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Ozone formation potential related to the release of volatile organic compounds (VOCs) and nitrogen oxide (NO_x) from a typical industrial park in the Pearl River Delta†

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Ozone (O₃) pollution has been recognized as the major air pollution in the Pearl River Delta (PRD) region, South China. Understanding O₃ formation sensitive to volatile organic compound (VOC)- and nitrogen oxide (NO_x)-limited regimes is a key step for alleviating O₃ pollution. Herein, measurements of VOCs, NO_x and O₃ were simultaneously performed at multi sampling sites in an industrial park of the PRD region during June, 2020. VOCs/NO_x ratios ranged from 0.5 to 5.7, suggesting that the O₃ formation was in the VOC-limited regime in the industrial park. The estimated O₃ formation potential (OFP) of VOCs showed that alkenes and aromatic hydrocarbons from motor vehicles and industrial sources contributed to 40% and 39% of the O₃ formation, respectively, in the industrial park. However, a low O₃ level (<50 ppb) was observed in the region where high OFP values (>194 ppb) were estimated. Further analysis found that the concentration of NO_x (25 ± 10 ppb) in the high O₃ region was lower than that (36 ± 6 ppb) in the low O₃ region, mostly due to the titration reaction of NO and O₃ to form NO₂, therefore leading to the consumption of O₃. This result implies that NO_x control was not conducive to the O₃ pollution in the study region. Thus, O₃ pollution control in the study region should be taken into consideration in terms of the effect of NO_x titration and control of VOC emissions.

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Environmental significance

O₃ pollution is a major air pollution in the Pearl River Delta region, China. Understanding O₃ formation sensitive to VOC- and NO_x-limited regimes is a key step for alleviating O₃ pollution. Herein, VOCs, NO_x and O₃ were simultaneously measured in a typical industrial park in this region. The ratio of VOCs to NO_x indicates that O₃ formation was in the VOC-limited regime in the industrial park. Alkenes and AHs principally from motor vehicles and industrial sources were the main potential contributors to the formation of O₃. However, no increase in O₃ was found in the abundant alkene and AH regions, mainly due to the titration reaction of NO and O₃ to form NO₂. These findings highlighted the role of NO_x-titration in O₃ formation under the VOC-limited regime.

1 Introduction

Ozone (O₃) has an inverse impact on the ecosystem and human health.^{1,2} High levels of O₃ can lead to a reduction in crop yields and abnormal growth of plants,^{3–5} and O₃ pollution can damage the health system of humans, leading to complications such as ocular discomfort, interruption of the skin's normal barrier

function and development and progression of obstructive airway diseases.^{6–8} More importantly, O₃ also occurs as the by-product in the complicated photochemical reaction involving NO_x and volatile organic compounds (VOCs).^{9,10} The area and frequency of O₃ pollution in China showed an increasing trend in recent years.¹¹ The mean O₃ concentration increased from 84.3 to 95.8 μg m⁻³ from 2015 to 2018 in China,¹² especially in the Beijing–Tianjin–Hebei, Fen Wei River Plain, and Pearl River Delta (PRD) regions. The O₃ formation is highly dependent on its precursors including VOCs and NO_x.^{9,10} However, non-linear relationships existed between the formation of O₃ and concentrations of VOCs and NO_x. Therefore, it is necessary to figure out the interaction between the production of O₃ and its precursors.

The formation of O₃ involves a series of complex photochemical reactions of NO_x and VOCs.¹³ The types and levels of VOCs and NO_x, their ratio as well as the intensity of solar radiation will influence the formation of the ground-level

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O₃.^{14,15} The O₃ formation was sensitive to its precursors in three regimes including the VOC-limited regime, the NO_x-limited regime and their transition regime.¹⁶ Therefore, the effectiveness of NO_x and VOC emission control in reducing O₃ needs to be discerned so that the O₃ pollution problem can be alleviated.

