteorology in the trend of PM and is attributed into three major factors:
18 deposition, vertical accumulation and horizontal transports. Based on the 2013-2019
19 PM_{2.5} observation data and re-analysis meteorological data in China, the contributions of
20 meteorology and control measures in nine regions of China were assessed separately by
21 the EMI-based framework. Monitoring network observations show that the PM_{2.5}
22 concentrations have been declined about 50% on national average and about 35% to 53%
23 for various regions. It is found that the nation-wide emission control measures were the
24 dominant factor in the declining trend of China PM_{2.5} concentrations, contributing to
25 about 47% of the PM_{2.5} decrease from 2013 to 2019 on the national average and 32% to



1 the 52% for various regions. The meteorology has a variable and sometimes critical
2 contribution to the year by year variations of $PM_{2.5}$ concentrations, 5% on annual average
3 and 10-20% for the fall-winter heavy pollution seasons.

4 **1. Introduction**

5 Recent observation data from the Ministry of Ecology and Environment of China (MEE)
6 has shown a steady improvement of air quality across the country, especially in particular
7 matter (PM) concentrations (Hou et al., 2019). According to 2013-2019 China Air Quality
8 Improvement Report issued by MEE, compared to 2013, the average concentration of
9 particulate matter with an aerodynamic diameter of less than $2.5 \mu m$ ($PM_{2.5}$)
10 concentrations in 74 major cities of China decreased by more than 50% in 2019. From
11 scientific and management point of views, a quantitative apportionment of the reasons
12 behind the trend is critical to assess the reduction strategies implemented by the
13 government and to guide future air quality control policy. However, the assessment of the
14 improvements of air quality is a complicated process that involves the quantification of
15 changes in the emission sources, meteorological factors, and other characteristics of the
16 $PM_{2.5}$ pollution, which are also interacting with each other. In order to separate the
17 degree of these factors, a comprehensive analysis, including observational data and
18 model simulation, is needed.

19 Researches have been done extensively on the impacts of weather systems on air
20 quality. Synoptic and local meteorological conditions have been recognized to influence



1 the PM concentrations at various scales (Beaver and Palazoglu, 2006;He et al., 2017a;He
2 et al., 2017b;Pearce et al., 2011a;Pearce et al., 2011b). For the atmospheric aerosol
3 pollution in eastern China, the dynamic effect of the downdraft in the "leeward slope" and
4 "weak wind area" of the Qinghai Tibet Plateau in winter is not conducive to the diffusion
5 of air pollution emissions in the urban agglomerations of eastern China (Xu et al., 2015;Xu
6 et al., 2002). The evolution of circulation situation is an important factor driving the
7 change of haze pollution (He et al., 2018). The local circulations, such as mountain and
8 valley wind and urban island circulation, have significant impact on local pollutant
9 concentration (Chen et al., 2009;Yu et al., 2016). Previous studies also revealed that PM_{2.5}
10 concentration is significantly correlated with local meteorological elements, such as
11 temperature, humidity, wind speed, and boundary layer height (He et al., 2017b;Bei et al.,
12 2020;Ma et al., 2019;He et al., 2016).

13 In the Beijing-Tianjin-Hebei (BTH) Region, a correlation analysis and principal
14 component regression method (Zhou et al., 2014) was used to identify the major
15 meteorological factors that influenced the API (Air Pollution Index) time series in China
16 from 2001-2010, indicating that air pressure, air temperature, precipitation and relative
17 humidity were closely related to air quality and resulting in a series of regression
18 formulas. Yet, the analysis was assumed a relatively unchanged emission whose impacts
19 were not taken into account. On a local scale, an attempt (Zhang et al., 2017) has been
20 made to correlate the air pollutant levels with a combination of meteorological factors
21 with the development of the Stable Weather Index (SWI) at CMA. The SWI is a composite
22 index which includes the advection, vertical diffusion and humidity and other



1 meteorological factors that are related to the formation of air pollutions in a specific
2 region or city. A higher value of SWI means a weaker diffusion of air pollutants. This index
3 had some success in assessing the meteorological impacts on air pollution, especially
4 calibrated for a specific region, i.e. Beijing. However, when applied to different areas
5 where the emission patterns and meteorological features are different, this index failed to
6 give a universal or comparable indication of meteorological assessment of pollution levels
7 across the nation.

8 Using the Kolmogorov-Zurbenko (KZ) wave filter method, Bai et al (2015) made an
9 effort to break the API time series in three Chinese cities into short-term, seasonal and
10 long-term components, and then used the stepwise regression to set up API baseline and
11 short-term components separately and establish linear regression models for
12 meteorological variables of corresponding scales. Consequently, with the long-term
13 representing the change of emissions removed from the time series, the meteorological
14 contributions alone were assumed and analyzed, pointing out that unfavorable conditions
15 often lead to an increase by 1-13 whereas the favorable conditions to a decrease by 2-6 in
16 the long-term API series, respectively. Though the contributions of emissions and
17 meteorological variations were separated by the research, it is only done by mathematical
18 transformations and far from the reality. The mechanisms behind the variation of the time
19 series were not investigated.

20 A chemical transport model (CTM) is an ideal tool to carry the task of assessment by
21 taking the meteorology, emissions and processes into considerations altogether.



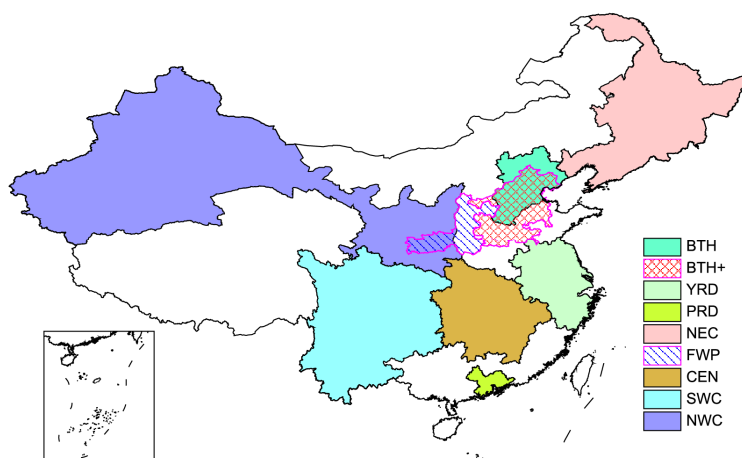
1 Andersson et al. (2007) used a CTM to study the meteorologically induced inter-annual
2 variability and trends in deposition of sulphur and nitrogen as well as concentrations of
3 surface ozone (O₃), nitrogen dioxide (NO₂) and PM and its constituents over Europe during
4 1958-2001. It is found that the average European interannual variation, due to
5 meteorological variability, ranges from 3% for O₃, 5% for NO₂, 9% for PM, 6-9% for dry
6 deposition, to about 20% for wet deposition of sulphur and nitrogen. A multi-model
7 assessment of air quality trends with constant anthropogenic emissions was also carried
8 out in Europe (Colette et al., 2011) and found that the magnitude of the emission-driven
9 trend exceeds the natural variability for primary compounds, concluding that that
10 emission management strategies have had a significant impact over the past 10 years,
11 hence supporting further emission reductions strategies. Model assessments of air quality
12 trends at various regions and time periods (Wei et al., 2017; Li et al., 2015) in China were
13 also done and yielded some useful results. For the BTH Region, Li et al. (2015) used the
14 Comprehensive Air Quality Model with extensions (CAMx) plus the Particulate Source
15 Apportionment Technology (PSAT) and simulated the contributions of emission changes in
16 various sectors and changes in meteorology conditions for the PM_{2.5} trend from 2006 to
17 2013. It was found that the change of source contribution of PM_{2.5} in Beijing and northern
18 Hebei was dominated by the change of local emissions. However, for Tianjin, and central
19 and southern Hebei province, the change of meteorology condition was as important as
20 the change of emissions, illustrating the regional difference of impacts by meteorology
21 and emissions. However, the emission changes in the simulations were assumed and did
22 not reflect the real spatio-temporal variations.



