Process analysis and sensitivity study of regional ozone formation over the Pearl River Delta, China, during the PRIDE-PRD2004 campaign using the Community Multiscale Air Quality modeling system

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Abstract. In this study, the Community Multiscale Air Quality (CMAQ) modeling system is used to simulate the ozone (O₃) episodes during the Program of Regional Integrated Experiments of Air Quality over the Pearl River Delta, China, in October 2004 (PRIDE-PRD2004). The simulation suggests that O₃ pollution is a regional phenomenon in the Pearl River Delta (PRD). Elevated O₃ levels often occurred in the southwestern inland PRD, Pearl River estuary (PRE), and southern coastal areas during the 1-month field campaign. Three evolution patterns of simulated surface O₃ are summarized based on different near-ground flow conditions. More than 75% of days featured interactions between weak synoptic forcing and local sea-land circulation. Integrated process rate (IPR) analysis shows that photochemical production is a dominant contributor to O₃ enhancement from 09:00 to 15:00 local standard time in the atmospheric boundary layer over most areas with elevated O₃ occurrence in the mid-afternoon. The simulated O₃ production efficiency is 2–8 O₃ molecules per NOₓ molecule oxidized in areas with high O₃ chemical production. Precursors of O₃ originating from different source regions in the central PRD are mixed during the course of transport to downwind rural areas during nighttime and early morning, where they then contribute to the daytime O₃ photochemical production. The sea-land circulation plays an important role on the regional O₃ formation and distribution over PRD. Sensitivity studies suggest that O₃ formation is volatile-organic-compound-limited in the central inland PRD, PRE, and surrounding coastal areas with less chemical aging (NOₓ/NOᵧ > 0.6), but is NOₓ-limited in the rural southwestern PRD with aged air (NOₓ/NOᵧ < 0.3).

1 Introduction

Tropospheric ozone (O₃) is a secondary pollutant produced through a series of photochemical reactions involving mainly nitrogen oxides (NOₓ) and volatile organic compounds (VOCs) in the presence of sunlight. Elevated concentration of ground-level O₃ is of great environmental concern due to its adverse impacts on human health and ecosystems, as well as its greenhouse effect (NARSTO, 2000). The O₃ pollution in megacities and regional areas has been investigated recently by many studies (Evyugina et al., 2006; Gonçalves et al., 2009; Lei et al., 2007; Kimura et al., 2008; Yu et al., 2009; Chang, 2008; Ran et al., 2009; Wang et al., 2009), in which the topics addressed include O₃ photochemical production, chemical sensitivity to precursors, roles of
meteorological conditions, O3 source attributions and development of control strategy.

The Pearl River Delta (PRD) is one of the most urbanized and industrialized regions in southern China. This area contains three megacities (Guangzhou, Shenzhen, and Hong Kong) and numerous medium and small cities, houses 4% of the total population of China, and produces about 19% of China's gross domestic product each year. As a consequence of the substantial economic development and increases in air pollutant emissions, in recent years, the PRD has experienced rapid deterioration of air quality. Photochemical smog has been one of the most severe air quality issues in the PRD, where surface O3 levels exceeding the national hourly standard of 200 µg/m³ (∼0.093 ppm) are frequently observed by air quality monitoring networks, especially in fall when northerly winds and clear sky conditions prevail (Wang et al., 2001, 2003; Zhang et al., 2007, 2008a).

The O3 pollution in Hong Kong has raised much attention over the past decade. Meteorological conditions have been found to be closely associated with O3 pollution in Hong Kong and the Pearl River estuary (PRE) area. The synoptic systems related to O3 episodes in Hong Kong have been classified into three patterns: tropical cyclones, continental anticyclones, and low-pressure troughs (Huang et al., 2005, 2006; Chan and Chan, 2000). The impacts of the complex topography, local circulations, and low mixing height on heavy O3 pollution were investigated (Liu and Chan, 2002; Ding et al., 2004; Lam et al., 2005), and a conceptual model was developed to explain the effect of land-sea breezes on pollutant transport, trapping, and accumulation (Lo et al., 2006). In addition, the local versus regional contributions to O3 episodes in Hong Kong have been widely investigated using chemical transport models (CTMs) and a backward trajectory method (Ding et al., 2004; Lam et al., 2005; Huang et al., 2005, 2006; Wang et al., 2006; Zhang et al., 2007). The relative importance of local production and of regional transport from inland parts of the PRD was found to change under different meteorological conditions and at different locations. Previous studies have also addressed the relationship between O3 and its precursors. In Hong Kong, the photochemical formation of O3 is generally believed to be VOC-limited (Chan and Yao, 2008).

However, O3 pollution in the PRD region outside of Hong Kong and PRE has not yet been investigated adequately, and only limited studies have been reported. Wang et al. (2005) evaluated the importance of different emission sources on the concentrations of O3 and other pollutants for the whole PRD in March 2001. Wei et al. (2007) investigated the impact of biogenic VOC emissions on O3 formation. Zhang et al. (2008b) and Shao et al. (2009) adopted an observation-based model to investigate O3 production sensitivity to precursors in Guangzhou. CTMs are useful tools for exploring the spatiotemporal evolution of O3 pollution and assessing the roles of different atmospheric processes in O3 formation. In spite of the works by Wang et al. (2005) and Wei et al. (2007), there are still very limited three-dimensional (3-D) modeling studies to address these issues over the inland PRD region. Moreover, to develop effective strategies for O3 pollution control, our understanding of O3-VOC-NOx chemistry over the whole PRD needs to be improved through more observational and modeling studies.

The Program of Regional Integrated Experiments of Air Quality over the Pearl River Delta, China, in October 2004 (PRIDE-PRD2004) obtained a comprehensive database of O3, particulate matter, and other air pollutant measurements in this region (Zhang et al., 2008a), providing an ideal testbed for CTM application to thoroughly investigate the spatiotemporal evolution and chemical characteristics of ground-level O3 pollution over the whole PRD region.

In this paper, the US Environmental Protection Agency (EPA) Community Multiscale Air Quality (CMAQ) modeling system is applied to simulate O3 episodes during the PRIDE-PRD2004 campaign. Our goal is to quantify the impacts of different chemical and physical processes on elevated O3 formation and to characterize the regional O3 chemical production over the entire PRD area. Section 2 describes the modeling methodology. Section 3 presents and discusses the model performance, the spatiotemporal evolution of elevated O3, the influence of different processes on O3 formation, and the characteristics of O3 chemical production. A summary of conclusions is provided in Sect. 4.