VOCs and NO_x can be from natural and anthropogenic sources. The biogenic emission is an important natural source of VOCs,¹⁷ while organic matter decomposition and lightning would generate natural NO_x.¹⁸ However, large amounts of VOCs and NO_x produced by human activities have become the main source recently.¹⁹ Industrial sources and vehicle exhaust are the important contributors to VOCs and NO_x emissions.^{20,21}

The PRD region is one of the O₃ pollution regions in China.²² In this region, the average O₃ concentration increased from 48 to 60 μg m⁻³ from 2006 to 2019, with an average growth rate of 0.80 μg (m³ a)⁻¹.²³ The formation of O₃ at urban sites was mainly in the VOC-limited region.²⁴ Moreover, industrial emission contributes to more than 40% of VOCs in this region.^{25,26} To characterize VOC_s from industrial emission, the measurement of VOCs in the industrial park *via* the fixed sampling site was reported previously.^{27,28} Compared to a single sampling site, observation of VOCs in industrial parks at multiple sampling sites would recognize spatial distribution characteristics and sources of VOCs.²⁹ Moreover, observation of VOCs at multiple sampling sites can be conducted to understand the dispersion of VOCs in an industrial park.

In this study, the observation of VOCs at 24 sampling sites was performed in an industrial park in PRD. Additionally, O₃ and NO_x were simultaneously monitored using a multipollutant sensor package. The objectives of this study were to (1) characterize the spatial distribution of VOCs, NO_x and O₃ in an industrial park, and (2) compare the spatial distribution of O₃ formation potential and observed O₃, and evaluate how the VOCs or NO_x are sensitive to O₃ formation.

2 Experimental section

2.1. Sampling sites

24 sampling sites are located in an industrial park in the PRD region, South China (Fig. S1†), which were decided by the principle of grid monitoring and practical situations, and the detailed description is listed in Table S1.† The industrial park consisted of more than 20 enterprises including raw material manufacturing, machinery manufacturing, chemical process, comprehensive utilization of waste resources, printing, textiles, food processing, pharmaceutical manufacturing, transportation, electronic and machine manufacturing, and research and experimental development (Fig. S1†). In addition, this industrial park is close to a main traffic road, and the traffic source therefore would be another contributor to the local air pollution. It should be noted that the industrial park is partly surrounded by a mountain, with a fully covering dense forest. Some natural VOCs (*e.g.*, isoprene) from the forest might release into the atmosphere in this industrial park. In addition, a village with approximately 15 000 residents is located in the central position of the industrial park.

2.2. Measurement of VOCs

A total of 24 sites with an interval of approximately 450 m were sampled in the industrial park area of 2.5 km × 1.6 km (Fig. S2†). The sampling period was from 10:40 to 15:00 on June 17, 2020, which the photochemical reaction was active due to enough sunlight. Wind speed was about 1 m s⁻¹ and the day was sunny during the sample period. Such a low wind was unfavourable for the dispersion of VOCs from the industrial park emission. Additionally, the industrial park is partly surrounded by a mountain. The presence of the mountain in the industrial park coupled with the low wind speeds would further enhance VOC accumulation from the industrial park emissions. VOCs were sampled using 2.7 L vacuum stainless Summa canisters (ENTECH Instruments Inc., Silonite™, CA, USA) with a restrictor valve (1 L min⁻¹) above ground level of 1.5 m. Prior to sampling, each Summa canister should be filled with high purity nitrogen (99.999%), vacated and washed repeatedly for more than five times. One out of every 20 cleaned canisters needs to be filled with high purity nitrogen and placed for more than 24 h. The sample was analysed to ensure that the target compounds in the canisters could not be detected or fell below the detection limit. The sampled VOCs were analysed as soon as possible within 20 days. The collected VOC samples were detected using an Entech 7100 pre-concentrator (Entech Instruments Inc., CA, USA) and a gas chromatography-mass spectrometer (7890A GC-5975C MS, Agilent Technologies, USA) based on the USEPA TO-15 method. The detailed procedure, and qualitative and quantitative analysis methods of VOCs are provided in the ESI (Text S1†).

