Influence of synoptic condition and holiday effects on VOCs and ozone production in the Yangtze River Delta region, China

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Abstract

Both anthropogenic emission and synoptic conditions play important roles in ozone (O3) formation and accumulation. In order to understand the influence of synoptic condition and holiday effects on ozone production in the Yangtze River Delta region, China, concentrations of speciated volatile organic compounds (VOCs) and O3 as well as other relevant trace gases were simultaneously measured at the Station for Observing Regional Processes of the Earth System (SORPES) in Nanjing around the National Day holidays of China in 2014, which featured substantial change of emissions and dominated by typical anticyclones. Different groups of VOC species and their chemical reactivities were comprehensively analyzed. We observed clear diurnal variations of short alkenes during the measurement period, considerable amount of short alkenes were observed during night (more than 10 ppb) while almost no alkenes were measured during daytime, which might be attributed to different chemical processes. The obvious enhancement of the VOC tracers during the National Day holidays (Oct. 1st-Oct. 7th) indicated that the holiday effect strongly influenced the distribution of VOC profile and chemical reactivity in the atmosphere. At the same time, two meso-scale anticyclone processes were also observed during the measurement period. The synoptic condition contributed to the accumulation of VOCs and other precursors, which consequently impacted the ozone production in this region. The integrated influence of synoptic and holiday effects was also analyzed with an Observation Based Model (OBM) based on simplified MCM (Master Chemical Mechanism) chemical mechanism. The calculated relative increment reactivity (RIR) of different VOC groups revealed that during the holidays, this region was in VOC-limited regime and the variation of RIR shows a close linkage to the development and elimination of anti-cyclones, indicating an in-negligible contribution of synoptic effect toward ozone production in this region.

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1. Introduction

Ozone (O3), one of the major components of photochemical pollutants in the troposphere, jeopardizes human immune system, nerve system as well as respiratory system (Brunekreef and Holgate, 2002). It also affects, negatively, the crop yield and forest growth (Fuhrer et al., 1997). Moreover, ozone is one of the major greenhouse gases influencing climate change on global scale (Seinfeld and Pandis, 2006; IPCC, 2013). The photochemical reactions of volatile organic compounds (VOCs) and nitrogen oxides (NOx) are the main sources of tropospheric ozone (Chameides, 1992; Atkinson, 1998; Arey and Atkinson, 2003).

Intensive anthropogenic VOCs and NOx emissions and their variations directly connect with ozone formation rate in urban and semi-urban areas (Zhang et al., 2008; Guo et al., 2009). It is a widespread phenomenon that, during holiday or weekend, daytime surface ozone concentrations in urban areas tend to have different variation pattern compared with normal days. Studies about holiday effects from Chinese New Year indicated a negative impact on the air quality. Ozone concentration in Taiwan was higher than normal days at every hour of the diurnal cycle (Tan et al., 2009,
The campaign covered the Chinese National Day holidays of 2014, when strong changes of human activities were expected, such as a reduced industrial production and increased traffic transportation. Meanwhile, anti-cyclone processes are quite typical in East Asia during this season. The anti-cyclones could provide favorable conditions of pollutant accumulation and the cold front tend to lead to the regional transport of pollutants (Ding et al., 2013a). The impacts of emission variation during the National Day holidays and the specific synoptic weather are addressed in this paper. In-situ observation of NO$_x$, carbon monoxide (CO), ozone, and VOCs combined with retroroule footprint analysis and an Observation based model (OBM) calculation based on the simplified MCM (master chemical mechanism) chemical mechanism CRI (Common Representative Intermediates) V2.1 (Jenkins et al., 2008) were applied in this work. The rest of this paper is structured as follows: Sect. 2 describes the site location and observation and modeling methodology. In Sect. 3, we analyze the VOC speciation, chemical reactivity, the ozone formation potential of different subgroups and its linkage with synoptic and holiday effects.

2. Field observations and modeling methodology

2.1. Observational station and instrumentation

The campaign was conducted at the SORPES station in Nanjing University Xianlin Campus, a suburban measurement site in the northeast to Nanjing downtown (118°57’10”E, 32°07’14”N), during Sept. 22nd–Oct. 7th, 2014. The easterly prevailing wind and synoptic condition makes it a representative background site of Nanjing and a regional, downwind site of the city cluster in the YRD region (Ding et al., 2013a, 2016).

Online measurement with GC-MS/FID was conducted to obtain consistent VOC concentration with time resolution of 1 h, which included 5-min sampling period, 40-min analysis period and 8-min flush and cooling down. Short alkanes and alkenes (C2-C4) were separated with an Al2O3 PLOT column (DM-PLOT:30m × 0.32 mm i.d.) and quantified with a flame ionization detector (FID). Larger molecular VOCs (C5-C12 alkanes, alkenes, aromatic compounds, halides and oxygenated VOCs) were separated on a semipolar column (DB624: 60M × 0.32 mm i.d. × 1.8 μm) and quantified with a quadruplet mass spectrometer (SHIMADZU QP2010-ULTRA). The initial column temperature of DM-PLOT and DB624 was set as 35 °C, kept for 3 min, raised to 180 °C with heating rate of 6 °C/min and then held for 5 min. Bromochloromethane, 1,4-difluorobenzene and 1-bromo-3-fluorobenzene were used as internal standards for MS calibration. Table S1 and Table S2 list the 97 VOC species that GC-FID and MS measured respectively.