2 Methodology

2.1 Model setup and inputs

We use CMAQ (version 4.5, Byun and Schere, 2006) with the Statewide Air Pollution Research Center version 99 (SAPRC-99) chemical mechanism (Carter, 2000) to simulate O3 formation during the whole month of October 2004. The model is configured to have triple-nested domains (Fig. 1). The outer domain with a horizontal grid spacing of 36 km covers the entire area of China, the 12-km grid-spacing inner domain covers Guangdong, Hong Kong, and Macao, and the innermost domain with a 4-km horizontal grid resolution focuses on the PRD. All the grids have 13 layers vertically extending from the surface to an altitude of ∼17 km above the ground, with seven layers below 1 km and the first layer thickness of ∼19 m. The outputs from the outer domains are used to provide boundary conditions for the inner ones by one-way nesting. A spin-up period of 3 days (1–3 October) is used to minimize the influence of initial conditions.

The fifth-generation Pennsylvania State University/National Center for Atmospheric Research (PSU/NCAR) Mesoscale Model (MM5, version 3.7) (Grell et al., 1994) is used to simulate the meteorological fields to drive CMAQ. MM5 is configured with the same nested horizontal domains as those for CMAQ (except that at least three grid cells are extended from each side of the
CMAQ domain) and with 34 vertical layers. The one-way nesting simulations are performed with the following physics options: the mixed-phase microphysics, the Grell cumulus scheme, the medium-range forecast model’s boundary layer scheme, the rapid radiative transfer model longwave scheme, and the Noah land-surface model. The National Centers for Environmental Protection (NCEP) \(1^\circ \times 1^\circ\) global reanalysis data, the NCEP global surface and upper air observation data, and observations measured in the atmospheric boundary layer (ABL) during the PRIDE-PRD2004 campaign are used to prepare the initial and boundary conditions for the MM5 simulations. The four-dimensional data assimilation technique (“grid nudging”) is used to nudge 3-D winds, temperature and humidity at six-hour intervals, and surface winds at three-hour intervals.

The gridded and speculated hourly emission inputs for CMAQ are prepared using the Sparse Matrix Operator Kernel Emissions (SMOKE) model (version 2.3, Houyoux et al., 2000). The TRACE-P anthropogenic emissions inventory (Streets et al., 2003) with \(1^\circ \times 1^\circ\) resolution is used for the outer 36-km domain. Inventory inputs (to SMOKE) of anthropogenic emissions for the inner domains are based on an inventory compiled by the Hong Kong Environmental Protection Department (P. Louie, personal communication, 2006), which has been applied in modeling studies such as Huang et al. (2005, 2006). Emission estimations in PRD region from several recent studies, including biomass burning emissions by Cao et al. (2005), vehicular emissions by Song and Xie (2006) and anthropogenic VOCs emissions by Liu et al. (2008a) are also included in the update of the inventory. In addition, the emissions are projected from the base year of original datasets to the year 2004 to include the impacts from the changes of economy, population and vehicle volume as well as other activity parameters relating to emissions. In this composite and high-resolution regional inventory, anthropogenic emissions of NO\(_x\), VOCs, CO, SO\(_2\), NH\(_3\), and PM are reported in three major source categories: point sources (mainly power generation and industrial sources), mobile sources (on-road and off-road vehicles, marine traffic, and aircraft), and area sources (e.g., domestic and commercial fuel combustion, industrial processes, solvent evaporation loss, storage and transport of petroleum products, and agricultural activities). The biogenic VOC and NO emissions are estimated by applying the Biogenic Emissions Inventory System version 3.09 (BEIS 3.09, Vukovich and Pierce, 2002) with a Chinese plantation survey dataset (Pearl River Delta environmental protection planning committee, 2006). The default BEIS emission factors and the hourly meteorological fields from MM5 simulations are used.

As shown in Table 1, mobile sources and power generation point sources contribute about 47% and 39%, respectively, of the total NO\(_x\) emissions in the PRD, whereas mobile sources, evaporation losses of solvents and petroleum, and biogenic sources are the three largest contributors to VOC emissions, accounting for 38%, 24% and 23%, respectively. Spatially, both NO\(_x\) and VOC emissions are found to concentrate over the inland urban areas of Guangzhou, Foshan, and Dongguan; the coastal areas of Dongguan and Shenzhen; and the urban core of Hong Kong (Fig. 2).

2.2 Model evaluation protocol

Predicted meteorological fields (including winds, temperature, and humidity) are examined against the hourly observations obtained during October 2004 for each domain. Performance statistics for the MM5 simulations are calculated using the Metstat statistical analysis package (Emery et al., 2001).

The simulation of \(O_3\) formation during 4–31 October is evaluated against observations made at two super sites of the PRIDE-PRD2004 campaign (Zhang et al., 2008a) and measurements collected simultaneously at the ten regular surface sites of the PRD air quality monitoring network (see site locations in Fig. 1b). Note that the platform of the super site at Guangdong Provinical Environmental Monitoring Center (GDEMC) sits on the roof of a 17-story building, \(\sim 50\) m above the ground, in the urban center of Guangzhou City. The other super site of Xinken is a seaside rural site. The levels of \(O_3\) and NO\(_x\) were measured by TECO commercial instruments TECO 49C and 42C, respectively, at the super sites and stations of PRD air quality monitoring network (Zhang et al., 2008b). Two kinds of measurement techniques for non-methane hydrocarbons (NMHCs) were adopted at the super sites. One was canister sampling followed by analysis.
using gas chromatography (GC) combined with mass spectrometry (MS) and flame ionization detection (FID) for C2-C12 species (Liu et al., 2008b), and the other was on-line GC-FID for C3-C12 species (Wang et al., 2008a). The canister measurements usually collected three samples per day at GDEMC and two samples per day at Xinken during the entire campaign, whereas the on-line observations are only available in the second half of the month. The simulated concentrations of all NMHC-related species in the SAPRC99 mechanism are summed together for comparison with canister NMHCs observations, and C2 species are not included in the simulated values when those values are compared with the on-line NMHCs observations.