2.3. Measurements of O₃ and NO_x

O₃ and NO_x were monitored using a multipollutant sensor package.³⁰ O₃ (OX-B431), NO (NO-B4) and NO₂ (NO₂-B42) sensors were commercially obtained from Alphasense Ltd (Essex, UK). All the gas sensors were calibrated completely with standard gases.³⁰ NO standard gas was from Shenkai Gases Tech. Ltd (Shanghai, China), O₃ standard gas was generated *via* an O₃ generator (Ozone Primary Standard, Model 49i-PS, Thermo Fisher Scientific Inc., USA) and NO₂ was produced from the reaction between NO and O₃. After calibration, the sensors were compared with field observation instruments including a UV photometric O₃ analyser (Model49C, Thermo Electron Corporation, USA) and a NO-NO₂-NO_x monitor (Model 42i, Thermo Scientific, USA), which were applied to a monitoring test together. The detailed calibration processes were described previously.³⁰

2.4. Ozone formation potential (OFP)

OFP was used to evaluate the contribution of VOCs to O₃ formation in the atmospheric photochemical reaction.²² Taking into account the reactivity of VOCs and their concentrations in the atmosphere, the Maximum Incremental Activity (MIR) method was used to calculate the contribution of VOCs to O₃ generation in near-ground level.³¹ The OFP can be expressed as:

$$\text{OFP}(i) = \text{Conc}(i) \times \text{MIR coefficient}(i)$$



where $\text{Conc}(i)$ is the concentration of compound i in the atmosphere (ppb: parts per billion by volume). MIR coefficient(i) is the MIR of compound i , which is updated previously.³² The values of MIR are summarized in Table S2.† A total of 70 VOC species contributed to the OFP in this study.

2.5. Observation-based chemical box model

An observation-based model (OBM) combined with the master chemical mechanism simulation online (AtChem Online, <https://www.atchem.york.ac.uk/>, Version 2.0) was used to explore the contribution of VOCs or NO_x to O_3 formation. Averaged air pollutants (O_3 , NO , NO_2 , and VOCs) and meteorological parameters (temperature, pressure, relative humidity) and boundary were input into the OBM-AtChem Online as constraints. The model was run at 00:00 local time as the initial time. Relative incremental reactivity (RIR) was an indicator to diagnose the sensitivity of O_3 reduction to its precursors NO_x or O_3 . The RIR can be expressed as:

$$\text{RIR} = \frac{\Delta P(\text{O}_3)/P(\text{O}_3)}{\Delta X/X}$$

Here, $\Delta P(\text{O}_3)/P(\text{O}_3)$ represents differences in the O_3 production rate (Text S2†), which was calculated in the OBM-AtChem Online. $\Delta X/X$ refers to differences in precursor concentrations, and the value was 20%.

3 Results and discussion

3.1. Characteristics of VOCs

The spatial distribution of the concentrations of total VOCs (TVOCs) at 24 sampling sites in the industrial park is shown in Fig. 1a. TVOCs ranged from 13 to 120 ppb with an average level of 41 ± 24 ppb (mean \pm standard deviation) in the industrial

park (Fig. S3†). High values (>54 ppb) can be found in the northwestern, southwestern and southeastern areas, while low values (<33 ppb) were observed in the central region. Low TVOCs in the central region may be mostly attributed to no industrialized source of VOCs (a village situated there). Table S3† lists the TVOCs, species number and top 10 species in the industrial park, urban, suburban and rural regions of the PRD region compared with that in this study. The mean TVOCs in this study were similar to the levels (35–89 ppb) in the urban area,^{33–35} but generally higher than that in suburban (8–60 ppb) and rural (13–23 ppb) areas in the PRD region.^{33–37} It should be noted that the sampling was performed during the COVID-19 pandemic, which negatively affected the industrial activity. This might lead to the reduction in VOCs emission from the industrial park. These indicate that industrial emissions have a great impact on the regional atmospheric environment. In addition, complex emission sources would lead to an increase in the VOC levels and the levels of VOCs in urban areas with industrial parks were generally higher.

Based on the functional groups, the detected VOCs were classified as five categories namely alkanes (19 species), alkenes (8 species), aromatic hydrocarbons (AHs) (18 species), halogenated hydrocarbons (HHs) (33 species) and oxygenated VOCs (OVOCs) (3 species) (Table S2†). The concentrations of alkanes, alkenes, AHs, HHs and OVOCs were 14 ± 16 , 6 ± 4 , 10 ± 5 , 9 ± 7 and 2 ± 2 ppb, respectively. Alkanes (35%) were the most contributing species, followed by AHs (24%), HHs (21%) and alkenes (14%), and the minimum contribution was from OVOCs (6%) (Fig. 2a).