Trace gases (O$_3$, sulfur dioxide (SO$_2$), NO, NO$_x$, total reactive nitrogen oxides (NO$_x^r$), and CO) at the SORPES station have been measured since 2011, with 90% data coverage and 1-min time resolution. Instruments (Thermo-fisher Scientific, TEI 49I, 43I, 42I, 42I-Y and 48I) were auto span and zero checked with frequency of one week and one day, respectively, and manually calibrated once a week in case of the dis-function of auto calibration program. The TEI 42I was coupled with a highly selective photolytic converter to measure NO$_2$ (Xu et al., 2013). The 5-min averaging data were adopted considering both the capability of describing the concentration variation and proper signal noise ratio. Because of the high ozone concentration, daytime NO concentration appeared to be equivalent to the detection limit of NO detector used at the SORPES station. The NO detection limit was set to be 3 times of the standard deviation of the zero calibration data, which was around 0.5–1.2 ppb.
Meteorological measurements including relative humidity (RH), wind speed, wind direction, and air temperature were recorded by Automatic Weather Station (CAMPEEL co., AG1000). Consistent observations have been started since 2010. UVB total radiation was measured by UVB radiometer (UVS-B-T UV Radiometer, KIPP & ZONEN).

2.2. Modeling methodology

2.2.1. Lagrangian backward dispersion modeling

In order to diagnose the transport and dispersion characteristic of the air masses during the campaign, Lagrangian dispersion model was conducted by using Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Stein et al., 2015). Briefly, for each hour during the study period, the model was run 2-day backwardly with 3,000 particles released every hour from the altitude of 100 m above the site. The model calculated the particle position with mean wind and turbulence transport component after being released at the receptor point. The residence time of particles below 100 m level was used to identify the “footprint”retroplume. The spatiotemporal distributions of these particles were used to evaluate the contribution from different regions to the pollutant transport. This method has been applied successfully to simulate transport of air pollution in eastern China and the Pearl River Delta region (Ding et al., 2013a,b). The reported results show that this method can well characterize the different transport processes of air pollution.

2.2.2. Observation-based chemical modeling

An observation-based chemical box model was used to investigate ozone chemistry and the sensitivity of ozone production to its precursors (Cardelino and Chameides, 1995, 2000; Cheng et al., 2010; Xue et al., 2013a). The model was based on a simplified MCM(CRI.2.1), a reduced mechanism describing ozone formation from the degradation of methane and 115 primary non-methane hydrocarbons and oxygenated volatile organic compounds, using the Master chemical Mechanism version 3.1(MCM v3.1) as a reference benchmark (Jenkin et al., 2008).

Before each simulation, the model was pre-run for seven days with constraints of observation data every hour, for the purpose that the unmeasured species would reach a steady state. The observed VOCs, other trace gases like CO, NO, NO2, temperature, RH, UVB total radiation were used as inputs for model calculation in order to ensure that the simulation was under proper ratio of NOx and VOCs and real-time meteorology and radiation conditions. The measured real-time UVB radiation data was used to scale photochemical rate constant fitted from solar zenith angle under clear sky condition, which is calculated within the model (Saunders and Jenkin, 2003). For the NO data used in the model, all afternoon and low concentration NO data were set to be 1 ppb. The sensitivity of ozone concentration to the changes of ozone precursors (10% reduction) was investigated following the concept of Relative Incremental Reactivity (RIR), which is defined as the percent change of ozone photochemical production, with per percent change of one of its precursor or certain combination of some precursors (Carter and Atkinson, 1989). For RIR calculation, the model read VOCs, NO and CO concentrations every hour to constrain the simulation, and read air temperature, RH and scaled radiation every time step to calculate the kinetic rate constants.

2.2.3. Positive matrix factorization (PMF)

PMF was a well-developed and widely used receptor model in identifying sources in the atmosphere (Ling et al., 2011; Zhang et al., 2013; Yan et al., 2016). The principle was based on the work of Paatero and Tapper (1994). The advantage of PMF was that it can provide source profiles and contribution matrices without any prior knowledge and constrain the solutions to be non-negative. The measured 97 primary VOCs including alkanes, alkenes, aromatic compounds and halogenated hydrocarbons observed from Sept. 22nd – Oct. 7th were used as input of PMF. The uncertainties for each species were set to be 0.2 and the detect limit to be 0.015 ppb.

3. Results and discussion

3.1. VOC concentration and speciation

Table 1 shows the average concentration and variation of 30 most abundant VOCs measured during the campaign compared with results from previous study of Guangzhou and other 43 Chinese cities (Barletta et al., 2005; Liu et al., 2008a,b). The concentration of alkanes ranged from tens of ppt (part per trillion) to tens of ppb (part per billion) level with an average of 8.39 ppb, which was comparable with the range of previous studies. But the concentration level of alkenes and aromatic compounds was much lower compared with the result from Guangzhou. Moreover, the upper range of alkanes measured at the SORPES station exceeded the previous study from 43 Chinese cities. These differences might be attributed to different measurement method, site representativeness, and the chemical reactivity of different groups of VOCs. The VOC measurement conducted at Guangzhou and 43 Chinese cities was based on canister sampling at city sites and off-line analysis, while the observation conducted at the SORPES station was continuous on-line measurement. The direct emission of vehicle and other local sources near these measurement sites might be the reason for a higher concentration level of aromatic compounds and alkenes. The air parcel measured at the SORPES station was aged and well-mixed after passing through the city cluster. The low-reactivity species (alkanes, Benzene, etc.) accumulated and most of highly reactive species (Toluene, Propylene, etc.) were oxidized during transport. The off-line measurement at these city sites tended to overestimate the concentration levels of reactive species and underestimate the effect of low reactive but much more abundant species.