The performance is judged by statistical measures, including the correlation coefficient, normalized mean bias (NMB), and normalized mean error (NME), defined as

$$NMB = \frac{\sum_{i=1}^{N} (C_i^s - C_i^o)}{\sum_{i=1}^{N} C_i^o} \cdot 100\%$$

$$NME = \frac{\sum_{i=1}^{N} |C_i^s - C_i^o|}{\sum_{i=1}^{N} C_i^o} \cdot 100\%$$

where $C_i^s$ and $C_i^o$ represent simulated and observed concentrations at a same monitoring site for the same hour, respectively, and $N$ is the total number of such data pairs of interest.

### 2.3 Integrated process rate analysis

Production of O$_3$ is the net result of interactions of the various atmospheric processes involved (e.g., chemistry, transport, deposition). The integrated process rate (IPR) analysis implemented in the CMAQ model is used here to investigate the influences of major physical processes and the net effect of chemistry on model predictions. The IPR analysis calculates hourly contributions of gas-phase chemistry, horizontal transport (including advection and diffusion), vertical transport (including advection and diffusion), dry deposition, and some other processes (such as cloud processes, aerosol processes, and emissions) at each model grid cell. Details of IPR analysis can be found elsewhere (Jeffries and Tonnesen, 1994; Jang et al., 1995; Gipson, 1999). Recent applications of the IPR in CMAQ model have been reported by Xu et al. (2008), Yu et al. (2009), Gonçalves et al. (2009), Wang et al. (2009), and Zhang et al. (2009a, b).

For the IPR analysis, we first assess the roles of various atmospheric processes in O$_3$ formation in ABL at the two super sites, GDEMC and Xinken, and at the Donghu station. The O$_3$ pollution at GDEMC represents a typical situation in Guangzhou urban areas (source region), whereas that at Xinken, a rural site with less local emissions, reflects the influences of pollutants transported from upwind areas (Guangzhou and Dongguan under northerly winds; Shenzhen and Hong Kong under southerly winds). Donghu is an urban site in Jiangmen City, located ~60 km downwind of the urban areas of Guangzhou and Foshan under northerly wind conditions. The maximum O$_3$ levels were recorded at Donghu during the campaign. We then investigate the influences of different processes (precursor emissions, physical transport, and gas-phase chemistry) on the formation and evolution of regional O$_3$ pollution over the PRD.

### 2.4 Ozone production efficiency calculation

Ozone production efficiency (OPE) is defined as the number of molecules of O$_3$ formed per NO$_x$ removed from atmospheric ozone-forming oxidation cycles [i.e., $P(O_3)/P(NO_x)$] (Seinfeld and Pandis, 2006). OPE is an important measure for determining the efficiency of the catalyst NO$_x$ in O$_3$ formation and for indicating the O$_3$-VOC-NO$_x$ sensitivity under certain polluted conditions (Sillman, 1995; Sillman and He, 2002). In this work, OPE for the time range and spatial region of interest is estimated based on the IPR results, which provide the net chemical production of O$_3$ and NO$_2$ at each grid cell on an hourly basis [i.e., $P(O_3)$ and $P(NO_2)$].

### Table 1. Summary of annual emissions of NO$_x$ and VOCs by source category over the PRD in 2004 (kilotons/yr).

<table>
<thead>
<tr>
<th>Source category</th>
<th>NO$_x$</th>
<th>VOCs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Power generation point source</td>
<td>294</td>
<td>34</td>
</tr>
<tr>
<td>Mobile source</td>
<td>349</td>
<td>545</td>
</tr>
<tr>
<td>Industrial source</td>
<td>52</td>
<td>20</td>
</tr>
<tr>
<td>Domestic &amp; commercial fuel combustion source</td>
<td>36</td>
<td>73</td>
</tr>
<tr>
<td>Solvent &amp; petroleum evaporation source</td>
<td>0</td>
<td>345</td>
</tr>
<tr>
<td>Agriculture source &amp; others</td>
<td>11</td>
<td>105</td>
</tr>
<tr>
<td>Biogenic source*</td>
<td>6</td>
<td>328</td>
</tr>
<tr>
<td>Total</td>
<td>748</td>
<td>1450</td>
</tr>
</tbody>
</table>

* The annual biogenic emissions are estimated based on the mean emission amounts in the October, 2004.

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**Fig. 2.** NO$_x$ (a) and VOC (b) emissions in the 4-km grid at 12:00 local standard time (LST) on 16 October 2004.
2.5 

O_3 sensitivity testing

Understanding of the non-linear relationship between O_3 formation and its precursors is critical for the development of an effective O_3 control strategy. We evaluate the O_3 response to precursors by perturbing the domain-wide anthropogenic emissions of NO_x and VOCs and then examining the resulting changes in O_3 concentrations and O_3 production rates. The perturbations of emissions include a 25% reduction in NO_x emissions, a 25% reduction in VOC emissions, and a 25% reduction in both emissions. The reductions are only applied to the anthropogenic emission sources, no changes to biogenic emissions. Also, the reductions are linearly distributed across all NO_x and/or VOC species and applied over all the time and space, at the same percentage of 25%. O_3 sensitivity to precursors may vary with the magnitude of emission reductions; here, the 25% emission-reduction scenarios are used because this reduction level is more feasible and probable in real controls than 50% or higher reductions. Furthermore, we examine the influence of the photochemical age of air mass (indicated by the ratio of NO_x/NO_y) on the sensitivity of O_3 formation to precursors.

3 Results and discussions

3.1 Evaluation of model performance

Simulated surface meteorological fields were examined against the surface hourly observations during the field campaign, and the MM5 model performance (shown in Table 2) is well within the typical range of meteorological modeling studies (Emery et al., 2001; Hanna and Yang, 2001).

Simulated O_3 concentrations compare well against the observations at the two super sites and at other network sites (Table 3), with a correlation coefficient of 0.73, NMB of −5.4%, and NME of 37.1%, comparable to the performance of other CMAQ applications (Zhang et al., 2006; Gonçalves et al., 2009). The simulations reproduce the diurnal variations and magnitude of O_3 reasonably well at most sites (Fig. 3), except at Tianhu, which is an upwind rural site. Comparisons of precursor concentrations (i.e., NO_2 and NMHCs) at the monitoring sites further demonstrate that the O_3 formation is captured reasonably well (and for the right reasons) over the domain and throughout the period (Table 3, Figs. 4 and 5).