Among these alkanes (Fig. 2b), undecane (27%), isopentane (23%), *n*-hexane (12%), 3-methylpentane (9%), and decane (4%)

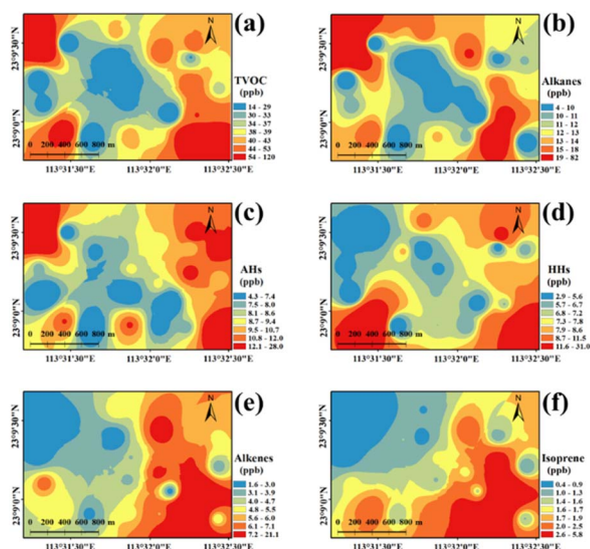


Fig. 1 The distribution of VOCs and air pollutants. The colours from blue to red represent a gradual increase in concentration. (a) The distribution of TVOCs; (b) the distribution of alkanes; (c) the distribution of AHs; (d) the distribution of HHs; (e) the distribution of alkenes; (f) the distribution of isoprene.

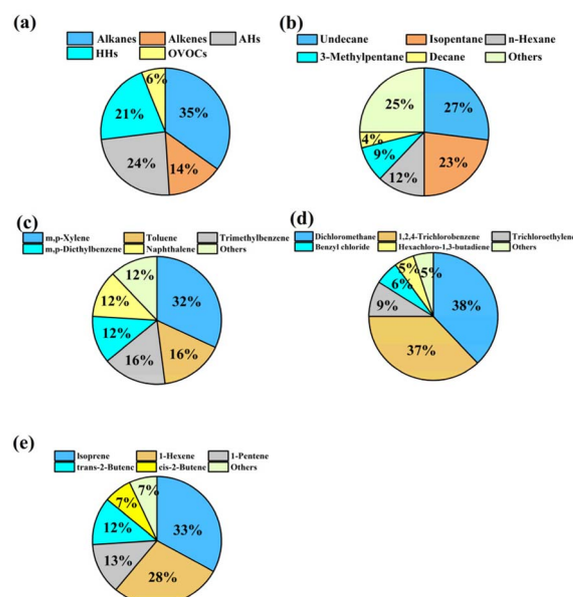


Fig. 2 The percentage of species and the contribution of dominant compounds: (a) the proportion of VOC species; (b) the ratio of the dominant species of alkanes; (c) the ratio of the dominant species of AHs; (d) the ratio of the dominant species of HHs; (e) the ratio of the dominant species of alkenes.



were the dominant species and their concentrations varied from 13 to 92 ppb. Furthermore, the spatial distribution of alkanes was illustrated (Fig. 1b). A high value (83 ppb) of alkanes was found in the northwestern region of the industrial park. It is worth noting that a highway toll station is located nearby, which might be the reason for the high concentration of alkanes in this area, since these alkanes (*e.g.*, undecane and isopentane) were frequently detected in the vehicle exhaust, tunnels or roadways.^{35,36} In addition, relatively high values (22–35 ppb) of alkanes were also found in the southeastern area, which had transportation, pharmaceutical manufacturing and machinery manufacturing, producing abundant alkanes.³⁷

Among AHs (Fig. 2c), toluene and *m*, *p*-xylene had a similar level of 36 ppb and they totally represented 32% of AHs. Naphthalene, *m*, *p*-diethylbenzene, and trimethylbenzene were the second most AHs, each of which accounted for 12% of AHs. The spatial distribution of AHs is shown in Fig. 1c, which shows a high value (27 ppb) of AHs in the northwestern region of the industrial park, and they mainly consisted of *m*, *p*-diethylbenzene and 1,2,3-trimethylbenzene. These AHs in the northwestern region of the industrial park might be released from vehicle exhaust³⁸ and the pharmaceutical industry.³⁹ In addition, a relatively high value (13–17 ppb) of AHs was observed in the eastern region, and toluene and *m*, *p*-xylene were primarily detected AH species. According to previous studies, they were common compounds from vehicle emission³⁸ and evaporative loss of solvents during various industrial processes.⁴⁰ In this study, in the eastern region of the industrial park, there are approximately five logistics companies with dense traffic, which were the main source of AHs.