Fig. 1c and d shows the time series of total VOCs (tVOCs) measured at the SORPES site and the ratio of different subgroups to tVOCs. The measured 97 hydrocarbons are classified into 9 subgroups based on the structure of molecules and reactivity with OH radical, which are light alkanes (C1-C4), long chain alkenes (C4>C5), short chain alkenes (C1-C4), long alkenes (C5>C6), BTEX (benzene, toluene, ethyl-benzene, xylene), other aro (other aromatic compounds), halides, alkylene and OVOCs. Halides, alkylene and short chain alkenes are low reactivity hydrocarbons, while alkenes, long chain alkanes, aromatic compounds (except benzene) and OVOCs are defined as reactive hydrocarbons (Zhang et al., 2007). The reason for grouping alkenes based on their molecular structure is that the two groups have different diurnal patterns. The separation of BTEX out of other aro is because the majority of aromatic compounds is constituted of these four species. Short alkenes are the most dominant group of VOCs, followed by BTEX, long chain alkanes and halides. Such speculation agreed with other studies in China (Barletta et al., 2005). Moreover, we observed clear diurnal variation of alkenes. The concentration level of short alkenes was relatively low during daytime while increase dramatically during nighttime, indicating different chemical processes. Specifically, the reactions with OH radical and ozone is the main sink during daytime, while NO3 radical related reactions and ozonolysis take the place of the most dominant chemical processes during night (Arey and Atkinson, 2003; Atkinson, 1998; Carter, 2007). Inferred from high surface concentration of NO during nighttime (Fig. 1b), NO3 radical was barely present in the atmosphere and the rate...
coefficient of alkenes with NO3 radical was much lower than that with OH radicals, which explained the high concentration of alkenes observed during night. One exception during the campaign was Sept. 23rd, 2014, considerable amount of ozone (~30 ppb) and short alkenes observed during the night. One exception during the campaign was Sept. 23rd, 2014, considerable amount of ozone (~30 ppb) and short alkenes observed during the night (Fig. 1b and c). Though no simultaneous measurement of radicals, we can infer from the ozone and NO concentration level that ozone and NO2 radical are the important oxidants that consume alkenes during night. The observed Sept. 23rd case could support the assumption that the diurnal variation of short alkenes is related to different pathways of oxidation reactions.