Note that the levels of peak O_3 tend to be underpredicted at a few sites (e.g., Tianhu, Wanqingsha, and Zimaling, see Fig. 3). This may be related to several factors, such as the uncertainties in the precursor emissions and meteorological parameters. In the current inventory, compared with the data for PRD urban areas, the emission estimations over the suburban and rural areas of the PRD and the areas outside the PRD have higher uncertainties due to the limited source information available and fewer reported studies. For instance, the evolution of observed O_3 at Tianhu, a rural site in the far north of Guangzhou, had an elevated background level of O_3 ranging from 60 to 70 ppb during the campaign and was less affected by local photochemical production (except during 29–31 October). Considering the prevailing northerly winds in October, the underestimated O_3 at Tianhu may be caused by the uncertainties in emissions in northern rural areas of Guangzhou and distant northern areas such as Qingyuan. The overestimation of surface wind speeds by MM5 (see Table 2) may result in more transport and less accumulation of O_3 and its precursors, which is a probable cause of the underestimated O_3 peaks at some other sites as well.

The measurements of NO_x and VOCs are greatly influenced by local emissions. October is harvest season for agricultural crops in PRD and biomass burning in open fields was significant and was observed during the campaign. Although the emissions by burning crop residues have been included in the inventory, an exact estimation of precursor emissions is difficult due to the limited information on the details of burning events (i.e., the place and duration of biomass burning, amount of burned crop residues). This may result in the uncertainties in the simulated precursors concentrations and O_3 levels as well, especially at rural sites (e.g., Tianhu,

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Fig. 3. Time series of simulated surface O_3 against that observed at PRD monitoring sites during 4–31 October 2004.
Wanqingsha and Xinken). Another significant uncertainty in estimating VOCs emissions comes from the lack of local representative emission factors of VOCs from industrial and domestic solvent use, which also accounts for the underprediction of ambient VOCs concentrations, especially for aromatic levels. Further nighttime vertical diffusions are not easily simulated by current mesoscale meteorological models, especially over the complex topography and land use in PRD. This may be a reason for higher deviations of precursor levels between model results and measurements during night and early morning.

In addition, studies have revealed that molybdenum converter (equipped in the TECO 42C chemiluminescence NO\(_x\) analyzers as used in the campaign) reduces not only NO\(_2\) but also other nitrogen-containing compounds (e.g., nitric acid, peroxyacetyl nitrate and other organic nitrates) to NO, hence can cause the NO\(_x\) analyzer overestimating the real NO\(_2\) concentrations (Steinbacher et al., 2007). Apparently this overestimation of real NO\(_2\) concentrations by measurements accounts for, at least in part, the low bias of the NO\(_2\) performance (Table 3).

The modeled and observed NO are reasonably correlated with a correlation coefficient of 0.54, similar to that of NO\(_2\) (see Table 3). Note that most of the NO low biases were for the measured NO peaks during the night time and the early morning rush hour (not shown). This may indicate that the NO\(_x\) emissions were under-estimated at those time periods. However, the low biases can also be explained by the mismatching of modeled volume concentrations in a 4-km grid cell with observations which were largely impacted by local primary emissions under stable boundary layer conditions.

### 3.2 Regional spatiotemporal evolution of surface ozone

Both the CMAQ simulation and air quality measurements demonstrate that the PRD experienced serious photochemical smog pollution at the regional scale during the campaign (Fig. 3). The simulated monthly average distribution

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**Table 2.** Performance statistics of the MM5-simulated meteorological variables against surface observations during 1–31 October 2004.

<table>
<thead>
<tr>
<th>Meteorological variable</th>
<th>Statistical parameter</th>
<th>D1(*)</th>
<th>D2(**)</th>
<th>D3(***)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface wind speed</td>
<td>RMSE** (m/s)</td>
<td>2.07</td>
<td>2.06</td>
<td>1.64</td>
</tr>
<tr>
<td></td>
<td>Bias (m/s)</td>
<td>0.32</td>
<td>0.89</td>
<td>0.27</td>
</tr>
<tr>
<td></td>
<td>IOA***</td>
<td>0.79</td>
<td>0.65</td>
<td>0.66</td>
</tr>
<tr>
<td>Surface wind direction</td>
<td>Gross Error (deg.)</td>
<td>39.49</td>
<td>42.15</td>
<td>38.11</td>
</tr>
<tr>
<td></td>
<td>Bias (deg.)</td>
<td>3.47</td>
<td>2.47</td>
<td>0.56</td>
</tr>
<tr>
<td>Surface temperature</td>
<td>Gross Error (K)</td>
<td>2.27</td>
<td>2.09</td>
<td>1.98</td>
</tr>
<tr>
<td></td>
<td>Bias (K)</td>
<td>−0.90</td>
<td>−0.88</td>
<td>0.39</td>
</tr>
<tr>
<td></td>
<td>IOA</td>
<td>0.97</td>
<td>0.90</td>
<td>0.85</td>
</tr>
<tr>
<td>Surface humidity</td>
<td>Gross Error (g/kg)</td>
<td>1.13</td>
<td>1.64</td>
<td>1.62</td>
</tr>
<tr>
<td></td>
<td>Bias (g/kg)</td>
<td>−0.50</td>
<td>−1.30</td>
<td>−1.24</td>
</tr>
<tr>
<td></td>
<td>IOA</td>
<td>0.94</td>
<td>0.78</td>
<td>0.64</td>
</tr>
</tbody>
</table>

\* D1, D2 and D3 stand for the domains with 36 km, 12 km and 4 km horizontal resolution, respectively.

** RMSE stands for the root mean square error.

*** IOA stands for the index of agreement (Emery et al., 2001).

**Table 3.** CMAQ performance statistics for the simulated hourly concentrations of surface O\(_3\), NO, NO\(_2\) and NHMC against observations over the PRD during 4–31 October 2004.

