Rates and regimes of photochemical ozone production over Central East China in June 2006: a box model analysis using comprehensive measurements of ozone precursors


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Abstract. An observation-based box model approach was undertaken to estimate concentrations of OH, HO2, and RO2 radicals and the net photochemical production rate of ozone at the top of Mount Tai, located in the middle of Central East China, in June 2006. The model calculation was constrained by the measurements of O3, H2O, CO, NO, NO2, hydrocarbon, HCHO, and CH3CHO concentrations, and temperature and J values. The net production rate of ozone was estimated to be 6.4 ppb h−1 as a 6-h average (09:00–15:00 CST), suggesting 58±37 ppb of ozone is produced in one day. Thus the daytime buildup of ozone recorded at the mountain top as ∼23 ppb on average is likely affected by in situ photochemistry as well as by the upward transport of polluted air mass in the daytime. On days with high ozone concentrations (hourly values exceeding 100 ppb at least once), in situ photochemistry was more active than it was on low ozone days, suggesting that in situ photochemistry is an important factor controlling ozone concentrations. Sensitivity model runs for which different NOx and hydrocarbon concentrations were assumed suggested that the ozone production occurred normally under NOx-limited conditions, with some exceptional periods (under volatile-organic-compound-limited conditions) in which there was fresh pollution. We also examined the possible influence of the heterogeneous loss of gaseous HO2 radicals in contact with aerosol particle surfaces on the rate and regimes of ozone production.

1 Introduction

Central East China (CEC) is regarded as one of the regions in the world where tropospheric ozone is present at high concentrations. Pochanart et al. (2009) revealed the seasonal variation in ozone concentrations over CEC from observation. They found that at the top of Mount Tai (36.26°N, 117.11°E, 1534 m a.s.l., Shandong Province of the People’s Republic of China) located in the middle of CEC, monthly averaged ozone concentrations regularly peak in June at ∼80 ppbv, with highest hourly values exceeding 120 ppbv in each year of 2004–2006. Considering that the site is not directly affected by plumes from large cities, a geographically wide area (with a scale of several hundred kilometers) is likely covered by air masses containing high concentrations of ozone. Zhu et al. (2004) indicated in their regional modeling study that a zonal band (between 35 and 40°N) of high ozone concentration (>70 ppb) emerged from the western boundary (70°E) of their model domain to 130°E in June, the latitudinal location of which was affected by seasonal transition associated with the Asian monsoon system. The regionally high ozone burden in this region is of interest from the viewpoints of the greenhouse effect, health effects, and agricultural damage.

The primary cause for the high ozone concentrations over this region would be photochemical production. Wang et al. (2006) simulated an episode of high ozone concentrations (around 110 ppbv) observed at Mt. Tai on 24 and 25 May 2004 with the Nested Air Quality Prediction Modeling System and found that the modeled ozone concentrations were...
at Mt. Tai had a large (20–50%) sensitivity to emissions in the Yangtze Delta region, suggesting that the photochemistry during the travel of the air mass from that region was important. Using a similar model, Li et al. (2007) calculated that photochemical production is the dominant term for the time derivative of ozone concentration for the grid surrounding Mount Tai station (surface area of 400×400 km with a 0–2.5 km altitude range) in June. However, an in situ production rate of ozone that is representative over this region has not yet been quantified on the basis of the observed concentrations of ozone precursors (NO\textsubscript{x}, CO, and hydrocarbons) and relevant parameters (\textit{J} values, temperature, and humidity).

Knowledge of the ozone production regime (NO\textsubscript{x}-limited vs. volatile organic compound (VOC)-limited) over this region is essential for the establishment of controlling strategies of ozone concentrations. These concentrations are expected to increase at least in the coming decade over CEC (e.g., Yamaji et al., 2008) owing to fast economic growth and the associated increase in the emission rates of ozone precursors (Ohara et al., 2007). In past modeling studies, Carmichael et al. (2003a) suggested that ozone production in the north-central region of CEC (north of 30–35° N) is VOC-limited at 440 m altitude in March, while it is NO\textsubscript{x}-limited south of ~30–35° N. Luo et al. (2000) similarly suggested that VOC limitation generally applies in late October–early November north of the Yangtze Delta. However, the precursor concentrations in the model simulations have not been validated through observations, making the analysis susceptible to large uncertainty in the emission rates of the precursors in this area (Streets et al., 2003). Martin et al. (2004) found that HCHO/NO\textsubscript{2} column concentration ratios derived from Global Ozone Monitoring Experiment satellite data are greater than 1 during summer over eastern China, indicating NO\textsubscript{x}-sensitive conditions. In this case, however, the observed column density of HCHO was used as a proxy to infer the VOC concentrations, which have not been observed directly. Considering that the regimes could change with seasons owing to variations in the production rates of radicals that mediate ozone production and in the amount of biogenic hydrocarbons, knowledge of the regimes over this region in this heavily polluted month is currently very limited.