Table 1

<table>
<thead>
<tr>
<th>Species</th>
<th>SORPES Range</th>
<th>SORPES Average ± std</th>
<th>CO2 Average ± std</th>
<th>43 cities ± std</th>
<th>Range ± std</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethane</td>
<td>1.04–9.36</td>
<td>2.65 ± 1.21</td>
<td>5.58 ± 3.34</td>
<td>3.7–17.0</td>
<td></td>
</tr>
<tr>
<td>Acetylene</td>
<td>0.30–7.32</td>
<td>1.56 ± 1.08</td>
<td>7.3 ± 5.2</td>
<td>2.9–58.3</td>
<td></td>
</tr>
<tr>
<td>Propane</td>
<td>0.23–9.87</td>
<td>1.48 ± 1.10</td>
<td>10.35 ± 8.53</td>
<td>1.5–20.8</td>
<td></td>
</tr>
<tr>
<td>n-Butane</td>
<td>0.15–18.22</td>
<td>1.42 ± 1.56</td>
<td>5.07 ± 4.42</td>
<td>0.6–14.5</td>
<td></td>
</tr>
<tr>
<td>1,2-Dichloroethane</td>
<td>0.16 ± 1.51</td>
<td>6.55 ± 4.28</td>
<td>2.1–34.8</td>
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<td></td>
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<tr>
<td>Iso-Butane</td>
<td>0.08–23.79</td>
<td>1.11 ± 1.86</td>
<td>2.7 ± 2.3</td>
<td>0.3–18.8</td>
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</tr>
<tr>
<td>t-Butane</td>
<td>0.12–11.02</td>
<td>0.90 ± 0.96</td>
<td>2.9 ± 2.6</td>
<td>0.4–4.6</td>
<td></td>
</tr>
<tr>
<td>Benzene</td>
<td>0.01–13.19</td>
<td>0.83 ± 1.28</td>
<td>1.19 ± 1.07</td>
<td>0.2–7.7</td>
<td></td>
</tr>
<tr>
<td>Toluene</td>
<td>0.15–4.44</td>
<td>0.72 ± 0.53</td>
<td>2.4 ± 1.9</td>
<td>0.7–10.4</td>
<td></td>
</tr>
<tr>
<td>MEK</td>
<td>0.13–2.38</td>
<td>0.65 ± 0.41</td>
<td>7.0 ± 7.3</td>
<td>0.4–11.2</td>
<td></td>
</tr>
<tr>
<td>Propylene</td>
<td>n.a.–3.37</td>
<td>0.53 ± 0.42</td>
<td>3.2 ± 3.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1,2-Dichloroethane</td>
<td>0.07–2.65</td>
<td>0.35 ± 0.24</td>
<td>0.1–2.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acetone</td>
<td>0.12–2.27</td>
<td>0.30 ± 0.15</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethylbenzene</td>
<td>0.03–2.43</td>
<td>0.30 ± 0.32</td>
<td>1.16 ± 1.22</td>
<td>0.0–5.6</td>
<td></td>
</tr>
<tr>
<td>Chloromethane</td>
<td>0.08–0.35</td>
<td>0.34 ± 0.03</td>
<td>1.18 ± 1.21</td>
<td>0.1–6.9</td>
<td></td>
</tr>
<tr>
<td>2-Methylpentane</td>
<td>0.03–0.80</td>
<td>0.19 ± 0.13</td>
<td>1.03 ± 0.94</td>
<td>0.2–10.1</td>
<td></td>
</tr>
<tr>
<td>1,2-Dichloropropane</td>
<td>0.02–2.22</td>
<td>0.18 ± 0.24</td>
<td>1.16 ± 1.21</td>
<td>0.0–5.6</td>
<td></td>
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<tr>
<td>m/p-Xylene</td>
<td>0.01–2.69</td>
<td>0.17 ± 0.23</td>
<td>1.46 ± 1.42</td>
<td>0.2–5.2</td>
<td></td>
</tr>
<tr>
<td>n-Hexane</td>
<td>0.02–0.91</td>
<td>0.17 ± 0.23</td>
<td>0.84 ± 0.80</td>
<td>0.1–5.0</td>
<td></td>
</tr>
<tr>
<td>O-Xylene</td>
<td>0.01–2.16</td>
<td>0.13 ± 0.15</td>
<td>0.52 ± 0.5</td>
<td>0.0–4.1</td>
<td></td>
</tr>
<tr>
<td>1-Butene</td>
<td>n.a.–2.11</td>
<td>0.11 ± 0.22</td>
<td>1.33 ± 0.91</td>
<td>0.0–4.1</td>
<td></td>
</tr>
<tr>
<td>3-Methylpentane</td>
<td>0.01–0.90</td>
<td>0.10 ± 0.09</td>
<td>0.67 ± 0.64</td>
<td>0.0–4.1</td>
<td></td>
</tr>
<tr>
<td>2,3-Dimethylbutane</td>
<td>0.02–1.00</td>
<td>0.10 ± 0.10</td>
<td>0.26 ± 0.24</td>
<td>0.0–4.1</td>
<td></td>
</tr>
<tr>
<td>Isopropene</td>
<td>n.a.–0.82</td>
<td>0.09 ± 0.12</td>
<td>0.22 ± 0.17</td>
<td>0.0–4.1</td>
<td></td>
</tr>
<tr>
<td>Acetonitrile</td>
<td>0.01–0.30</td>
<td>0.09 ± 0.04</td>
<td>0.66 ± 0.29</td>
<td>0.0–4.1</td>
<td></td>
</tr>
<tr>
<td>Carbon Terachloride</td>
<td>0.04–1.35</td>
<td>0.08 ± 0.10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n-Dodecane</td>
<td>0.01–0.43</td>
<td>0.08 ± 0.10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1-Pentene</td>
<td>n.a.–1.26</td>
<td>0.07 ± 0.12</td>
<td>0.18 ± 0.14</td>
<td></td>
<td></td>
</tr>
<tr>
<td>cis-1,1-Dichloroethene</td>
<td>n.a.–0.41</td>
<td>0.06 ± 0.07</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trans-2-Butene</td>
<td>0.01–1.68</td>
<td>0.05 ± 0.08</td>
<td>0.4 ± 0.36</td>
<td>0.01–3.4</td>
<td></td>
</tr>
<tr>
<td>MTBE</td>
<td>n.a.–1.30</td>
<td>0.05 ± 0.05</td>
<td>0.96 ± 0.94</td>
<td>0.02–3.7</td>
<td></td>
</tr>
<tr>
<td>cis-2-Butene</td>
<td>n.a.–0.47</td>
<td>0.01 ± 0.01</td>
<td>0.38 ± 0.33</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a Data from (Barletta et al., 2005).

b Data from (Liu et al., 2008b).

c Under detection limit.

The averaged mixing ratios (ppb) of VOCs measured by GC-FID/MS at the SORPES station and corresponding values in other studies.

During the campaign, alkanes was the most abundant group, followed by BTEX, alkylene, and short alkenes (only during night time).The averaged concentration of nighttime tVOCs was much higher than that during daytime, implying a substantial role of photochemical process in VOC consumption and dilution effect of boundary layer development. Moreover, the variation of CO correlated well with tVOCs variation, indicating CO and VOCs could have similar sources. NOx concentration rose during the holiday (Oct. 1st-Oct. 7th), and the highest VOC concentration level during the campaign was observed on Oct 3rd. It may link to the holiday effects and the impact of VOCs and NOx on ozone production will be discussed in the following section. Fig. 2 displays the diurnal variation of ozone, the total oxidants Ox (=O3 + NO2), NOx (=NO + NO2), and tVOCs. The diurnal variation of tVOCs has a negative correlation with the variation of Ox than with ozone, indicating that Ox is a better indicator of atmospheric oxidizing capacity than ozone. The validation of using CO:NOx proxy to investigate the NOx-VOC chemical mechanism is proved with VOCs observations, which correlated well with CO.

The ratio of ambient concentrations of VOCs with similar chemical reactivity equals to their ratio of original emission rate (Goldan et al., 2000; Jobson, 2004). When this theory is applied to real situations, complicated emission sources would obscure the
Fig. 1. The time series of [a] meteorology parameters (air temperature, RH, wind speed and direction), [b] trace gases (O3, CO, NO, and NOx), [c] different groups of VOCs. [d] Hourly concentration of total VOCs (tVOCs, ppb) and the ratios of different VOC subgroups to tVOCs during the measurement period.
inter-species comparison results. The combined use of photochemical age analysis would reveal more information about the source of specific VOCs.