<table>
<thead>
<tr>
<th>Species</th>
<th>Number of data pairs</th>
<th>Correlation coefficient</th>
<th>NMB(\ast) (%)</th>
<th>NME(\ast) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>O(_3)</td>
<td>6856</td>
<td>0.73</td>
<td>−5.4</td>
<td>37.1</td>
</tr>
<tr>
<td>NO</td>
<td>4591</td>
<td>0.54</td>
<td>−69.6</td>
<td>86.0</td>
</tr>
<tr>
<td>NO(_2)</td>
<td>4706</td>
<td>0.52</td>
<td>−26.8</td>
<td>52.1</td>
</tr>
<tr>
<td>NHMC</td>
<td>759</td>
<td>0.66</td>
<td>−24.2</td>
<td>47.7</td>
</tr>
</tbody>
</table>

\* NMB stands for normalized mean bias; NME stands for normalized mean error.
indicates that elevated levels of surface O$_3$ usually occurred in the western and southern PRD. As shown in Fig. 6, areas with O$_3$ higher than 0.090 ppm include southern Foshan, Jiangmen, Zhongshan, the PRE and surrounding areas, and the southwestern coastal area. Regional O$_3$ pollution was also recorded at stations located in the western PRD and around the PRE, such as Donghu, Zimaling, Wanqingsha, Xinken, and Tung Chung, where the O$_3$ nonattainment (i.e. hourly O$_3$ concentration exceeding the national ozone air quality standard of 200 $\mu$g/m$^3$ or $\sim$0.093 ppm) days accounted for 88%, 95%, 77%, 88%, and 61% of valid observation days during the campaign, respectively. The maximum hourly O$_3$ concentration of 0.179 ppm was measured at Donghu station (Zhang et al., 2008b). Although no surface observations of O$_3$ concentrations were conducted in central and western Jiangmen during the campaign, aircraft measurements in the ABL reported average O$_3$ levels of around 0.080 ppm and a maximum O$_3$ value of 0.101 ppm over Kaiping in the afternoon (close to the levels in Zhongshan, but higher than those in Foshan and Huizhou) (Wang et al., 2008b), suggesting the occurrence of O$_3$ pollution in the rural western PRD where the local precursor emissions are low.

The formation and distribution of O$_3$ pollution over the PRD is greatly influenced by synoptic weather conditions and local circulations. During the campaign, a high-pressure system dominated over the PRD and resulted in light-to-moderate northerly or northeasterly synoptic winds. Three cold air masses from the north intruded into the PRD on 1, 17, and 25 October. After the influence of cold air weakened, both meteorological observations and simulated results showed the evolution of sea-land breezes in the southern PRD. Under weak synoptic conditions, northeasterly/northerly/easterly winds were dominant over the PRD from early morning to midday, and southerly or southeastern sea breezes usually began in afternoon and ended around midnight in the southern PRD (Fan et al., 2008).

Based on the influences of the different near-ground flow patterns, we summarize three evolution patterns of simulated surface O$_3$ over the PRD during the studied month. In the first pattern, named as “O$_3$-South”, the flows in lower layers were characterized by northerly or northeasterly winds prevailing over the PRD during the whole day due to moderate or strong synoptic forcing resulting from the intrusion of northern cold air. Elevated O$_3$ during mid-afternoons was usually distributed over southern or southwestern inland and coastal regions, as well as the PRE area. At dusk, O$_3$ concentrations in the urban areas of Guangzhou, Foshan, Shenzhen, and Hong Kong dropped to low levels due to the decrease in photochemical production and the removal of O$_3$ by NO titration and dry deposition, while continuous transport by northerly winds moved the O$_3$ plume to the southern water area (e.g. Fig. 7a).

The other two patterns occurred when synoptic winds were weak and sea–land breezes showed significant influence on the transport of air pollutants. Weak synoptic winds and stagnant conditions usually resulted in more severe photochemical pollution in the PRD. In the second pattern, named as “O$_3$-Southwest”, northeasterly winds were dominant in the daytime over the PRD. Daytime elevated O$_3$ was mainly distributed in the southwestern inland regions, PRE, and southern coastal areas (e.g. Fig. 7b). Because a southeasterly sea breeze developed along the coastal areas and strengthened gradually during the afternoon and at dusk, transport of the O$_3$ plume to the southern water area was delayed, and O$_3$ remained at high levels in the southern rural land areas of Jiangmen and Guangzhou even after sunset. In the third pattern, named as “O$_3$-West”, the easterly winds prevailed in lower layers during the daytime over most areas, and elevated O$_3$ usually occurred in the western PRD in the afternoon. In this pattern, Jiangmen experienced a relatively light O$_3$ pollution compared with the episodes on O$_3$-Southwest days, and high levels of O$_3$ were seldom found along the southern coastal area except for the PRE (e.g. Fig. 7c).

We group all days during 4–31 October into the aforementioned three patterns (Table 4). The classification of pollution days reflects the influence of the interaction between synoptic forcing and local circulation on the evolution of O$_3$ pollution over the PRD during the campaign.

3.3 Process analysis of O$_3$ formation

Here, 16–22 October is used for the IPR analysis because the PRD experienced all three O$_3$ pollution patterns during this period (see Table 4), and the CMAQ simulation shows good performance at most sites on these days. The observed maximum height of the daytime ABL was about 1200 m during the campaign (Fan et al., 2008), corresponding to the lowest seven layers in the CMAQ simulation. Therefore, our process analysis mainly focuses on the IPR results from layers 1 to 7.

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**Fig. 6.** Average hourly simulated surface O$_3$ at 15:00 LST during October 2004.
Fig. 7. Simulated surface O₃ concentrations superimposed with the wind fields at 15:00 LST on (a) 19 October, (b) 16 October, and (c) 29 October, taken as examples of the three O₃ evolution patterns, respectively.

Table 4. Classification of O₃ pollution days into three evolution patterns in the PRD during 4–31 October 2004.

<table>
<thead>
<tr>
<th>No.</th>
<th>Pattern</th>
<th>Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>O₃-South</td>
<td>4, 5, 8, 18, 19, 25, 26</td>
</tr>
<tr>
<td>2</td>
<td>O₃-Southwest</td>
<td>6, 7, 9, 10, 11, 12, 14, 15, 16, 17, 20, 24, 27</td>
</tr>
<tr>
<td>3</td>
<td>O₃-West</td>
<td>13, 21, 22, 23, 28, 29, 30, 31</td>
</tr>
</tbody>
</table>

3.3.1 Process analysis at the selected site locations

Figures 8 and 9 reveal the different roles of atmospheric processes in the evolution of O₃ at the grid cells of the GDEMC, Xinken, and Donghu sites. In the lower layers of the urban GDEMC site, gas-phase chemistry exhibited a significant consumption of O₃ during the whole day (especially during traffic rush hours) due to O₃ titration by high NOₓ emissions, whereas horizontal and vertical transports were the main contributors to compensate for the chemical loss and to enhance the O₃ levels during the daytime (Fig. 8a). During the build-up of daytime maximum O₃ in the ABL from 09:00 to 15:00 local standard time (LST), the top chemical contributions occurred in the layers 5–7 (150–1000 m above ground level – a.g.l.); the produced O₃ was then horizontally transported to downwind areas and vertically transported to lower layers to balance the O₃ removal by NO titration and dry deposition (Fig. 9a).