1 There is no surprise that previous studies could not systematically catch the
2 meteorological impacts across the whole nation as the controlling meteorological factors
3 involving the characteristics of planetary boundary layers (PBL), wind speed and turbulence,
4 temperature and stability, radiation and clouds, underlying surface as well as pollutant
5 emissions, vary greatly from region to region. A single index or correlation cannot be
6 applied to the entire nation. Obviously, in order to systematically assess the impacts of
7 meteorology on air pollution, these factors have to be taken into consideration in a
8 framework and be assessed simultaneously. This paper presents a methodology to assess
9 the individual impacts of meteorology and emission changes, based on a model-derived
10 index EMI, i.e., Environmental Meteorology Index, and observational data, providing a
11 comprehensive analysis of the air quality trends in various regions of China, with
12 mechanistic and quantitative attributions of various factors.

13 **2. Methodology**

14 The assessment is carried out through the combination of observational data and EMI
15 index from model analysis. Since the emission and air quality characteristics vary greatly
16 from region to region in China, the analysis is divided into 9 focused regions (Figure 1).
17 Regional air quality data ($PM_{2.5}$) provides the basis for the trend analysis. Separating the
18 trend contribution from regional emission reduction and meteorological variation needs a
19 framework, which is discussed below.



Note: **BTH**: Beijing, Tianjin and Hebei; **BTH+**: BJ, TJ + 26 cities; **YRD**: Shanghai, Jiangsu, Zhejiang and Anhui; **PRD**: 9 cities in Guangdong; **NEC**: Heilongjiang, Jilin and Liaoning; **FWP**: 11 cities in Shanxi, Shannxi and Henan; **CEN**: Hubei, Hunan and Jiangxi; **SWC**: Yunnan, Guizhou, Sichuan, Chongqing; **NWC**: Shannxi, Gansu, Ningxia and Xinjiang

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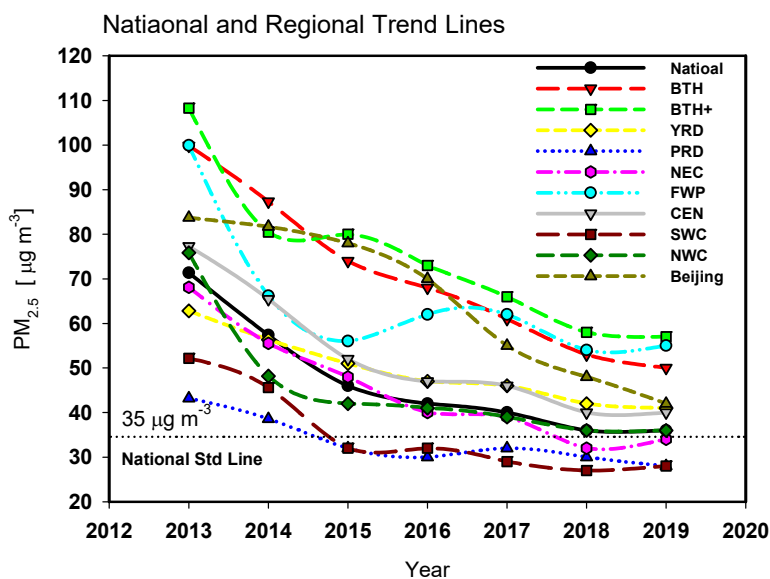
2 Figure 1: Analysis region separation and definition.

3 **2.1. Particular Matter (PM) Observation Data**

4 The observational pollution data of $PM_{2.5}$ concentrations used in this study were from
5 the monitoring network of the Ministry of Ecology and Environment (MEE) of China
6 ([http:// english.mee.gov.cn/](http://english.mee.gov.cn/)). From 2013 to 2019, the concentrations have shown a large
7 change in the country where most regions see a declined trend in the annual
8 concentrations. Data show that from 2013 to 2019, the national annual averaged $PM_{2.5}$
9 concentrations have dropped about 50% (Fig. 2), where the haze days have been
10 shortened by 21.2 days from the China Meteorological Administration (CMA) monitoring
11 data (Table 1), with some regional differences. Regionally, by 2019, the $PM_{2.5}$ reduction



- 1 rate from 2013 ranges from 35 to 53%. Detailed analysis will be given in the Results and
- 2 Discussion section.



3

4 Figure 2: National and regional trend lines of $PM_{2.5}$ in China from 2013 to 2019.

5 It is noted that the $PM_{2.5}$ mass concentrations by MEE are now reported under
6 observation site's actual conditions of temperature and pressure from September 1, 2018
7 before which the values were reported under the standard state (STP), i.e. 273 K and
8 101.325 kPa. In order to maintain the consistence of the data series, the $PM_{2.5}$
9 concentrations used in this study have all been converted according to the new standard
10 (MEE, 2012)(GB3095-2012) under actual conditions. Research has shown that after the
11 change of reporting standard, the $PM_{2.5}$ concentration in most cities decreased, and the
12 number of good days to meet the standard increased (Zhang and Rao, 2019).



1 2.2. Meteorological Data

2 Conventional meteorological data can provide qualitative assessment of the
3 contributions of meteorological factors to the changes of air quality. The data used in this
4 study are from 843 national base weather stations of the CMA from 2013 to 2019. The
5 wind speed (WS), day with small wind (DSW), relative humidity (RH) and haze days are
6 used to analyze the pollution meteorological conditions. When the daily average wind
7 speed is less than 2 m s^{-1} , a DSW day is defined. Since the haze formation is always related
8 to stable meteorological conditions and high aerosol mass loading, haze observation from
9 CMA is also used to analyze the haze trends and the impact of air quality on visibility. A
10 haze day is defined with daily averaged visibility less than 10 km and relative humidity less
11 than 85% (Wu et al., 2014), excluding days of low visibility due to precipitation, blowing
12 snow, blowing sand, floating dust, sandstorms and smoke.

13 Data show that from 2013 to 2019, the national annual averaged WS has increased
14 by 12.9%, DSW dropped by 15.1%, and RH almost unchanged (Table 1), with regional
15 differences. It can be seen that the annual haze days have a certain degree of correlations
16 positively with WS and negatively with DSW. Detailed analysis linking $\text{PM}_{2.5}$ and
17 meteorology will be given in the Results and Discussion secession.

