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Enhanced summertime PM_{2.5}-suppression of O₃ formation over the Eastern U.S. following the O₃-sensitivity variations†

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The suppression of ozone (O₃) formation due to the presence of fine particulate matter (PM_{2.5}) has recently been highlighted for further O₃ pollution controls in regions that suffer high ozone concentrations. Here we derive multiple PM_{2.5}-suppression factors for the Eastern United States (U.S.) major cities based on a non-linear fitting of the PM_{2.5} and O₃ relationship from the multiyear surface observations. Our results show that these PM_{2.5}-suppression factors are increasing with time and generally follow the transition of the O₃-sensitive regime towards NO_x-limited chemistry. A spatial discrepancy of this suppression factor is seen currently with a higher value in the Southeastern U.S. than in the Northeastern U.S. A spatial similarity between urban regions and their downwind locations was observed for the New York City metro area. This more extensive formulation of the PM_{2.5}-suppression factor will further improve the ability of models to help guide O₃ and PM_{2.5} concentration pollution controls.

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Environmental significance

Ozone and fine particulate matter (PM_{2.5}) remain troublesome air pollution problems for a large number of areas, including metropolitan areas in the Northeastern U.S. like New York City. The paper's findings shed light on the interplay of these pollutants, namely the role that enhanced PM_{2.5} can play in suppressing ozone formation. The magnitude of ozone suppression from PM_{2.5} provides an additional indicator of the sensitivity of ozone formation from its VOC and NO_x precursors. The PM_{2.5} suppression of ozone formation per unit of PM_{2.5} mass concentration has been increasing over the period 2004–2019 in the Northeast U.S., indicating a transition from VOC-limited to NO_x-limited ozone formation sensitivity. It also provides guidance for further O₃–PM_{2.5} studies and pollution control regulations.

1. Introduction

The Eastern United States (U.S.) has been marked as a region persistently suffering from the co-occurrence of summertime ozone (O₃) and particulate matter (with a diameter under 2.5 μm, PM_{2.5}) pollution during the summertime.¹ However, these summertime concentrations of O₃ and PM_{2.5} in the region's major cities have shown decreasing trends since the 1970s as a result of the implementation of emission control policies.^{2–6} While the region has seen reductions in both pollutants, extreme concentrations of O₃ and PM_{2.5} (defined as the top 5%

of measured values in a given year) in New York City (NYC) have shown different overall reductions with a more significant reduction for PM_{2.5} than O₃. This can be attributed to the different reduction rates of their precursors, with the control policies targeting sulfur dioxide (SO₂) and primary PM_{2.5}, than volatile organic compounds (VOCs) and nitrogen oxides (NO_x).⁷

Co-occurrence of summertime maximum daily 8 h average O₃ (hereafter: MDA8 O₃) and the daily 24 h average PM_{2.5} (hereafter: DA24 PM_{2.5} for simplification) based on ground measurements in NYC has shown a direct relationship between the pollutants,⁷ with a monotonically increasing near-linear relationship for low PM concentrations. A leveling-off or even decreasing relationship for high PM concentrations was observed in megacity clusters in China.^{7–10} This flat or declining relationship has been partly attributed to the scavenging of hydroperoxyl (HO₂)/nitrate radicals (NO₃) by high concentrations of PM_{2.5} that inhibits the photochemical production of O₃,^{11–15} or reduced photolysis rates with PM_{2.5} increasing.¹⁶ A number of model simulations have used a uniform reactive uptake coefficient for HO₂ on aerosols (γ_{HO₂} = 0.2)^{9,17–21} to focus

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at SP3 ($\text{HCHO}/\text{NO}_2 \sim 2.7$). However, the variation of the HCHO/NO_2 ratio comparing SP2 to SP1 (SP2 vs. SP1: 2.2 vs. 1.5, 46% increasing compared to SP1) was larger than the variation of the $\text{PM}_{2.5}$ -suppression factor (both near 0.1, Fig. 1a and b), and this generally matches the current model simulation result about the O_x - NO_x relationship considering the $\text{PM}_{2.5}$ effect over Chinese urban regions from Li *et al.* (2022),³¹ which indicated that the $\text{PM}_{2.5}$ -suppression effect was weaker at higher NO_x concentrations in the VOC-limited regime, but strengthened as the O_3 -sensitivity approached the transitional regime. The weak $\text{PM}_{2.5}$ -suppression effect in the VOC-limited regime could be due to the competition for the consumption of HO_x by NO_x rather than $\text{PM}_{2.5}$, making the NO_x concentration the dominant factor for O_3 concentration sensitivity under these conditions.³¹

For the NYC downwind sites (Eisenhower Park site, Fig. 1c; Holtsville site, Fig. 1d, and Criscuolo Park site, Fig. 1e), their $\text{PM}_{2.5}$ -suppression factors for each subperiod were similar to the NYC urban sites (IS51 site, Fig. 1a and Queens College site,