The photochemical age of air masses was estimated by the ratio of observed toluene and benzene. The photochemical age was calculated using the method of de Gouw et al. (2005):

\[ t = \frac{1}{[\text{OH}]} \times [\text{toluene}] - [\text{benzene}] \times \left[ \text{ln} \left( \frac{[\text{toluene}]}{[\text{benzene}]} \right) \right], \]

where \([\text{OH}]\) is the 24-h averaged concentration of OH radicals \((3.0 \times 10^4 \text{ molecules cm}^{-3})\) (Warneke, 2004), and \(k_{\text{toluene}} (5.63 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})\) and \(k_{\text{benzene}} (1.22 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})\) are the rate coefficients for the reaction of OH with toluene and benzene (Arey and Atkinson, 2003). The emission ratio of toluene to benzene was calculated based on the linear fit of the canister sampling data from tunnel at Nanjing \(\text{[toluene]} \text{[benzene]} = 3.16\). Fig. 3a shows the scatterplot of ethyl benzene \((k = 7.0 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})\) versus acetylene \((k = 1.22 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1})\), colored by the photochemical age derived from equation (1). The slope of ethyl benzene/acetylene decreases with the increase of photochemical age. The rate constant of ethyl benzene is about six times larger than that of benzene which indicates that removal rate of ethyl benzene is much faster. The change of slope with photochemical age can be explained by their different reactivities, which indicates that the photochemical age derived from toluene and benzene ratio can, on some extent, describe the process of the removal of VOCs with different rate coefficients.

Benzene and acetylene have similar rate coefficient with OH radical, about \(1.22 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}\). Fig. 3b shows the scatterplot of benzene versus acetylene, also colored by the photochemical age. It shows a good correlation of benzene and acetylene with the correlation coefficient of 0.726 indicating that benzene and acetylene have similar sources. It is well acknowledged that acetylene is the product of incomplete combustion processes (Goldan et al., 2000; Liu et al., 2008b), while benzene is associated with vehicles emission (Wang et al., 2002), thus the source of benzene and acetylene is mainly related to fuel combustion of vehicular. The scatterplot also shows that the correlation of benzene and acetylene decreases with increase of photochemical age indicating the mixed air masses bring benzene from other sources such as industry emission and solvent vaporization. Figs. 3c and 3d display scatterplot of \(n\)-butane and \(iso\)-butane versus propane. \(n\)-butane is the tracer of LPG evaporation and LPG powered vehicular emission (Liu et al., 2008a,b). A good correlation of both \(n\)-butane and \(iso\)-butane with propane was observed, with correlation coefficient of 0.796 and 0.804, respectively. The slope of \(n\)-butane and \(iso\)-butane versus propane was 1.212 and 0.75, respectively, different from the result from Guangzhou (0.48 and 0.28) (Liu et al., 2008b), and Mexico City (0.458 and 0.210) (Bravo and Sosa, 2002). The dispersed dots with longer photochemical age in Fig. 3c and d may be explained by long-range transport. The PMF results (Figs. S6–S8) resolved sources of LPG usage, which also indicated that propane, butanes and pentanes were the main contributions of LPG related sources. Based on the diurnal variation of the contributions, we further distinguished traffic emissions with a clear two-peak shape corresponding to rush-hours. These results imply that LPG evaporation and LPG vehicular emission were important sources of light alkanes at the SORPES station. The different butane/propane ratio from previous studies suggests a different fuel usage in Nanjing. In fact, a program of use both natural gas and LPG to drive vehicles involving all taxis and part of private cars started since 2012. When vehicles drive with low speed, the fuel supply system would use natural gas instead. This would increase the evaporation of LPG.

To further understand the VOC sources, 6 factors were resolved for the VOC measurement during Sept. 22nd–Oct. 7th with PMF analysis. Because of short data span, the uncertainties of PMF results were inevitable. Factor profiles and contributions are shown in Figs. S6–S8. The factors were identified as LGD usage, traffic emission, biogenic emission, transported plume, PX-related industry and other industry. Factor 1 was dominated by acetylene, propane and \(n/i\)-butanes. Propane and butanes are traces of LDG (Blake and Rowland, 1995; Ling et al., 2011), and acetylene is the main constituents of internal combustion. Consequently, factor 1 was identified as LPG usage. Factor 2 was similar with Factor 1, but with much higher \(i/n\) pentane constitution. The diurnal variation of its contribution showed two peaks during rush hours. Therefore, Factor 2 was identified as traffic emission. Factor 3 showed a clear diurnal pattern with the contribution increased with the increase of temperature and solar radiation and was characterized by isoprene and its oxidant products (methyl vinyl ketone (MVK) and methacrolein (MACR)). Thus, Factor 3 was assigned to biogenic emissions. Factor 5 was characterized with mixed VOC groups including alkanes, OVOCs and aromatic compounds. The
contribution of this factor increased only during Sept. 25th–Sept. 29th. Thus, Factor 5 was characterized as transported plume based on the contribution time series. Factors 4 and 6 were defined as industry emissions because both of them had considerable fraction of benzene and toluene. The difference was that Factor 6 was also characterized with xylenes and the diurnal pattern indicated a stronger emission during nighttime. Since there is a p-Xylene (PX) chemical plant in Xianlin district, Factor 6 was further assigned as PX related industry.