18



- 1 Table 1: National and regional environmental meteorology in 2019 and comparisons with
- 2 2013 and 2016

Region	Wind Speed			Days with Small Wind			Relative Humidity			Haze (days)		
	AVG	vs 2016 (%)	vs 2013 (%)	Days	vs 2016 (%)	vs 2013 (%)	%	vs 2016 (%)	vs 2013 (%)	Days	vs 2016	vs 2013
National	1.9	+3.6	+12.9	129.8	-0.8	-15.1	60.1	-6.3	-3.9	25.7	-12.1	-21.2
BTH	1.9	-2.7	-2.2	131.0	+5.8	+9.0	56.7	-3.3	-4.2	45.2	-15.2	-26.1
BTH+	2.1	-2.2	+0.8	114.4	+4.3	-5.6	58.3	+0.2	+0.6	54.5	-21.2	-30.3
YRD	2.1	-2.2	-4.7	114.1	+1.9	+5.2	76.3	-1.5	+5.5	34.0	-36.9	-54.9
FWP	2.0	-1.6	+10.9	122.8	+7.2	-25.2	59.9	-1.5	+3.3	51.6	-32.7	-43.8
PRD	2.0	+0.9	-10.4	118.5	+12.7	+14.4	79.7	-3.0	+10.3	3.1	-6.4	-34.3
NEC	2.7	+3.6	+12.9	55.8	-10.1	-38.4	61.6	-3.4	-5.8	13.6	-19.4	-12.4
CEN	1.8	-2.1	+0.4	172.1	+2.2	-2.8	77.9	+0.4	+6.9	30.3	-19.7	-23.2
SWC	1.7	+4.4	+12.2	180.7	-6.9	-16.3	74.7	+1.5	+5.7	11.1	-13.7	-12.4
NWC	1.9	-3.3	+4.3	146.8	+6.3	-9.5	58.5	-0.1	+2.8	20.2	-9.6	-6.6

- 3 Note: "+" increased; "-" decreased



1 **2.3. EMI – the Environmental Meteorological Index**

2 Due to the complicated interactions of emissions, meteorology and atmospheric
3 processes, a single set of meteorological factors or a combination of them cannot
4 quantitatively attribute the individual factor to the changes of concentration
5 observed.

6 In order to quantitatively assess the impacts of meteorological conditions to the
7 changes of air pollution levels, an index EMI (Environmental Meteorological Index) is
8 defined as follows. For a defined atmospheric column (h) at a time t , an EMI is
9 defined as an indication of atmospheric pollution level:

$$10 \quad \text{EMI}(t) = \text{EMI}(t_0) + \int_{t_0}^t \Delta \text{EMI} * dt$$

11 (1)

12 where the ΔEMI is the tendency that causes the changes of pollution level in a time
13 interval dt defined as:

$$14 \quad \Delta \text{EMI} = i\text{Emid} + i\text{Tran} + i\text{Accu} \quad (2)$$

15 where the $i\text{Emid}$ is the difference between emission and deposition, and $i\text{Tran}$ and
16 $i\text{Accu}$ are the net (in minus out) advection transports and the vertical accumulation
17 by turbulent diffusion in the column, respectively. A positive sing of each factor
18 indicates a net flow of pollutants into the column, and vice visa.

19 Mathematically, these factors are expressed as:

$$20 \quad i\text{Tran} = \frac{1}{C_0} \int_0^h \left(u \frac{\partial C}{\partial x} + v \frac{\partial C}{\partial y} + w \frac{\partial C}{\partial z} \right) dz$$



1

$$2 \quad iAccu = \frac{1}{C_0} \int_0^h \left[\frac{\partial C}{\partial x} \left(K_x \frac{\partial C}{\partial x} \right) + \frac{\partial C}{\partial y} \left(K_y \frac{\partial C}{\partial y} \right) + \frac{\partial C}{\partial z} \left(K_z \frac{\partial C}{\partial z} \right) \right] dz$$

$$3 \quad iEmid = \frac{1}{C_0} \int_0^h [Emis - (V_d + L_d)] dz$$

4

(3)

5 where the tendency is normalized by a factor C_0 . For an application of EMI to the
6 $PM_{2.5}$, C_0 is set to equal $35 \mu\text{g m}^{-3}$, the national standard for $PM_{2.5}$ in China (MEE,
7 2012), and the $EMI(t)$ is written as $EMI(t)_{2.5}$. If the $EMI_{2.5}$ is less than 1, the
8 concentration level will reach or be better than the national standard.

9 It can be seen here that these key parameters account for the major
10 meteorological factors which control the air pollutant levels, including wind speed
11 and directions (u, v, w), turbulent mixing (K_x, K_y, K_z) as well as dry and wet
12 depositions (V_d and L_d). Therefore, under the conditions of an unchanged emissions
13 ($Emis$), the EMI variation reflects the impacts of meteorological factors on the levels
14 of atmospheric pollutants. Furthermore, because of the inclusion of individual
15 factors such as $iTran$, $iAccu$ and $iEmid$, the variation of $EMI(t)_{2.5}$ can be attributed to
16 the variation of each factor, which gives more detailed information on the
17 meteorological influence to the ambient pollutant concentration variations.

18 For a period of time p (t_0 to t_1) when the averaged pollutant level (e.g. $PM_{2.5}$)
19 is compared with $EMI(t)_{2.5}$, the time integral has to be done to obtain the averaged
20 index for the period, such as:

$$21 \quad \overline{EMI(p)_{2.5}} = \frac{1}{t_1 - t_0} \int_{t_0}^{t_1} EMI(t)_{2.5} dt$$



$$\begin{aligned} \overline{EMI(p)}_{2.5} &= \frac{1}{t_1 - t_0} [nEMI(t_0)\Delta t + n\Delta EMI(1)\Delta t^2 + (n-1)\Delta EMI(2)\Delta t^2 \\ &+ (n-2)\Delta EMI(3)\Delta t^2 + (n-3)\Delta EMI(4)\Delta t^2 + \dots + \Delta EMI(n)\Delta t^2] \end{aligned} \quad (5)$$

where n is the time steps in the period and the averaged EMI has been linked to the starting point $EMI(t_0)$ and the changing rates of EMI, i.e. $\Delta EMI(n)$, at each time step.

2.4. Assessment Framework

The $EMI_{2.5}$ index provides a way to assess the meteorological impacts on the changes of $PM_{2.5}$ concentrations at two time periods, i.e. January 2103 (p_0) and January 2016 (p_1) under the assumption of unchanged emissions. However, due to the national efforts of improving air quality, the year-by-year emissions are changing rapidly and unevenly across the country. The changes in both emissions and meteorology tangled together to yield the observed changes in ambient concentrations. For policy makers, the emission reduction quantification is critical to guide the further air quality improvements. The framework proposed here is to combined changes in the observed concentration levels and meteorology factors $\overline{EMI(p)}_{2.5}$ to quantify the changes in emission changes at two time periods.