Among HHs (Fig. 2d), dichloromethane (38%), 1,2,4-trichlorobenzene (9%), trichloroethylene (6%), (chloromethyl)benzene (5%) and hexachloro-1,3-butadiene (5%) were the dominant HH species (Fig. 2d). High levels of HHs were distributed in the southeastern (31 ppb) and southwestern (31 ppb) regions of the industrial park (Fig. 1d). These regions were mostly affected by multiple pollution sources. HHs are mostly used as industrial solvents in certain sections of the pharmaceutical chemical industry³⁹ and chemical industry⁴¹ (*e.g.*, coatings, fragrances and flavors, printing, textiles). Further, the contributions of solvent use, refrigeration and chemical manufacturing to HHs were remarkable in the PRD region.⁴² Additionally, dichloromethane is mainly consumed in the pharmaceutical industry, metal cleaning agents and chemical production.⁴³ In the southwest and southeast regions of the study area, printing, textiles, raw material manufacturing, pharmaceutical manufacturing industries located here might be the main sources of HHs.

Among alkenes (Fig. 2e), the five dominant species were isoprene (33%), 1-hexene (28%), 1-pentene (13%), *trans*-2-butene (13%) and *cis*-2-butene (7%). The high-value (8–21 ppb) region of alkenes was distributed in the eastern region (Fig. 1e). Abundant isoprene was measured in the eastern region of the industrial park (Fig. 1f), which was adjacent to a mountain covered with plentiful trees. Therefore, isoprene in the eastern region was likely due to the vegetation emission in the mountain.

Comparatively, the measured VOCs were dominated by alkanes (42%), followed by aromatics (20%), alkenes (18%), and

halo-hydrocarbons (13%) in a typical petrochemical area in the Yangtze River Delta, China.⁴⁴ The most dominant VOCs were alkanes (45%), followed by alkenes (25%) and AHs (22%) in a complicated industrial area of Nanjing, China.³⁴ That is, the contribution ratio of these species in this study varied from other industrial parks, but alkanes and AHs were mainly from industrial emission.

3.2. Characteristics of NO_x and O₃

NO_x and O₃ were simultaneously monitored at sampling sites in the industrial park. As shown in Fig. 3 and S4†, the average concentration of NO_x was 30 ± 10 ppb, ranging from 7 to 47 ppb. High levels (33–47 ppb) of NO_x appeared in the western region, which was close to a roadway. However, a different distribution between NO and NO₂ was found in the western region. A higher value of NO was measured in the northwestern region (19–21 ppb) compared to the southeastern region (5–8 ppb). The spatial distribution of NO₂ showed an opposite feature. Such an opposing feature of NO and NO₂ can be attributed to the efficient conversion NO to NO₂, as discussed below.

For the O₃, it can be found that the concentration of O₃ ranged from 36 to 85 ppb, with a mean value of 55 ± 16 ppb (Fig. S4†). Interestingly, high values of O₃ were found in the eastern and northern regions of the industrial park (Fig. 3d), where the level of NO₂ showed low values. In contrast, a low level of O₃ and a high concentration of NO₂ were observed in the southwestern or southeast region of the industrial park. This difference in spatial distribution between O₃, NO, and NO₂ suggests that the formation of O₃ in these regions was strongly related to the variation in the levels of NO and NO₂. However, relatively low values of O₃ and high values of VOCs were distributed in northeastern or southeastern regions in the industrial park, as previously discussed.