Fig. 1b). This can be attributed to urban plume transport, which has been discussed in some detail from recent and current studies based on the 2018 Long Island Sound Tropospheric Ozone Study (LISTOS),^{22–24} and the formed O_3 and $\text{PM}_{2.5}$ and some unreacted precursors can be carried to downwind regions. Meanwhile, based on the fact that the averaging time period for MDA8 O_3 (8 hours) and DA24 $\text{PM}_{2.5}$ (24 hours) is generally much longer than the time scale of the photochemical reactions, it is reasonable to believe that the urban plume transport could result in the similar $\text{PM}_{2.5}$ -suppression factors for both urban and downwind sites. However, except for the Eisenhower Park site which had an HCHO/NO_2 ratio for each subperiod similar to the Queens College site as their proximity to each other (<20 km), the HCHO/NO_2 ratios of each subperiod for the Holtsville site (around 70 km from Queens College site) and the Criscuolo Park site (around 100 km from the Queens College site) were much higher than the values of the NYC urban sites (Fig. 1f). Based on satellite observations, this indicates that Holtsville

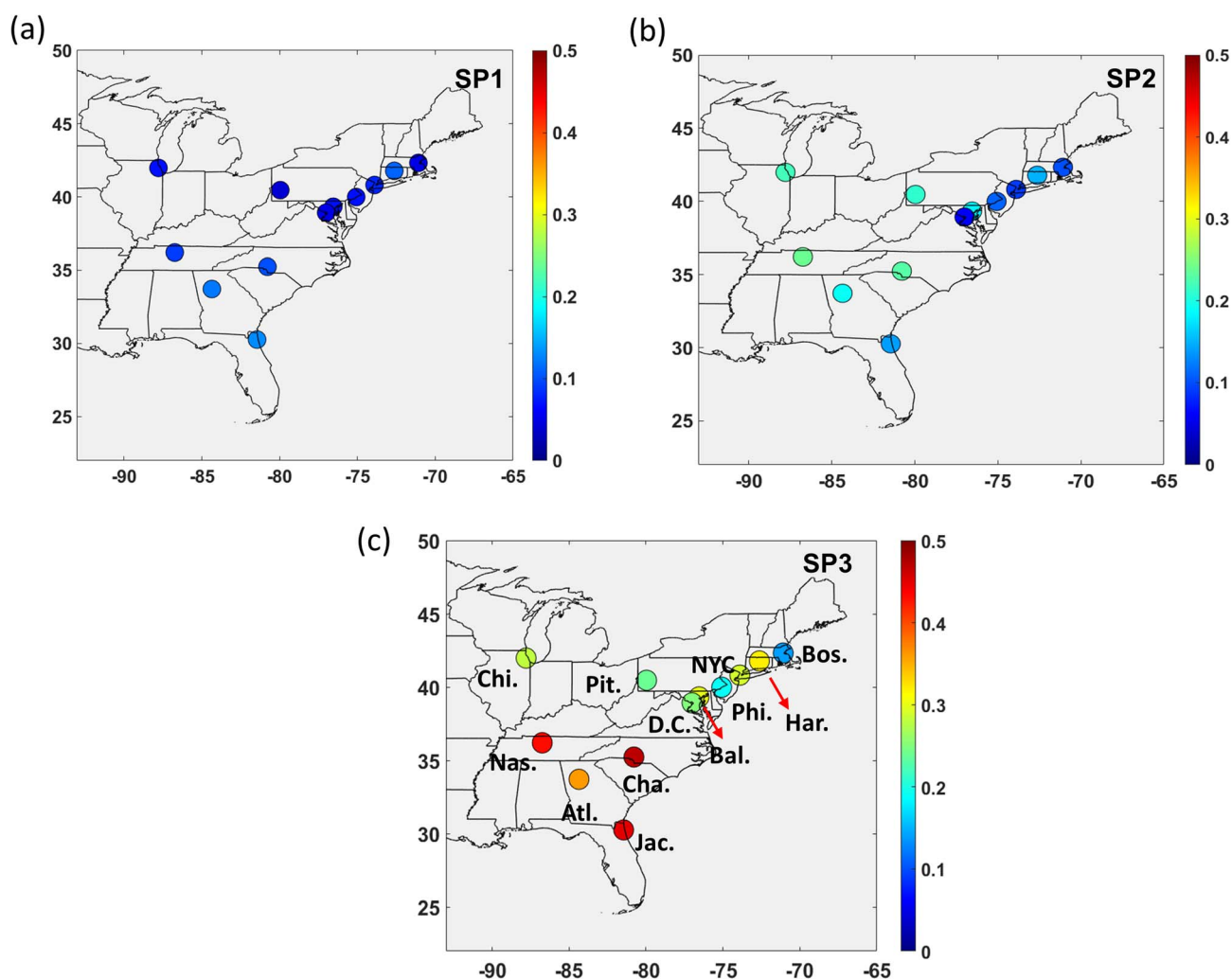


Fig. 2 The $\text{PM}_{2.5}$ -suppression factor distribution map for the twelve major cities in the Eastern U.S. for (a) Subperiod 1 (SP1: 2004–2008), (b) Subperiod 2 (SP2: 2009–2013), and (c) Subperiod 3 (SP3: 2014–2019). These twelve major cities include Chicago (Chi., Illinois), Pittsburgh (Pit., Pennsylvania), Boston (Bos., Massachusetts), Hartford (Har., Connecticut), NYC (New York), Philadelphia (Phi., Pennsylvania), Baltimore (Bal., Maryland), Washington, D.C. (District of Columbia), Charlotte (Cha., North Carolina), Atlanta (Atl., Georgia), Jacksonville (Jac., Florida), and Nashville (Nas., Tennessee).