The observed concentrations at p_0 and p_1 are defined as PM (m_0, e_0) and PM (m_1, e_1) where (m_0, e_0) and (m_1, e_1) indicate the meteorology and emission status at p_0 and p_1 , respectively. The ratio of $\overline{EMI(p_0)}_{2.5} / \overline{EMI(p_1)}_{2.5}$ reflects the impact ratio of sole meteorology variations on the concentrations from p_0 to p_1 . Therefore:



$$1 \quad \frac{PM(m0, e1)}{PM(m1, e1)} = \frac{\overline{EMI(p0)}_{2.5}}{\overline{EMI(p1)}_{2.5}} \quad (6)$$

2 which gives an assumed concentration of pollutant under the conditions of
3 unchanged meteorology at $p0$ but with new emission at $p1$. In a simple statement,
4 the PM ($m0, e1$) is the concentration at $p1$ only under the influence of emission
5 change from $e0$ to $e1$ but with the same meteorology ($m0$). Consequently, the
6 impact of only emission changes from $e0$ to $e1$ on the concentration changes can be
7 expressed as:

$$8 \quad \Delta EMIS = \frac{PM(m0, e1) - PM(m0, e0)}{PM(m0, e0)} \times 100\% \quad (7)$$

9 which can be used to assess the impact of emission changes (control measures) on
10 the air pollutants.

11 **2.5. Quantitative Estimate of EMI**

12 Finally, a process-based method is developed to calculate the EMI and its
13 components, i.e. *iEmid*, *iTran* and *iAccu*. The main modeling frame-work used is the
14 chemical weather modeling system MM5/CUACE, which is a fully coupled
15 atmospheric model used at CMA for national haze and air quality forecasts (Gong
16 and Zhang, 2008; Zhou et al., 2012). CUACE is a unified atmospheric chemistry
17 environment with four major functional sub-systems: emissions, gas phase
18 chemistry, aerosol microphysics and data assimilation (Niu et al., 2008). Seven aerosol
19 components, i.e. sea salts, sand/dust, EC, OC, sulfates, nitrates and ammonium salts
20 are sectioned in 12 size bins with detailed microphysics of hygroscopic growth,
21 nucleation, coagulation, condensation, dry depositions and wet scavenging in the



1 aerosol module (Gong et al., 2003). The gas chemistry module is based on the
2 second generation of Regional Acid Deposition Model (RADM II) mechanism with 63
3 gaseous species through 21 photo-chemical reactions and 121 gas phase reactions
4 applicable under a wide variety of environmental conditions especially for smog
5 (Stockwell et al., 1990) and prepares the sulfate and SOA production rates for the
6 aerosol module and for the aerosol equilibrium module ISORROPIA (Nenes et al.,
7 1998) to calculate the nitrate and ammonium aerosols. This is the default method to
8 treat the secondary aerosol formations in CUACE. For the EMI application of CUACE,
9 another option was also adapted to compute the secondary aerosol formations by a
10 highly parameterized method (Zhao et al., 2017), that computes the aerosol
11 formation rates directly from the pre-cursor emission rates of SO₂, NO₂ and VOC.
12 This option was added to facilitate timely operational forecast requirements for
13 CMA. Both primary and pre-cursor emissions of PM are based on the 2016 MEIC
14 Inventory (<http://www.meicmodel.org/>) developed by Tsinghua University for China.

15 In order to quantitatively obtain each term defined in Equation 3, the CUACE
16 model was modified to extract the change rates for the processes involved. Driven
17 by the re-analysis meteorological data, the new system CUACE/EMI can be used to
18 calculate each term in ΔEMI at each time step (Δt).

19 In summary, this section presents a systematic platform to separate and assess
20 the impacts of the meteorology and emissions on the ambient concentration
21 changes. The $\overline{EMI(p)}_{2.5}$ and ΔEMI form the basis for the assessment. In the Results
22 and Discussions section, the application of the platform is presented to assess the
23 fine particular matter (PM_{2.5}) changes in China.

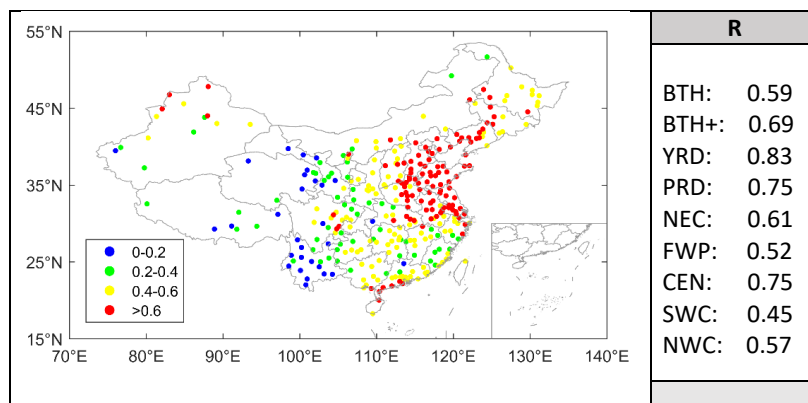


1 **3. Results and Discussions**

2 **3.1. Validation of EMI by Observations**

3 Under the conditions of no changes in annual emissions for PM_{2.5} and its
4 precursors, the daily EMI_{2.5} was computed by CUACE from 2013 to 2019 on a 15×15
5 km resolution across China and accompanied by its contribution components: *iTran*,
6 *iAccu* and *iEmid*. However, in order to reflect the significant changes of industrial and
7 domestic energy consumptions within a year in China, a monthly emission (Wang et
8 al.) variation was applied to the emission inventory for computing the EMI_{2.5}, which
9 is more realistically reflecting the meteorology contributions to the PM_{2.5}
10 concentrations.

11 To evaluate the applicability of EMI_{2.5}, the index was compared with the
12 observed PM_{2.5} concentrations. Figure 4 shows the spatial distribution of correlation
13 between PM_{2.5} and EMI_{2.5} for 2017 for all China. The correlation coefficients between
14 EMI_{2.5} and PM_{2.5} concentrations are greater than 0.4 for most of the Eastern China
15 and greater than 0.6 for most of the assessment regions. Less satisfactory correlation
16 was found in Western China, possibly due to complex terrain and less accurate
17 emission data over there. Furthermore, due to the uncertainty in emissions and the
18 difference in model performance for year-to-year meteorology simulations, the
19 correlation coefficients may differ for different years. Overall, the good correlation
20 between them merits the application of EMI_{2.5} to quantify the meteorology impact
21 on PM_{2.5}.