3.3. Formation of O₃ related to VOCs and NO_x

The contribution of precursors to the formation of O₃ can be described as from VOC-limited, NO_x-limited or their transition regimes. A ratio of VOCs to NO_x has been widely employed to

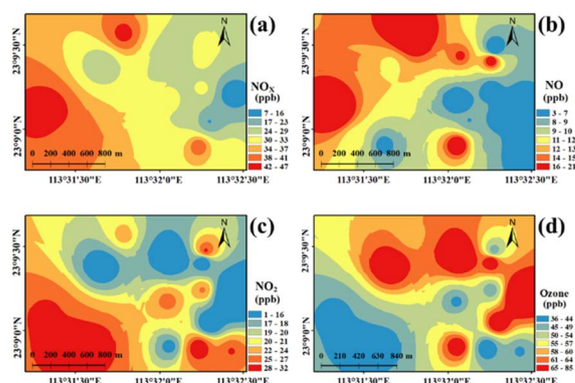


Fig. 3 The distributions of NO_x and ozone. The colours from blue to red represent a gradual increase in concentration: (a) the distribution of NO_x; (b) the distribution of NO; (c) the distribution of NO₂; (d) the distribution of ozone.



evaluate the sensitivity of O₃ formation to the VOC-limited or NO_x-limited regime in various studies.^{45–47} In this study, the VOC/NO_x ratio varied from 0.5 to 5.7, with a mean value of 1.8. These VOC/NO_x ratios were much lower than the 8 : 1 threshold for a transition from the VOC-limited to the NO_x-limited regime,^{48,49} suggesting that the formation of O₃ in the industrial park was probably in the VOC-limited regime. It must be pointed out that these results exhibited here were on the basis of the threshold value of the VOC/NO_x ratio reported for Los Angeles.⁵⁰ Based on the observation-based chemical box model, the RIR values of NO_x and VOCs were −0.45 and 0.21, respectively, suggesting that the reduction of O₃ in the industrial park was VOC-sensitive. The RIR values of alkanes, alkenes (not including isoprene), isoprene, aromatic hydrocarbons, halogenated hydrocarbons and OVOCs were 0.01, 0.07, 0.06, 0.05, 0.01 and 0.01, respectively.

To further evaluate the formation of O₃ sensitive to VOCs in the industrial park, high (above 50 ppb) and low (below 50 ppb) concentrations of O₃ with change in concentration and species of VOCs were discussed. The concentration of TVOCs in the high O₃ region (35 ± 12 ppb) was comparable to that in the low O₃ region (48 ± 21 ppb) (Fig. 4a). Similar values of alkanes, alkenes, AHs and OVOCs in the industrial park were also found in the high and low O₃ regions (Fig. S5†). However, a lower level of HHs in the high O₃ region (14 ± 11 ppb) was measured compared to that in the low O₃ region (7 ± 2 ppb). Thus, the concentration of the generated O₃ is closely related to the concentration and species of VOCs.

Due to the difference in the reactivity of individual VOC species to generate O₃, the OFP of individual VOC species was also used to evaluate the contribution of VOCs to the generation of O₃ in the high and low O₃ regions in this study. As previously discussed, the increase in the reactive VOCs would result in the formation of O₃ due to the VOC-limited regime.^{51,52} Fig. 5 shows the OFP values of reactive VOCs and their proportion. The maximum value of OFP for VOCs was contributed by alkenes (40%), followed by AHs (39%), alkanes (12%) and OVOCs (7%), and the minor contribution was from HHs (2%) (Fig. 5a). More specifically, isoprene, *trans*-2-butene and 1-hexene contributed to 36%, 20% and 16% of OFP, respectively, in the alkenes. *Cis*-2-butene and 1-pentene showed a similar proportion of 10% in

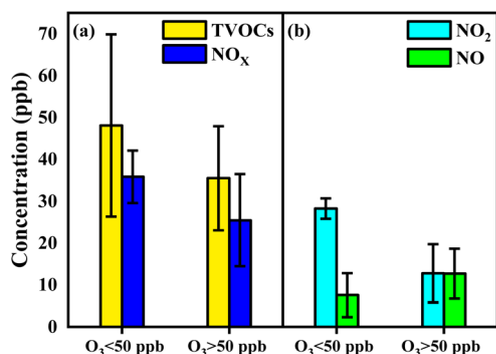


Fig. 4 (a) The concentration of TVOCs in the high O₃ region and the low O₃ region; (b) the concentration of NO_x in the high O₃ region and the low O₃ region.