At the rural Xinken site, vertical transport contributed mainly to the O₃ levels and was then decreased on a similar magnitude by horizontal transport during the daytime (Fig. 8b). Such influences of transport processes may be associated with the circulations of sea–land breezes over the PRE. The analysis of the simulated 3-D air flows showed that local air circulations often occurred at Xinken and were characterized by downdraft in the upper layers and divergence in the lower layers (Detailed discussion of this issue is presented in another manuscript in preparation by Zhou et al., 2009: land-sea breezes over Pearl River Estuary and their impact on local air quality in October 2004). Gas-phase chemistry was also an important contributor to daytime O₃ enhancement, characterized by more O₃ production in lower layers (Fig. 9b).

Among the three sites (GDEMC, Xinken, and Donghu) shown in Fig. 9, Donghu experienced the maximum increase of O₃ concentration in the ABL from 09:00 (0.046 ppm) to 15:00 LST (0.121 ppm) (Fig. 9c). Chemical production dominated the O₃ enhancement, especially in upper layers (layers 3–7). In lower layers, although the influences of horizontal and vertical transports exhibited day-to-day variations due to different meteorological conditions, the net effects of transport mainly showed a positive contribution to near-ground O₃ in most hours of a day (Fig. 8c).
Dry deposition was a sink for surface O$_3$ at GDEMC and Donghu, but showed weaker effects at Xinken because water covers more than 90% of the grid cell in which this coastal site is located. The wet deposition and cloud processes were also negligible due to the low cloudiness and the absence of precipitation during this period.

In many O$_3$ chemistry studies, the total oxidant O$_x$ (estimated by O$_3$+NO$_2$) is frequently used in data analysis because O$_x$ is not affected by the rapid photodissociation of NO$_2$ and the titration of O$_3$ with NO, being a better measure of the real photochemical production of ozone (Chou et al., 2006; Zhang et al., 2008b). In layers 1–7, the overall average O$_x$ chemical productions at GDEMC, Donghu, and Xinken were 0.102 ppm/6 h, 0.082 ppm/6 h, and 0.048 ppm/6 h, respectively, indicating that the photochemical oxidant production rate in the urban area was greater than that in the rural region, similar to findings from a study of O$_3$ pollution in Taipei (Chou et al., 2006). As shown in Fig. 10, an interesting phenomenon is that the vertical profiles of O$_x$ changes derived from gas-phase chemistry and accumulated from 09:00 to 15:00 LST were similar at GDEMC, Xinken, and Donghu (O$_x$ chemical production decreases with height), in contrast to the significant differences among the vertical profiles of O$_3$ chemical formation illustrated in Fig. 9.

As pollutants are generally well mixed in the ABL during daytime conditions, we summarize the process budgets integrated over the depth of the daytime ABL (corresponding to the vertical range of layers 1 to 7) in Table 5. In the ABL, the process contributions to daytime maximum O$_3$ at GDEMC, Xinken, and Donghu share common features regardless of their urban vs. rural locations: (1) although horizontal and vertical transports accounted for a considerable portion of process budgets, the net effects of transport processes exhibited negative contributions to the build-up of maximum O$_3$, and (2) gas-phase chemistry dominated the O$_3$ enhancement from morning to mid-afternoon, when O$_3$ concentrations usually reached the maximum level of the day.

### 3.3.2 Process analysis applied to regional O$_3$ pollution

In this section, the process analysis focuses on the whole PRD region during its daytime O$_3$ build-up in the ABL from 09:00 to 15:00 LST. On 16, 19, and 22 October (characterized by different near-ground flow conditions and O$_3$ pollution patterns; refer to Table 4), gas-phase chemistry played a dominant role in the O$_3$ enhancements over most of the area of the PRD, especially in the southern and western PRD where elevated O$_3$ was distributed around mid-afternoon. In contrast, the transport process (the sum of horizontal and
vertical transports) exhibited less influence on the regional scale, except in isolated areas, where the large power plants or the busy harbors were located and intensive NO\textsubscript{x} emissions resulted in strong O\textsubscript{3} titration (Fig. 11). The dominant contribution of chemical production not only occurred on the days with different near-ground flow conditions, but also over the areas with different emission strengths (i.e., urban vs. rural).

The lower precursor emissions in Jiangmen and southwestern coastal areas (Fig. 2) indicate that the transport process might play an important role in maintaining the necessary levels of precursors involved in the elevated O\textsubscript{3} chemical production there. The regional distribution of NO\textsubscript{x} contributed by the transport process, integrated over the depth of layers 1 to 7 and accumulated from late afternoon (17:00 LST) to the next morning (09:00 LST), is used to reflect the redistribution of NO\textsubscript{x} from the source regions to the whole PRD (Fig. 12). The IPR results show that the transport process functions as a sink of NO\textsubscript{x} in areas with intensive NO\textsubscript{x} emissions (such as in urban areas of Guangzhou, Dongguan, Shenzhen, and Hong Kong), but as the dominant source of NO\textsubscript{x} over the southern or southwestern PRD (especially in Jiangmen and the PRE), even if the accumulated time range is extended to 15:00 LST of the next afternoon.

The flow conditions influence the transport and redistribution of NO\textsubscript{x} during the night and next morning (Fig. 12). From late afternoon to midnight on 15 and 21 October, southeasterly sea breezes developed and prevailed in the southern part of the PRD, and wind convergence appeared over Panyu, Jiangmen, and Zhongshan (Fan et al., 2008), which mixed the NO\textsubscript{x} plumes from northern source areas (i.e., urban Guangzhou and Foshan) with plumes from southern areas (i.e., Dongguan, Shenzhen, and Hong Kong) inside the PRD; this hindered the transport of NO\textsubscript{x} outside the PRD. After midnight, northerly or northeasterly wind again prevailed over the PRD, and the delayed NO\textsubscript{x} plume was transported to the south and west, providing the majority of NO\textsubscript{x} involved in O\textsubscript{3} enhancement by the daytime photochemical process in Jiangmen and southwestern coastal areas. The transport of NO\textsubscript{x} from the late afternoon of 18 October to the morning of 19 October was mainly controlled by northerly winds and was not affected by the sea breeze, which resulted in less NO\textsubscript{x} accumulation inside the PRD than what happened on 15–16 and 21–22 October.