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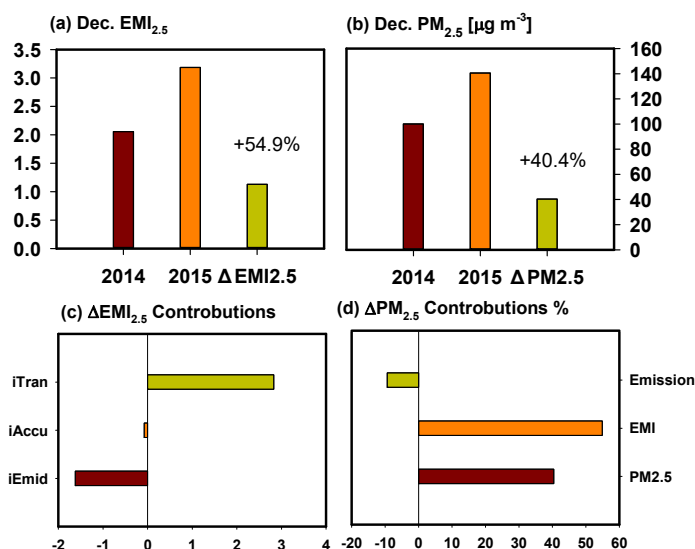
2 Figure 4: Correlation coefficients (R) between the $EMI_{2.5}$ and the observed $PM_{2.5}$
3 concentrations across China for 2017 and for typical regions averaged between 2013 and
4 2019.

5

6 To further illustrate the applicability of $EMI_{2.5}$, the difference of various
7 conditions between December 2014 and December 2015 in BTH region was also
8 analyzed when significant change of air quality and meteorological conditions
9 occurred. The winter of 2015 was accompanied by a strong El Niño (ENSO) event,
10 resulting in significant anomalies for meteorological conditions in China. Analysis
11 shows that the meteorological conditions in December 2015 (compared to
12 December 2014) had several important anomalies, including that the surface
13 southeasterly winds were significantly enhanced in the North China Plain (NCP) and
14 the wind speeds were decreased in the middle-north of eastern China, while slightly
15 increased in the south of eastern China. Study suggests that the 2015 El Niño event
16 had significant effects on air pollution in eastern China, especially in the NCP region,
17 including the capital city of Beijing, in which aerosol pollution was significantly
18 enhanced in the already heavily polluted capital city of China (Chang et al., 2016).



1 Figure 5 shows the monthly average $EMI_{2.5}$, $PM_{2.5}$ and the contribution of
2 sub-index to total $EMI_{2.5}$ in December 2014 and 2015 over BTH region. The monthly
3 average $EMI_{2.5}$ increases about 54.9% from 2.1 in December 2014 to 3.2 in December
4 2015, indicating worsening meteorological conditions for $PM_{2.5}$ pollution. The
5 increase of $EMI_{2.5}$ is mainly contributed by adverse atmospheric transport conditions
6 (Fig. 5c), which results in the increase of $EMI_{2.5}$ reaching 3.2. With the increase of
7 background concentration, the deposition and vertical diffusion also increase, and
8 offset the impact of adverse transport conditions to some extent.



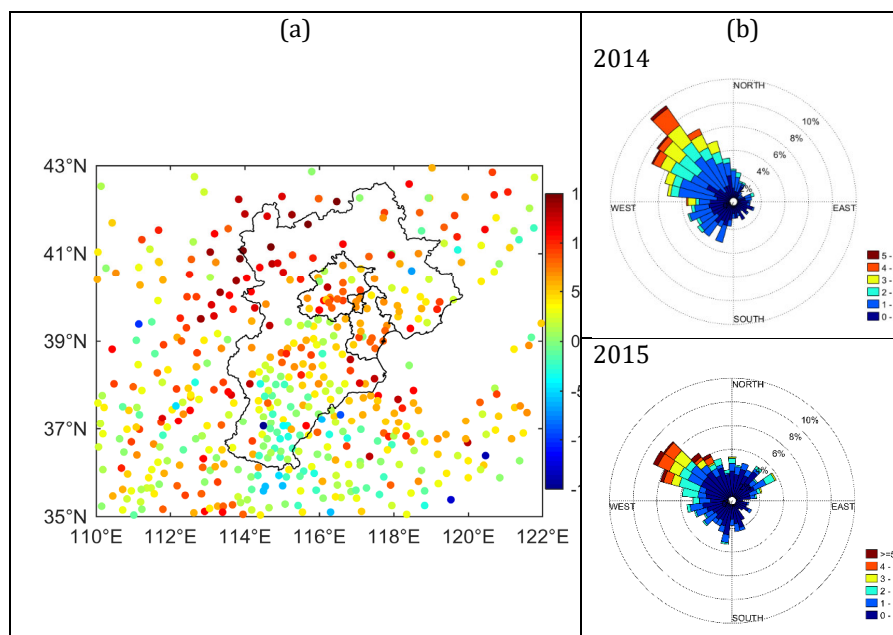
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10 Figure 5: The monthly average $EMI_{2.5}$ (a), $PM_{2.5}$ (b) and the contribution of sub-index (c) and
11 contributions to the $PM_{2.5}$ changes (d) in Decembers 2014 and 2015 over BTH.

12

13 The worsening meteorological conditions represented by $EMI_{2.5}$ were also
14 supported by the observations for the two periods. The observed day with small
15 wind (DSW, wind speed less than 2 m s^{-1}) reveals that, except for part of southern



1 Hebei province, the DSW increases 5-15 days for 2015 in most meteorological
2 stations in BTH region (Fig. 6a), which indicates a large decrease of local diffusion
3 capability. The comparison of wind rose map shows that the decrease of northwest
4 wind and the increase of southwest and northeast wind occurred in December 2015
5 (Fig. 6b). The change of wind fields indicates more pollutants were transported to
6 BTH region from Shandong, Jiangsu, Henan, and Northeast China. These variations
7 indirectly validate the conclusions of adverse atmospheric transport conditions with
8 high iTran in December 2015.



9
10 Figure 6: (a) The change of DSW (days) from December 2014 to December 2015 (December
11 2015 – December 2014) and (b) Wind rose maps in December 2014 and December 2015
12 over BTH region.

13

14 Based on the assessment method of emission contribution to the observed
15 trend (Eqs. 6 and 7), the emissions reduction in December 2015 as compared to 2014



1 was estimated to contribute about 9.4% (Fig. 5d) to the $PM_{2.5}$ concentration decrease,
2 compensating the large increase caused by meteorology, which is comparable with
3 previous studies of about 8.6% reduction in emissions (Liu et al., 2017; He et al., 2017a)
4 for the same two months. In other words, without the regional emission reduction
5 efforts, the observed $PM_{2.5}$ concentration in December 2015 would have had a similar
6 rate of 54.9% increase as the worsening meteorology conditions would bring about as
7 compared with December 2014. This assessment of emission reduction is supported
8 by the estimate of emission inventories for the BTH region in the Decembers of 2014
9 and 2015 by Zheng et al. (2019) who found out that the monthly emission strengths
10 for $PM_{2.5}$, SO_2 , NO_x , VOCs and NH_3 in 2015 were reduced by 22.0%, 6.9%, 2.5%, 2.5%
11 and 2.5%, respectively, as compared with 2014. The sensitivity and the nonlinear
12 response of $PM_{2.5}$ concentrations to the air pollutant emission reduction in the BTH
13 region (Zhao et al., 2017) have been estimated to be about 0.43 for both primary
14 inorganic and organic $PM_{2.5}$, 0.05 for SO_2 , -0.07 for NO_x , 0.15 for VOCs, 0.1 for NH_3 .
15 Combining the emission reduction percentages between Decembers 2014 and 2015
16 and the nonlinear response of emissions to the $PM_{2.5}$ concentrations results in an
17 approximately 10.2% ambient $PM_{2.5}$ concentration reduction due to the emission
18 changes. This is very close to the estimate of emission reduction contribution to the
19 December $PM_{2.5}$ concentration difference of about 9.4% between 2014 and 2015 by
20 the EMI framework.