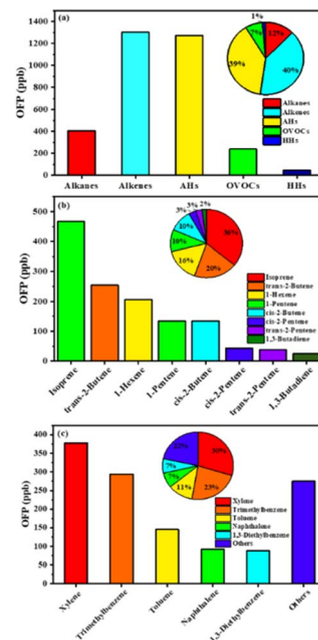


Fig. 5 The OFP values of reactive VOCs and their proportion. (a) VOC species; (b) the dominant species of alkenes; (c) the dominant species of AHs.

the alkenes (Fig. 5b). Xylene and trimethylbenzene accounted for 30% and 23% of OFP in the AHs, respectively. Approximately 12% of OFP in the AHs was attributed to toluene (Fig. 5c).

Further, the distribution of OFP values in the investigated industrial park was demonstrated. There was no increase in OFP values in the high O₃ region (132 ± 52 ppb) compared with the low O₃ region (153 ± 58 ppb) (Fig. S6†). Moreover, the spatial distribution of OFP showed that a high value of OFP appeared in the southeastern region of the industrial park (Fig. S7†), where existed a low value of O₃. These results imply that the enrichment of O₃ has not occurred in the high OFP value of VOCs.

The concentration of NO_x in the high O₃ region was 25 ± 10 ppb, which was lower than 36 ± 6 ppb in the low O₃ region, as shown in Fig. 4b. This indicates that the formation of O₃ in the industrial park readily occurred in the low NO_x region. In the meantime, a lower level of NO₂ (13 ± 7 ppb) was observed in the high O₃ region relative to that (28 ± 2 ppb) in the low O₃ region (Fig. 4b). In contrast, a higher level of NO (13 ± 5 ppb) in the high O₃ region was compared with that (8 ± 5 ppb) in the low O₃ region. This diversity suggests that O₃ is enriched in the low NO₂ region or high NO region in the industrial park.

The low O₃ region in the industrial park was mostly due to the titration reaction of NO and O₃ to form NO₂, in turn, showing high values of NO₂ in the low O₃ region.^{53,54} This reaction also leads to the consumption of NO, therefore reducing NO in the low O₃ region. The reduction of O₃ in the high OFP value region in the industrial park can be attributed to the presence of the titration reaction, in spite of the production of O₃ in the VOC-limited regime. In contrast, abundant NO and the decrease in NO₂ in the high O₃ region relative to the low O₃



region suggest that the effect of the titration reaction on the consumption of O₃ was minor, therefore leading to the accumulation of O₃.

The mean concentration of O₃ was 55 ± 16 ppb in this study, which was higher than the averaged O₃ concentration of 35–50 ppb from 57 national monitoring sites in the PRD region.⁵⁵ Generally, higher averaged O₃ levels (about 20–60 ppb) in the suburban or rural sites were compared with those in the urban sites (about 10–45 ppb) in the PRD region.⁵⁶ The NO titration effect likely leads to the decrease in O₃ levels in urban sites,⁵⁵ which agreed with the finding in the current study. However, higher levels of O₃ in the industrial park relative to urban sites in the PRD region might be also related to meteorological conditions. Low wind and sunny day during the sampling period would also accumulate precursors and thus lead to O₃ formation in the small industrial park.

It was found that the VOC-limited region was mostly located at urban cores and large city clusters in the PRD region through the analysis of the whole PRD.^{57,58} Lu *et al.* mentioned that the O₃ formation was in a VOC-limited regime in Guangzhou and Hong Kong, South China.⁵⁹ Several studies also showed that the formation of O₃ in urban sites of Guangzhou was mainly limited by VOCs.^{60–62} Oppositely, the generation of O₃ in the NO_x-limited or transition regime was observed in the suburban or rural sites of the PRD region.^{58,63} For example, Wang *et al.* indicated that the formation of O₃ was frequent in the NO_x-limited regime at a rural site in the PRD.⁵⁸ Wang *et al.* found differently that a rural site in Hong Kong was a transition regime.⁶⁴ Li *et al.* pointed out that the O₃ formation regime would change over complicated spatial and diurnal variations.⁶⁵