3.2. Co-effect of synoptic condition and national holiday on VOCs

The distinct air masses transport characteristic and the holiday effect on the VOC speciation were observed during 2014 campaign.

![Fig. 3. Scatterplots of (a) acetylene versus ethyl benzene, (b) acetylene versus benzene, (c) propane versus n-butane and (d) propane versus iso-butane. The scatter points are colored by photochemical age derived from observed toluene/benzene ratio, based on equation (1).]

![Fig. 4. Averaged retroplume (footprint residence time) of air parcels during working days (a) before the holiday, i.e. Sept. 24th–Sept. 30th and (b) during the National Day holidays of China, i.e. Oct. 1st–Oct. 7th.]

Fig. 4 shows the averaged retroplume during working days before (Sept. 24th—Sept. 30th) and during the National Day holidays (Oct. 1st—Oct. 7th). Fig. 4a displays the average retroplume of Sept. 24th—Sept. 30th, the footprint shows two branches with one transport northeastward from south Japan, East China Sea and northern part of Jiangsu province, and another from adjacent southeastern provinces (Zhejiang, Fujian and Jiangxi). The average footprint distribution of Fig. 4a suggests a long footprint residence time over East China Sea and southeastern provinces of China. The clean air masses from the ocean and the low anthropogenic emission rate in southeast China (Fig. S1) leads to the relatively low concentration level of the observed NO and VOCs.

Fig. 4b shows the average retroplume of Oct. 1st—Oct. 7th, air masses mainly transported through northern part of Jiangsu province, Shandong province and adjacent Anhui and Henan provinces before arriving at the SORPES station. The effect of northern China was dominant, and the high footprint residence time reveals that the northern part of Jiangsu province is the region that contributes most to the observations during this period. The hourly mean footprint shows two anti-cyclone processes on Oct. 2nd and Oct. 5th, which transported air masses from northern region at a very fast speed. The development of these two anticyclones could be seen from the daily averaged retroplume of each day's transport characteristic in the support information (Fig. S1). The two anti-cyclones developed during Oct. 2nd—3rd and Oct. 5th—6th (Fig. S3), transporting air masses to the observational site from northern Jiangsu province and adjacent provinces. The first one was stronger, which brought air masses from nearly all Jiangsu and Anhui provinces and even from Henan and Shandong provinces, accompanied with sharp decrease of temperature, strong wind change and precipitation. The second one was much weaker that only northern Jiangsu province and Shandong province had a high footprint residence time, and the measurement site only experienced strong north wind. The accumulation of air pollutants during anti-cyclone was observed (Fig. 1). The tVOCs concentration increased sharply from 5.5 ppb to 71.2 ppb, and CO concentration increased from 0.33 ppm to 1.68 ppm, accompanied with strong wind shift from near static weather condition to more than 5 m/s and the highest daily temperature decrease from 26.7 to 19.6 Celsius degree. There was no much difference of NOx concentration during the anti-cyclone processes; the high NO/NOx ratio indicated that local emissions dominated the variation of NOx. These two convergent processes together with severe increase of traffic emission may play an important role in the ozone chemistry in the YRD region. When anti-cyclone weakened and vanished, the transport characteristic shifted to northeastward, which was quite typical during this season (Oct. 4th and 7th).

Significant increase of NOx was observed during the National Day holidays (Fig. 5d) and the highest VOC concentration level during the campaign was observed on Oct. 3rd (Fig. 1). Since family vacation is a tradition during long holidays in China, we prefer to use transportation-related VOC species to estimate the holiday effect. As discussed above, acetylene measured at the SORPES station is mainly from vehicle emission and alkenes such as propene, ethylene and 1,3-butene are the characteristic products of vehicles, thus acetylene and alkenes were chosen as tracers to estimate the human activity intensity. Fig. 5a shows diurnal variations of short alkenes/tVOCs ratio during the holidays and pre-holiday working days. Both periods show clear diurnal variations, alkene ratio kept deceasing from morning and were barely measured during the midday due to its high chemical activity. Compared with working days, during nighttime, the alkene ratio increased significantly, especially during 18:00—23:00 local time, corresponding to the variation of human activity. Fig. 5b uses acetylene as tracer to investigate the holiday effect. The diurnal variation of working days shows clearly the signal of morning rush hour around 06:00—10:00 local time. During national holiday, the acetylene ratio generally increased, and the obvious increase of acetylene ratio during 08:00—23:00 local time corresponded to the time of intensive human activities indicating an enhanced traffic emission. Figs. 5c and d6 demonstrate the diurnal variations of benzene/toluene ratio and NOx before and during the holidays, respectively. The benzene/toluene ratio was applied to estimate the relative importance of traffic and industrial emissions as used in Liu et al. (2008b). During working days, the two peaks during rush hours (4:00—6:00, 18:00—22:00 local time) indicated that the benzene/toluene ratio could describe the variation of traffic emission. During the holiday, the ratio increase about 50%, compared with the value during working days. The higher benzene/toluene could be explained by enhanced vehicle emission and favorable convective condition during the holidays. Moreover, PMF analysis (Fig. S7) also showed an increase trend of LPG usage and traffic emissions at Oct. 3rd, 5th and 6th, consistent with high VOC episodes.