The above process analysis not only exhibits the redistribution of precursors from night to the next morning, but also reveals the significance of O\textsubscript{3} chemical production on a large regional scale in the daytime under such close interactions among the precursor emissions, physical transport, and photochemical process.

### 3.4 Ozone production efficiency

As illustrated in Fig. 13, OPE shows significant spatial variations over the PRD. The calculated OPEs are 2–8 in most regions of the middle and western PRD, corresponding to the areas of high O\textsubscript{3} chemical production (Fig. 11a). Less efficiency (i.e., OPE values of 1–5) is shown in the urban areas of Guangzhou, Foshan, Shenzhen, and Hong Kong, characterized by intensive NO\textsubscript{x} emissions, as well as downwind areas of the urban plumes under prevailing northeasterly flow. In contrast, high OPEs (larger than 11) are shown over upwind or rural areas with less NO\textsubscript{x} source emissions (e.g., Huizhou, Zhaoqing, and Qingyuan).

For the period of 16–22 October, the average OPE values during 09:00–15:00 LST were 3.6 (2.9–4.0) at GDEMC, Xinken, and Donghu sites (ppm/6 h).

<table>
<thead>
<tr>
<th>Site</th>
<th>CHEM*</th>
<th>HTRA</th>
<th>VTRA</th>
<th>DDEP</th>
<th>NETC</th>
</tr>
</thead>
<tbody>
<tr>
<td>GDEMC</td>
<td>0.072</td>
<td>0.004</td>
<td>−0.030</td>
<td>−0.002</td>
<td>0.044</td>
</tr>
<tr>
<td>Xinken</td>
<td>0.055</td>
<td>−0.031</td>
<td>0.025</td>
<td>0.000</td>
<td>0.049</td>
</tr>
<tr>
<td>Donghu</td>
<td>0.082</td>
<td>−0.021</td>
<td>0.016</td>
<td>−0.002</td>
<td>0.075</td>
</tr>
</tbody>
</table>

* CHEM: gas-phase chemistry; HTRA: horizontal transport, the net effect of horizontal advection and diffusion; VTRA: vertical transport, the net effect of vertical advection and diffusion; DDEP: dry deposition; NETC: net change due to all processes.
Fig. 12. (a), (b) and (c) The accumulated contributions to NO\textsubscript{x} by the transport process over the depth of layers 1 to 7 from 17:00 to 00:00 (left), 09:00 (center), and 15:00 LST (right) of the next day. The arrows show the near-ground dominant wind directions during the periods of 17:00–24:00 LST (left), 00:00–09:00 LST (center), and 09:00–15:00 LST (right). (d) The accumulated NO\textsubscript{x} emissions in the same time range and vertical layers during 15–16 October are shown for comparison.

4.9 (4.2–5.9) at Xinken, and 5.0 (3.3–7.3) at Donghu. NO\textsubscript{x} exhibits lower catalysis efficiency at the urban GDEMC site because of intensive local NO\textsubscript{x} emissions by vehicles and industrial activities. For the rural Xinken site, despite lower local emissions, the significant NO\textsubscript{x} sources in neighboring areas (such as power plants along the southern coast of Dongguan and intense NO\textsubscript{x} emissions in the western coastal area of Shenzhen and in Hong Kong) provide NO\textsubscript{x} for local photochemical reactions, resulting in a similar level of OPE as that found at Donghu (urban site). The simulated OPEs at the super sites are comparable to those observed in urban Beijing, ranging from 3.9 to 9.7 (Chou et al., 2009), and those observed or modeled in US cities (Nashville, 2.5–4.0, Nunnermacker et al., 1998; New York City, 2.2–4.2, Kleinman et al., 2000) and in the Mexico City Metropolitan Area (4–10, Lei et al., 2007).

3.5 Sensitivity of O\textsubscript{3} to precursors

In this section, we first examine the response of surface O\textsubscript{3} to precursor emission reductions, averaged over 12:00–17:00 LST from 16 to 22 October (Fig. 14). Over the central inland PRD and the PRE and surrounding coastal areas (i.e., the region marked by the red ellipse in Fig. 14), O\textsubscript{3} levels decrease (by 0.006–0.019 ppm) with the reduction of only VOC emissions; however, O\textsubscript{3} levels increase (by 0.003–0.017 ppm) in most of the areas with the reduction of only NO\textsubscript{x} emissions. Such O\textsubscript{3} changes suggest that O\textsubscript{3} formation was under a VOC-limited regime and was depressed by high NO\textsubscript{x} levels in these urban areas and the immediate downwind areas during the campaign period, which is consistent with Zhang et al.’s (2008b) findings from an observation-based model at the GDEMC and Xinken sites. The O\textsubscript{3} levels are more sensitive to the perturbation of NO\textsubscript{x} emissions but exhibit less changes with VOCs reductions in the rural southwestern PRD (i.e., the region marked by blue ellipse in Fig. 14), where less NO\textsubscript{x} and low-to-moderate VOC emissions are distributed (see Fig. 2). The O\textsubscript{3} response supports the important role of NO\textsubscript{x} regional transport to O\textsubscript{3} formation in the southwestern PRD, as obtained by the process analysis in Sect. 3.3.2. With reductions in both NO\textsubscript{x} and VOCs emissions, O\textsubscript{3} levels decrease over most of the central and western PRD.