In the present work, the O₃ formation was VOC-limited in the industrial park, which is in agreement with the findings at urban sites in the PRD region reported previously.^{60,63} However, due to the variation in the reactivity or sources of VOCs among different urban or industrial sites, changes in VOCs would alter the contribution of VOCs to the production of O₃ at different sites. Louie *et al.* observed three sites in the PRD from 2008 to 2009, and found that AHs and alkenes together contributed to about 80% of the total OFP of VOCs, while alkanes contributed to 12%–14%.⁶⁶ Ou *et al.* also found that AHs and alkenes become predominant with contributions of 59% and 26%, respectively, of the total OFP.⁶⁷ Similar to our research, we found that AHs and alkenes made up 79% of the total OFP. Vehicle exhaust and industrial processes (*e.g.* pharmaceutical industry, raw material manufacturing, electronic and machine manufacturing) were the factors that mainly contributed to the OFP in PRD, making up more than 40% of the total OFP.⁶⁸ It is worth noting that even though the industrial park was in a VOC-limited regime, due to the different sources and compositions of VOCs, the targets for ozone reduction and control of VOCs are also different. In this park, vehicle exhaust and industrial processes mentioned above were the important control sources.

3.4. Atmospheric implication

Multi-site observation presented here found that abundant NO_x (33–47 ppb) and TVOCs (71–120 ppb) in the western region of

the industrial park were related to vehicle emissions. The enrichment of TVOCs (65–76 ppb) in the eastern region of the industrial park was likely attributed to the industrial emissions. Multi-site observation can provide a clearly spatial distribution of atmospheric pollution in a study area, which was a supplement to the measurement at an individual site. Our results also showed that the reduction in O₃ in the high OFP region was due to the consumption of O₃ *via* the titration reaction. Therefore, the multi-site observation will be a useful method to better understand the source and evolution of atmospheric pollution, especially in the small scale, such as an industrial park.

The O₃ abatement should be achieved by the reduction of its precursors. Our results suggest that O₃ formation in the industrial park was in the VOC-limited regime due to the ratios of VOCs/NO_x ranging from 0.5 to 5.7. Xylene (12%), trimethylbenzene (9%), and *trans*-2-butene (8%) emitted from vehicular or industrial emissions in the industrial park have been considered as the significant contributors to O₃ formation. Thus, reduction of these VOC emissions in the industrial park might be helpful for alleviating O₃ pollution.

Although it showed high OFP values for the O₃ formation, the stronger titration reaction led to a significant decrease in O₃. The NO_x concentration changed from 25 ± 10 to 36 ± 6 ppb, corresponding to the high O₃ region and low O₃ region in the industrial park, respectively. Several studies also observed that reducing a small amount of NO_x in the PRD region would lead to an increase in O₃ concentration in the VOC-limited regime, and it was called as the NO_x titration trap.^{65,69} Such a NO_x emission reduction was counterproductive for reducing VOC-limited O₃ concentrations only if the reduction of NO_x was large enough or the emission of VOC was eliminated altogether.⁷⁰ Tang *et al.* found that Beijing successfully bypassed the NO_x titration trap *via* elimination of scattered emission sources as well as the implementation of the China National VI (A) standard for motor vehicles after 2016.⁷¹ Therefore, these control strategies might not only result in the decrease of VOC emission, but also reduction of NO_x emission in the PRD region, while, bypassing the NO_x titration trap in the future.

4 Conclusions

The simultaneous measurements of VOCs, NO_x and O₃ were conducted at multi-sample sites in an industrial park of the PRD region. The ratio of VOCs to NO_x ranging from 0.5 to 5.7 indicates that the formation of O₃ was in the VOC-limited regime in the industrial park. Alkenes and AHs principally from motor vehicles and industrial sources have been considered as the main potential contributors to the formation of O₃ in the industrial park. However, no increase in O₃ was found in the abundant alkene and AH regions, which showed high NO₂ or low NO values in the industrial park. This was mostly due to the titration reaction of NO and O₃ to form NO₂, therefore leading to the consumption of O₃. These findings highlighted the role of NO_x-titration in the formation of O₃ under the VOC-limited regime. Thus, the policy for O₃ abatement in the study region might consider the control of VOC emission as well as NO_x-titration.



Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

Taicheng An: conceptualization, writing–review & editing, supervision, funding acquisition. Jiajia Li: investigation, methodology, data curation. Qin hao Lin: methodology, investigation. Guiying Li: writing–review & editing, data curation.

Conflicts of interest

There are no conflicts to declare.

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