1 **3.2. PM_{2.5} Trends and Meteorological Contributions**

2 The annual averaged PM_{2.5} concentrations in China have been decreased
3 significantly from 2013 to 2019. Figure 7 shows the observed spatial distribution of
4 national PM_{2.5} concentrations from 2103 to 2019, respectively. These spatial
5 distributions are consistent with those of primary and precursor emissions of PM_{2.5}
6 (Wang et al.), pointing out to the fundamental cause of the air pollution in China.
7 From the spatial distributions, it is clear that the regions of BTH, FWP, CEN and NWC
8 had the highest PM_{2.5} concentrations among the 9 regions. Even though the national
9 concentrations have been reduced significantly from 2013 by reducing emissions,
10 the pollution center of particular matters has not been changed very much, locating
11 at the southern Hebei Province and indicating the macroeconomic structure has not
12 been gone through a great change yet. Another phenomenon can be seen from the
13 distribution is that in the North-west China, especially in some cities of the Xinjiang
14 and Ningxia Provinces, the PM_{2.5} concentrations were on an increasing trend, due to
15 certain migrating industries from developed regions in East China.

16

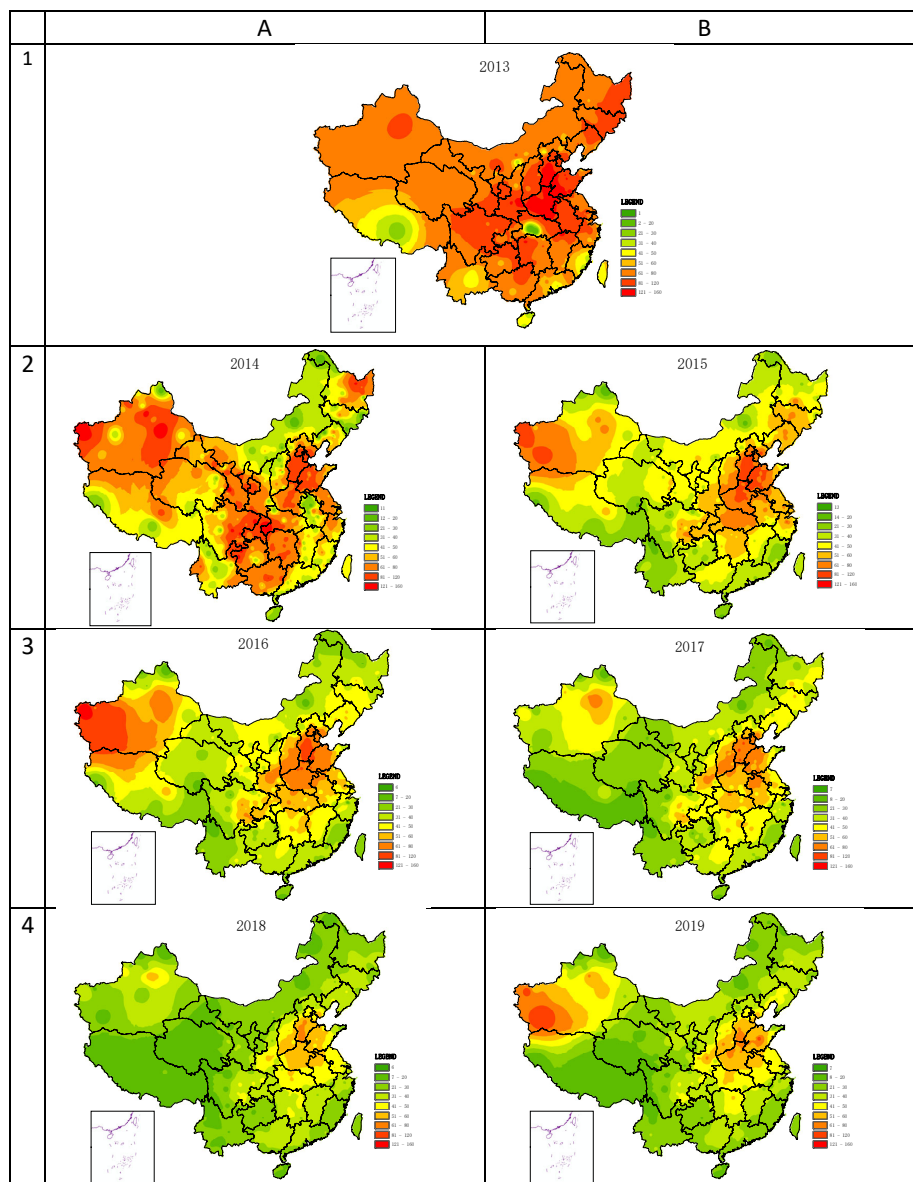
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1

2 Figure 7: Regional annual PM_{2.5} concentration distributions from 2013 to 2019.

3

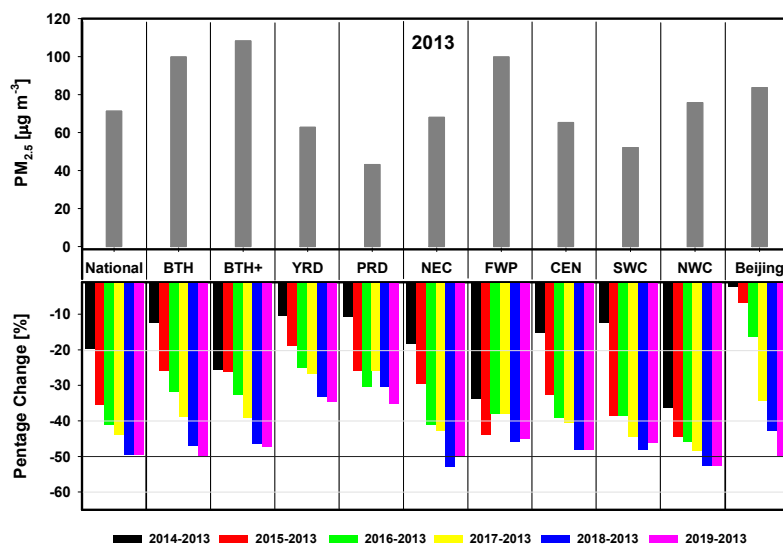
4 Averaged for the nation, 9 focused regions and Beijing, the PM_{2.5} trend lines

5 were shown in Figure 2. It is seen that all regions have had a large reduction of more

6 than 35% in surface PM_{2.5} concentrations in 2019 as compared with those in 2013.



1 The averaged national annual concentration at $36 \mu\text{g m}^{-3}$ has been very close to the
2 national standard of $35 \mu\text{g m}^{-3}$ while the concentrations in PRD, SWC and NEC
3 regions have been below the standard. Regions above the standard are BTH+, BTH,
4 YRD, CEN and FWP. Regionally, the largest drop percentage of $\text{PM}_{2.5}$ was seen in NEC
5 and NWC regions (Fig. 8), reaching over 50% compared with 2013. In the BTH, BTH+,
6 FWP and CEN regions, the reduction was in the range of 45% to 50% while in YRD
7 and PRD the reduction was around 35%.