3.3. Chemical reactivity of VOCs and ozone production

The most important and direct process that increases ozone level in troposphere is the photolysis of NO2 and O3 tends to be decreased by the titration reaction of NO with O3. If sufficient NOx is present in the atmosphere, which is the normal state in urban and suburban areas of China, the reactions of VOCs with atmospheric oxidants (mainly with OH radicals, if exist) forming peroxy radicals, which either consume NO or convert NO to NO2, determine the ozone formation rate (Carter, 1994). OH (kOH) reactivity is used here to estimate the initial formation rate of peroxy radical and to evaluate the chemical reactivity during different time of a day. This method is also used in previous studies (e.g. Liu et al., 2008b), which considers the first chemical step of full atmospheric chemistry with OH radical, and gives a general estimation of the contribution of this VOC species to atmospheric chemistry during daytime. kOH is calculated by multiplying the OH reaction rate coefficient and the mixing ratio of the given compounds:

\[
    k_{OH} = [VOC]_i \times k_{OH}^{i}.
\]

Where [VOC] is the mixing ratio of the ith VOC compound and \(k_{OH}^{i}\) is the corresponding OH reaction rate coefficient. Here, the rate coefficients published by Arey and Atkinson (2003) were used in this work.

The same method is used on the nighttime data to estimate the relative contribution of each group of VOCs to the chemical reactivity during nighttime, using the chemistry reaction rate coefficient with NO3 radical (\(k_{NO3}^{i}\)) as a substitute to \(k_{OH}^{i}\). It should be noted that though NO3 radical is very important in the nighttime chemistry, only a few studies have reported the rate coefficient with alkenes, aromatic hydrocarbons, and oxygenated hydrocarbons. In this work we used NO3 reaction rate coefficient with alkenes and aromatic hydrocarbons reported by Atkinson et al. (1984). In terms of aldehydes, values suggested by Cababas et al. (2001) were applied.

Fig. 6 shows the averaged OH reactivity (during daytime) and NO3 reactivity (during nighttime) of different groups of VOCs. Short chain alkenes contributed most to the chemical reactivity, followed by short alkanes and BTEX during morning, and the OH reactivity of short chain alkenes showed a peak during morning rush hours and gradually decreased during daytime, which might be explained by the diurnal variations of vehicular emissions. At rush hours, i.e. 07:00—09:00 and 16:00—20:00 local time, the increase of the chemical reactivity of alkenes (both short chain and long alkenes) and BTEX was also observed, indicating the effect of traffic emission.
on the total chemical reactivity of VOCs. NO3 reactivity was used to estimate the chemical contribution during nighttime, since NO3 radical reactions are important during nighttime in the troposphere (Atkinson, 1998). The NO3 loss rate is about two factors smaller compared with OH loss rate, which could explain the diurnal variation of reactive alkenes (Fig. 1). Since the reactions with NO3 radical are much slower, more short-chain alkenes could be observed during nighttime. Alkene-NO3 reactions were the dominant processes during nighttime, contributed to more than 90% of NO3 reactivity, with the relative NO3 reactivity of OVOCs and BTEX being less than 10% and 1%, respectively. Discrepancies of calculated OH reactivity and total OH reactivity have been reported from multiple studies, which might be related to the unmeasured and unidentified VOCs and the uncertainties of measurements (Yang et al., 2016). Unfortunately, there was no measurement of total OH reactivity during this campaign. However, the calculated OH reactivity and NO3 reactivity does provide some information on contribution from specific VOC to the intensity of photochemistry. The variation of the OH reactivity of short chain alkenes indicated that the chemical processes after sunrise would be important. The accumulated reactive VOCs during night react with OH radical to produce considerable amount of peroxy radicals, which would not only enhance the ozone production but also secondary organic aerosol formation. Moreover, the NO3 reactivity analysis during nighttime also revealed that short alkene reactions were the dominated processes. The night time reactions of alkenes with ozone and NO3 radicals would be an important source of organic nitrate (Atkinson, 1998), especially in urban and suburban areas where the high concentration of alkenes was observed (Fig. 1) due to heavy traffic and industry load.

RIR was used as an index to assess the relative importance of the precursors to ozone production. The diagnosed RIRs of major cities during several ozone pollution episodes were studied by Xue et al. (2013b). The OBM results indicated that Shanghai and Guangzhou...
were VOC-limited regions with the dominant precursors to be aromatics and alkenes. Lanzhou, as an inland city with heavy petrochemical industry load, was a mixed regime, with alkenes, especially light olefins such as propene and ethene to be the most important VOC precursors. The RIRs diagnosed in Nanjing were consistent with the results in Shanghai.

The co-effect of synoptic condition and holiday effects on ozone production and the sensitivity of different precursors were discussed in this section. Fig. 7 shows the RIRs of major ozone precursors (upper panel) during the campaign and the RIRs of subgroups of VOCs (lower panel), which is categorized as reactive aromatic compounds (R-ARO), reactive olefins (R-OLE), alkanes which have more than 4 carbon atoms (C≥4), and oxygenated VOCs (OVOCs). Generally, during the holiday the site was in VOC-Limited regime and the effect of CO cannot be neglected. The negative RIR value of NO suggested that the reduction of NO may cause the increase of ozone pollution in this area. While the control of both CO and VOCs emission would, in some extent, alleviate ozone issue. On Oct. 2nd, Oct. 5th and Oct. 6th when convergent process dominated, the RIR of CO increased compared with Oct. 1st and Oct. 7th when air masses were transported from northeast. The sensitivity test of the subgroups of AHC indicated that OVOC was the least important group contributed to ozone production. The other three subgroups competed to be the dominant precursor of ozone pollution under different synoptic conditions.