Next, we check the response of O\textsubscript{3} photochemical production rates to the perturbations of precursor emissions because of the critical influence of gas-phase chemistry on O\textsubscript{3} enhancement from low concentrations in morning to maximum levels in mid-afternoon over most parts of the central and western PRD. Figure 15a and b show that the responses of the net chemical production rates of O\textsubscript{3} \([P(O_3)]\) are similar to the changes of O\textsubscript{3} in the central and western PRD (see Fig. 14a and b), except for the obvious decrease in the eastern PRD with the reduction of NO\textsubscript{x} emissions (i.e., a clear
NO\textsubscript{X}-sensitive response). The distribution of the NO\textsubscript{X}/NO\textsubscript{Y} ratio is also presented for understanding the responses of $P(O\textsubscript{X})$ (Fig. 15c). The VOC-sensitive area over the PRD generally corresponds to the areas of higher NO\textsubscript{X}/NO\textsubscript{Y} ratios, whereas the decrease of $P(O\textsubscript{X})$ due to the reduction of NO\textsubscript{X} emissions usually occurs in the regions of lower NO\textsubscript{X}/NO\textsubscript{Y} ratios, similar to results from the Mexico City Metropolitan Area (Lei et al., 2007). Generally, in and near urban cores, air plumes are exposed to fresh pollutants (high NO\textsubscript{X}/NO\textsubscript{Y} ratios) and tend to fall into the VOC-limited regime and even the NO\textsubscript{X} disbenefit regime (i.e., an increase in $P(O\textsubscript{X})$ resulting from a decrease in NO\textsubscript{X}). However, in suburban and rural regions, air plumes tend to maintain low NO\textsubscript{X}/NO\textsubscript{Y} due to the rapid removal of NO\textsubscript{X} by the chemical process in the course of transport to those regions, and O\textsubscript{3} production also tends to shift from VOC-limited to NO\textsubscript{X}-limited chemistry for the same reason (NARSTO, 2000).

We then further investigate the relationships between $P(O\textsubscript{X})$ changes and NO\textsubscript{X}/NO\textsubscript{Y} ratios at the GDEMC, Xinken, Donghu, Kaiping, and Duanfen sites (Fig. 16). Kaiping and Duanfen are located in the central and southern Jiangmen city, respectively (see Fig. 1), which are selected here to investigate O\textsubscript{3} photochemical sensitivity in the southwestern inland and coastal areas of the PRD. Although the distribution of NO\textsubscript{X}/NO\textsubscript{Y} ratio shows significant difference (i.e., GDEMC is characterized by high ratios, but Kaiping and Duanfen are dominated by low values), the responses of $P(O\textsubscript{X})$ share some common features at the five sites. In the case of 25% reduction of NO\textsubscript{X} emissions, the percentage change in $P(O\textsubscript{X})$ tends to increase from about $-30\%$ to $30\%$ with increasing NO\textsubscript{X}/NO\textsubscript{Y} ratio, whereas a decreasing tendency is obtained in the condition of VOCs emission reduction. The VOC benefit (i.e., a decrease in $P(O\textsubscript{X})$ resulting from a reduction in VOCs) and the NO\textsubscript{X} disbenefit suggest an obvious VOC-limited chemistry under higher NO\textsubscript{X}/NO\textsubscript{Y} conditions.
(>0.6, corresponding to fresh pollutants), whereas the opposite responses of $P(O_3)$ indicate the NO$_x$-limited regime occurring under lower NO$_x$/NO$_y$ conditions (<0.3, corresponding to chemically aged pollutants). The range of the NO$_x$/NO$_y$ ratio from about 0.4 to 0.6 seems to be a common transition between NO$_x$- and VOC-limited chemistry at all five sites.

### 4 Summary

The MM5/SMOKE/CMAQ modeling system was applied to investigate O$_3$ pollution over the PRD region during the PRIDE-PRD2004 campaign. Model performance was assessed by comparing simulated O$_3$, NO$_x$, and NMHC concentrations with measurements from two super sites (GDEMC and Xinken) of the campaign and from PRD regional air quality monitoring networks. The model reasonably reproduced the ozone episodes observed during the 1-month campaign.

Elevated O$_3$ levels were usually observed in the southwestern inland PRD, PRE, and southern coastal areas, resulting from the intensive precursor emissions in the central PRD and the dominant northerly or easterly winds during October. Three evolution patterns of simulated surface O$_3$ are categorized based on the different near-ground flow conditions, and more than 75% of the days occurred under the interaction between weak synoptic forcing and local sea-land circulation.

For daytime O$_3$ evolution at the urban GDEMC site, gas-phase chemistry serves as sink in lower layers (0–80 m a.g.l.) but as a source in upper layers (80–1000 m a.g.l.). The significant chemical removal of O$_3$ near the surface is compensated by the contributions of vertical transport from upper layers and horizontal advection from upwind areas. The rural Xinken site is characterized by strong local circulations, resulting in significant vertical inflow and comparable horizontal outflow. In this area, photochemical production serves as a steady contributor during the daytime. The IPR results on overall O$_3$ evolution in the ABL over the PRD suggest that photochemical production is the dominant contributor to O$_3$ enhancement from 09:00 to 15:00 LST, and regions of high chemical production generally coincide with those of elevated O$_3$ levels. The range of simulated OPE is 2–8 O$_3$ molecules per NO$_x$ molecule oxidized in most areas of high O$_3$ chemical production.

Compared with previous studies, an in-depth understanding of the regional O$_3$ formation over inland PRD area is obtained by process analysis. Through the transport process during nighttime and morning, O$_3$ precursors originating from northern source areas (i.e. urban Guangzhou and Foshan) and from southern areas (i.e. Dongguan, Shenzhen and Hong Kong) are usually mixed and transported to western or southern rural areas, where they are then involved in the daytime O$_3$ photochemical production. Such close interactions among precursor emissions, physical transport, and gas phase chemistry resulted in significant O$_3$ chemical production on a large regional scale in the daytime. The sea-land circulations played an important role on the regional O$_3$ formation and distribution over PRD during the campaign.

The O$_3$ sensitivity to precursors was investigated by comparisons of simulated O$_3$ levels between the base case and emission-control scenarios. Formation of O$_3$ is VOC-limited and NO$_x$-depressed in the central inland PRD, PRE, and surrounding coastal areas, whereas O$_3$ levels show more sensitivity to NO$_x$ in the rural southwestern PRD. The significant spatial variations of O$_3$ production sensitivity show a close relationship to the chemical aging of air plumes. VOC-limited chemistry usually dominates in less chemically aged air (NO$_x$/NO$_y$ >0.6), whereas NO$_x$-limited chemistry generally occurs within chemically aged plumes (NO$_x$/NO$_y$ <0.3).

The spatial variation of O$_3$ formation sensitivity suggests a non-uniform precursor emission-reduction strategy for O$_3$ pollution control in sub-regions of the PRD. Considering the source emissions concentrated in urban areas and the significant precursor transport to downwind rural areas, reductions in both VOC and NO$_x$ emissions, combined with more emphasis on VOC controls, appear to be practical for lowering the O$_3$ levels over the PRD region.

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