8

9 Figure 8: Annual averaged $\text{PM}_{2.5}$ concentrations in 2013 (top) and corresponding changing
10 rates (bottom) from 2014 to 2019 as compared with 2013 for the nation, 9 regions and
11 Beijing City.

12

13 As one of the key factors in controlling the ambient $\text{PM}_{2.5}$ concentration
14 variations, the annual meteorological fluctuations, i.e. $\text{EMI}_{2.5}$, from 2014 to 2019
15 with 2013 as the base year, are shown in Figure 9 for nine regions. Generally, the



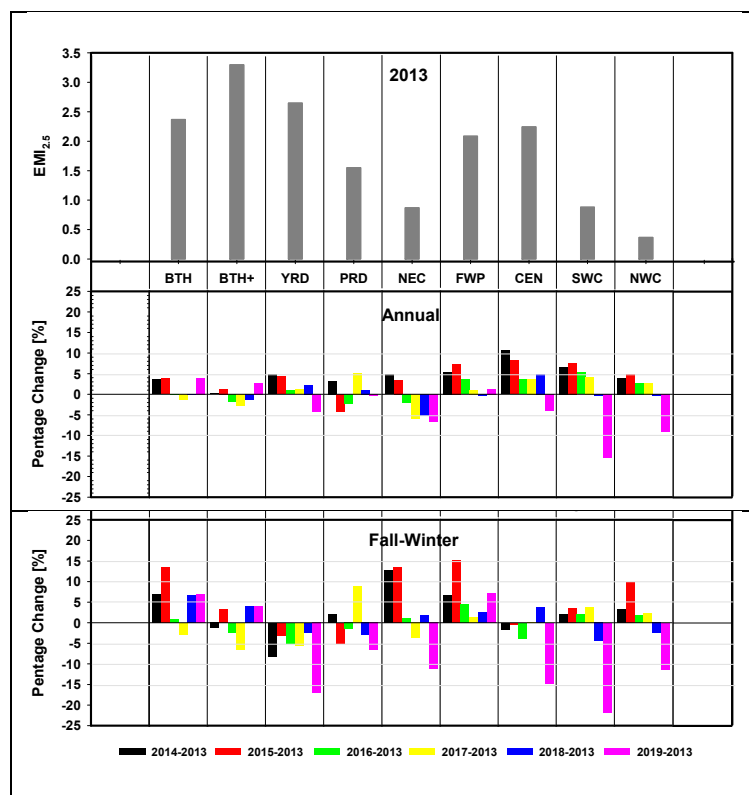
1 annual $EMI_{2.5}$ shows a positive or negative variation, reflecting the meteorological
2 features for that specific region. Except for a couple of regions or years, most of the
3 fluctuations are within 5% as compared with 2013 and have a no definite trend. It
4 can be inferred that the meteorological conditions are possibly responsible for about
5 5% of the annual $PM_{2.5}$ averaged concentration fluctuations from 2013 to 2019 (Fig.
6 9 middle). This is consistent with what has been assessed in Europe by Andersson et
7 al. (2007).

8 The variations in meteorological contributions ($EMI_{2.5}$) to $PM_{2.5}$ for the heavy
9 pollution seasons of fall and winter (October 1 to March 31) generally follow the
10 same fluctuating pattern as the annual average but are much larger than the average
11 (Fig. 9 bottom), over 5% for most of the regions and years. For specific regions and
12 years, e.g. BTH, YRD, NEC, SWC and CEN, the variations are between 10-20% as
13 compared with 2013. Since the $PM_{2.5}$ concentrations are much higher in the
14 pollution season, the larger meteorology variations in fall-winter would exercise
15 more controls to the heavy pollution episodes than the annual averaged
16 concentrations, signifying the importance of meteorology in regulating the winter
17 pollution situations.

18 It is found that though most of the regions have a fluctuating $EMI_{2.5}$ in the
19 pollution season during the 2014-2019 period (Fig. 9 bottom), the YRD and FWP
20 show a consistent favorite and un-favorite meteorological conditions, respectively.
21 BTH has witnessed the same un-favorite conditions as FWP except in 2017. In other
22 words, in BTH and FWP, the decrease in ambient concentrations of $PM_{2.5}$ from 2014



- 1 to 2019 has to overcome the difficulty of worsening meteorological conditions with
- 2 larger control efforts.



- 3
- 4 Figure 9: Annual averaged $EMI_{2.5}$ in 2013 (top) and corresponding changing rates for annual
- 5 average (middle) and for fall-winter seasons (bottom) from 2014 to 2019 as compared with
- 6 2013 in 9 regions.
- 7



1

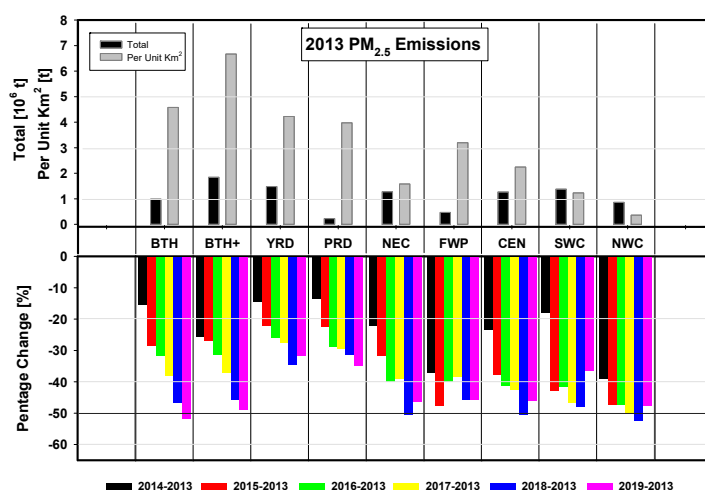
2 **3.3. Attribution of Control Measures to the PM_{2.5} Trend**

3 As it is well known that the final ambient concentrations of any pollutants are resulted
4 from the emission, meteorology and atmospheric physical and chemical processes.
5 Separating emissions and meteorology contributions to the pollution level reduction entails
6 a combined analysis of them. The analysis in Section 3.2 shows that from 2013 to 2019, the
7 national averaged PM_{2.5} as well as those for 9 separate regions were all showing a gradual
8 decline trend (Fig. 8). By 2019, 45% - 50% of reductions in surface PM_{2.5} concentrations
9 were achieved while the meteorology contributions did not show a definite trend as from
10 2013, clearly pointing out the contribution of emission reductions in the trend. Using the
11 analysis framework for separating emissions from meteorology based on the monitoring
12 data of PM_{2.5} and EMI_{2.5} (Section 2.4), the emission change contributions are estimated.