The simulated RIR results showed that on Sept. 30th, ozone...
production rate was not sensitive to the reduction of its precursors (Fig. 7). The measurement data showed that on Sept. 30th the air mass was clean, with the lowest ozone concentration observed during the campaign (the ozone concentration was less than 40 ppb during the midday) and VOC concentration was also very low, with tVOCs concentration being less than 10 ppb and low active alkanes constituting more than half of the total amount. Furthermore, the UVB measurement showed that the radiation intensity was quite weak during the midday. The footprint of air mass suggested that the branch transported from the East China Sea and southeastern China was the dominant sources during the week before the holidays. The clean air mass and weak radiation might be the reason of low sensitivity of the precursors towards ozone formation.

During the National Day holidays, air mass mainly transported from North China, and anti-cyclone processes led to the accumulation of VOCs and other precursors. The RIR variation also reflected connections between transport and ozone production characteristics. On Oct. 2nd and Oct. 6th, during which time the anti-cyclone was most intensive, the effect of CO overweighed VOCs showing a non-neglectful effect of long-life species brought by regional transport (Fig. 7). The retroplume analysis indicated that the first anti-cyclone started and developed at Oct. 2nd, weakened and vanished at Oct. 3rd. The sharp increase of the RIR of CO was most likely attributed to the strong long-range transport. Similarly, the regional transport from north and northwest during Oct. 5th afternoon and Oct. 6th morning brings high concentration of CO over 1.6 ppm, and the increasing RIR of CO from Oct. 4th to Oct. 6th was corresponded to the development of the anti-cyclone. On both cases, high CO concentration was observed during the night, the accumulated CO would react with OH radical during daytime to produce considerable amount of HO2 radical, which would directly enhance the ozone production.

Along with the increase RIR of CO, the RIRs of VOC subgroups also changed with the development of anti-cyclone. On Oct. 1st and Oct. 4th air masses were transported from northeast, and the RIR results showed that aromatic compounds were the most important precursors that led to ozone production in these two days which was consistent with the simulation results (Tie et al., 2013; Zhang et al., 2015). During Oct. 2nd—Oct. 3rd and Oct. 5th—Oct. 6th, when two strong regional transport processes from north and northwest were observed, the different RIR distributions of VOC subgroups were found to be in consistent with the change of transport pattern. RIR of both cases showed significant increase of alkenes and alkanes compared with Oct. 1st and Oct. 4th, and alkenes became the most dominant precursors of ozone production. As showed in Fig. 5, during the holidays the traffic emission was strongly enhanced. The co-effect of enhanced human activities and suitable synoptic weather condition made alkenes the most sensitive precursor towards ozone formation, and the role of alkanes, which were also related to motor vehicle emission, was also enhanced. During Oct. 6th, the effect of alkenes even outweighed aromatics compounds, indicating that under strong regional transport condition, the effects of low reactive VOC species were also important.

The low RIR of both VOC and CO on Oct. 3rd could be explained by the weather condition. The high RH of whole day and weak diurnal variation of temperature (Fig. 1) indicated that precipitation during daytime hindered the photochemical processes, and titration effect led to the high negative RIR of NO.

4. Conclusions

For the purpose of understanding the influence of synoptic condition and holiday effects on ozone production in the Yangtze River Delta region, concentrations of speciated volatile organic compounds (VOCs) and ozone were simultaneously measured at the Station for Observing Regional Processes of the Earth System (SORPES) around the National Day holidays of 2014. When compared with canister samples and off-line measurements from other studies, here we observed high alkanes and low aromatics compounds and alkenes, which was related to the chemical reactivity and the age of air mass. Clear diurnal variations of short alkenes were observed, high concentration level of short alkenes could only be observed during the night and the boundary layer effect was excluded by the diurnal variations of the ratio of subgroups to total VOCs. This diurnal variation of short chain alkenes could be attributed to the relatively low reaction rate coefficient with NO3 radical during nighttime.

The inter-species comparison coupled with photochemical age derived from toluene/benzene ratio indicated that benzene and acetylene were related to vehicle emissions, while benzene had other sources, such as industry emissions and solvent usage. The results also showed that LPG usage was the main sources of propane and butane and other short chain alkanes. Among all the species, short alkenes were the most important subgroup of VOCs during nighttime and contributed to more than 90% of chemical reactivity, while during daytime clear signal of rush hour were observed from VOC chemical reactivity.

Relative Incremental Reactivity (RIR) was calculated based on OBM, and the co-effects of synoptic condition and holiday effects were investigated. Acetylene, short alkenes and Toluene/Benzene ratio were used as tracers of traffic emissions to investigate the holiday effect during national holiday. Strong signals of human activity during “non-sleep” hours were observed compared with working days. Meanwhile, pollutant accumulation was observed during anti-cyclone processes: The tVOCs concentration increased from 5.5 ppb to 71.2 ppb, and CO concentration increased from 0.33 ppm to 1.68 ppm. The relative contributions of CO, NO and subgroups of VOCs to ozone production were estimated with OBM. When industrial emission dominated, corresponding to the northeastern retroplume, aromatic compounds were the most important precursors of ozone production. When air masses shifted to a more regional transport pattern under the effect of anti-cyclone processes, the co-effect of intensive human activities during national holidays and regional transport of the air pollutants from north and northwest of Nanjing led to a significant enhancement role of CO, alkenes and alkanes on ozone production.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http:// dx.doi.org/10.1016/j.atmosenv.2017.08.035.

References


Further reading