13 Figure 10 shows the 2013 base emissions of PM_{2.5} (Zhao et al., 2017) and the annual
14 changes in the emission contributions to the PM_{2.5} concentrations from 2014 to 2019 as
15 estimated from the EMI_{2.5} and observed PM_{2.5}. For the emissions, it is found that the unit
16 area emissions match better with ambient concentrations of PM_{2.5} in regions than the total
17 emissions and the high emission regions are BTH, BTH+, YRD, PRD and FWP in 2013.
18 Nationally by 2019, the emission reduction contributions to the ambient PM_{2.5} trend
19 accounted for ranging from 32% to 52% of the total PM_{2.5} decrease percentage, while in BTH
20 and BTH+ regions the reduction was more than 49% from 2013 base year emissions, leading
21 the national emission reduction campaign. The emission reduction rates clearly illustrate the
22 effectiveness of the national-wide emission control strategies implemented since 2013 and
23 the emission reduction is the dominate factor for ambient PM_{2.5} declining trend in China.



- 1 Taking the analysis data of $PM_{2.5}$ and $EMI_{2.5}$ from this study for BTH+ region from 2013 to
- 2 2017, it is found that control strategy contributed more than 90% to the $PM_{2.5}$ decline. Chen
- 3 et al (2019) has estimated that the control of anthropogenic emissions contributed to 80%
- 4 of the decrease in $PM_{2.5}$ concentrations in Beijing from 2013 to 2017.



- 5
- 6 Figure 10: Annual $PM_{2.5}$ emissions (total and per unit Km^2) for 2013 (top) and corresponding
- 7 changing rates (bottom) from 2014 to 2019 as compared with 2013 in 9 regions.
- 8
- 9 Regionally, the emission reduction trends from 2014 to 2019 display some unique
- 10 characteristics. For the regions of BTH, BTH+ and PRD, the year-by-year reduction rate is
- 11 consistent, indicating that regardless of fluctuations in meteorology, these regions have had
- 12 an effective emission control strategy and maintained the emission reduced year by year
- 13 since 2014. However, in some regions such as FWP, NEC, SWC and NWC, the emission
- 14 reduction rates were fluctuating from 2014 to 2019, implying the emissions in these regions
- 15 were increased in certain years. Especially in FWP from 2016 to 2017, the emissions were
- 16 estimated to be increased by about 10%, and then decreased in 2018 and 2019, despite of



1 the factor that FWP has experienced un-favorite meteorological conditions during this
2 period.

3 Table 2 summarizes the $PM_{2.5}$ difference between 2019 and 2015 and the relative
4 contributions of meteorology and emission changes to the difference for all China, Beijing
5 and nine regions. Once again, as of the end of 2019, the $PM_{2.5}$ concentrations are all
6 reduced from 2015, ranging from -1.8% in FWP to -46.2% in Beijing. During this period of
7 time, regions of BTH, BTH+, PRD and Beijing had encountered un-favorite meteorological
8 conditions with positive $EMI_{2.5}$ changes, which indicated that for these regions, emission
9 reductions were not only to maintain the decline trend but also to offset the un- favorite
10 meteorological conditions in order to achieve the observed reductions in ambient $PM_{2.5}$
11 concentrations. On the contrary, for the regions of FWP and SWC, the emission control
12 impacts were to deteriorate the concentrations, implying an increase in emissions to
13 restrain the $PM_{2.5}$ concentration decrease by favorite meteorological conditions. For other
14 regions, both meteorology and emission controls contributed to $PM_{2.5}$ decrease from 2105
15 to 2019, with the control measures contributing from -7.9% in NWC to -68.4% in NEC (Table
16 2).

17 Therefore, due to the diversity of meteorological conditions and emission distributions
18 in China, their impacts on ambient $PM_{2.5}$ concentrations display unique regional
19 characteristics. Overall, the emission controls are the dominant factor in contributing the
20 decline trend in China from 2013 to 2019. However, in certain regions or certain period of
21 years, emissions were found to increase and the meteorological dominance did occur,
22 which means the design of national control strategies has to take both meteorology and
23 emission impacts simultaneously in order to achieve maximum results.



- 1 Table 2: Observed PM_{2.5} difference between 2019 and 2015 as well as its attributions to
- 2 meteorology and control measures for all China, Beijing and nine regions.

Regions	Observed PM _{2.5} Difference		Attributions			
			Meteorology (EMI)		Emission Controls	
	(μg m ⁻³)	(%)	(μg m ⁻³)	Relative %	(μg m ⁻³)	Relative %
National	-10	-21.7	-4.1	-40.9	-5.9	-59.1
BTH	-24	-32.4	+0.1	+0.4	-24.1	-100.4
BTH+	-23	-28.8	+1.2	+5.4	-24.2	-105.4
YRD	-10	-19.6	-4.0	-39.7	-6.0	-60.3
PRD	-4	-12.5	+1.4	+36.0	-5.4	-136.0
NEC	-14	-29.2	-4.4	-31.6	-9.6	-68.4
FWP	-1	-1.8	-3.6	-362.2	+2.6	+262.2
CEN	-12	-23.1	-5.5	-45.5	-6.5	-54.5
SWC	-4	-12.5	-8.5	-211.5	+4.5	+111.5
NWC	-6	-14.3	-5.5	-92.1	-0.5	-7.9
Beijing	-36	-46.2	+3.4	+9.4	-39.4	-109.4

- 3 Note: "+" increased; "-" decreased

4 4. Conclusions

5 Based on a 3-D chemical transport model and its process analysis, an Environmental
 6 Meteorological Index (EMI_{2.5}) and an assessment framework have been developed and
 7 applied to the analysis of the PM_{2.5} trend in China from 2013 to 2019. Compared with
 8 observations, the EMI_{2.5} can realistically reflect the contribution of meteorological factors to
 9 the PM_{2.5} variations in the time series with impact mechanisms and can be used to as an
 10 index to judge the meteorological conditions whether are favorite or not to the PM_{2.5}
 11 pollutions in a region or time period. In conjunction to the observational trend data, the
 12 EMI_{2.5}-based framework has been used to quantitatively assess the separate contribution of
 13 meteorology and emission changes to the time series for 9 regions in China. Results show



1 that for the period of 2013 to 2019, the $PM_{2.5}$ concentrations have been dropped
2 continuously throughout China, by about 50% on national average. In the regions of NWC,
3 NEC, BTH, BEIJING, CEN, BTH+, SWC, the reduction was in the range of 46% to 53% while in
4 FWP, PRD and YRD, the reduction was from 45% to 35%. It is found that the control
5 measures of emission reduction are the dominant factors in the $PM_{2.5}$ declining trends in
6 various regions. By 2019, the emission reduction contributes about 47% of the $PM_{2.5}$
7 decrease from 2013 to 2019 on the national average, while in BTH region the emission
8 reduction contributes more than 50% and in YRD, PRD and SWC regions, the contributions
9 were between 32% and 37%. For most of the regions, the emission reduction trend was
10 consistent throughout the period except for FWP, NEC, SWC and NWC where the emission
11 amounts were increased for certain years. The contribution by the meteorology to the
12 surface $PM_{2.5}$ concentrations from 2013 to 2019 was not found to show a consistent trend,
13 fluctuating positively or negatively about 5% on annual average and 10-20% for the fall-
14 winter heavy pollution seasons.

15

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