In June 2006, we conducted the Mount Tai Experiment 2006 field campaign at the top of Mount Tai and measured ozone, its precursors (CO, NO, NO\textsubscript{2}, hydrocarbons, HCHO, and CH\textsubscript{3}CHO), and relevant meteorological/radiative parameters (\textit{J} values, temperature, and humidity). Using the observed concentrations and parameters as input to a photochemical box model employing a tropospheric chemistry mechanism, concentrations of OH, HO\textsubscript{2}, and RO\textsubscript{2} (organic peroxy) radicals were calculated first and then net in situ photochemical production rates of ozone were estimated on the basis of the radical concentrations. We also performed sensitivity model runs where different NO\textsubscript{x} or VOC concentrations were assumed to study the ozone production regimes at the location. Another set of model runs including heterogeneous loss of HO\textsubscript{2} on aerosol surfaces was executed to estimate its impact on the ozone production rate and regimes.

2 Experimental

\textit{O}_{3} and CO were measured by commercially available instruments based on ultraviolet and nondispersive infrared absorption (Thermo, models 49C and 48C) on a year-round basis (Pochanart et al., 2009). Purified air produced by a zero air generator (Thermo, model 111) was periodically introduced to the CO-measuring instrument to determine the zero level. NO, NO\textsubscript{x}, and NO\textsubscript{2} were sequentially detected by a customized instrument based on a commercially available instrument (Thermo, model 42CTL). An air sample for this instrument passed through one of three gas lines: a line with a molybdenum converter, a line with a blue light (light emitting diode) converter (Droplet Measurement Technology, USA), and a line without converters. The two converters were located at the entrance of the sampling tube, such that NO\textsubscript{y} and NO\textsubscript{2} are converted to NO, a relatively inert molecule, at an early part of the inlet line with minimum loss. The conversion efficiency of NO\textsubscript{2} to NO by the blue light converter during the campaign was 50%. The sensitivity to NO was determined against a premixed gas of NO/NO\textsubscript{2} (2.004 ppmv, Taiyo Nippon Sanso Corporation). The sensitivity agreed with that determined with a cylinder with NIST-traceability to within 2%. The detection limit of the instrument is specified to be 0.1 ppbv for NO and 0.2 ppbv for NO\textsubscript{2} and NO\textsubscript{y}.

VOC measurements were made with canister samplings (performed on 30 occasions during the campaign) followed by analysis using gas chromatography-flame ionization detection (GC-FID) and gas chromatography-mass spectrometry (GC-MS) (Suthawaree et al., 2009). The sampling was made typically once per day (typically at 14:40 CST), with a sampling duration of 2 min. Proton-transfer reaction-mass spectrometry (PTR-MS) was conducted in the latter part of the campaign (13–30 June) (Inomata et al., 2008, 2009). We used concentrations of HCHO, CH\textsubscript{3}OH, CH\textsubscript{3}CHO, isoprene, methylvinylketone (MVK) + methacrolein (MACR), methylethylketone (MEK), benzene, and \Sigma-monoterpene measured by PTR-MS for analysis in this paper. The detection limits were estimated to be 0.37–0.83, 0.74, 0.29–0.35, 0.28, 0.1, 0.08, 0.13, and 0.06 ppbv, respectively. It was assumed that the signal corresponding to m/z 71 in the PTR-MS was from MACR and MVK with 50% contributions. The spectral actinic flux was measured by a single-monochromator/photodiode array instrument (Meteorologie Consult Inc.) covering a wavelength region from 274 to 698 nm. The instrument was calibrated against a standard halogen lamp. The uncertainty in \textit{J}(O\textsuperscript{1}D) for a similar instrument has been estimated to be 14% (Heard et al., 2006).
Table 1. Hydrocarbons taken into account in the model calculation and their categorization in the RACM.

<table>
<thead>
<tr>
<th>RACM category</th>
<th>Hydrocarbons taken into account in the model calculation</th>
<th>Measured by GC, taken into account in the model by relationship with CO</th>
<th>Measured by PTR-MS, taken into account in the model by relationship with CO</th>
<th>Biogenic species measured by PTR-MS, averaged diurnal variations used in the model</th>
</tr>
</thead>
<tbody>
<tr>
<td>ETH ethane</td>
<td>ethane</td>
<td></td>
<td></td>
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<tr>
<td>HC3 propane,</td>
<td>i-butane, n-butane, acetylene</td>
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<td></td>
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<tr>
<td>HC5 i-pentane,</td>
<td>n-pentane, 2-methylpentane, 3-methylpentane, n-hexane</td>
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<td></td>
</tr>
<tr>
<td>HC8 methylcyclopentane, cyclohexane, 2-methylhexane, n-heptane, n-octane, n-decane</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>ETE ethene</td>
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<tr>
<td>OLT propene,</td>
<td>1-butene</td>
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<tr>
<td>OLI trans-2-butene, cis-2-butene, trans-2-pentene, cis-2-pentene, 2-methyl-2-butene, cyclopentene, trans-2-hexene, cis-2-hexene</td>
<td></td>
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<tr>
<td>DIEN 1,3-butadiene</td>
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<tr>
<td>ISO isoprene</td>
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<tr>
<td>API alpha-pinene, beta-pinene, camphene</td>
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<tr>
<td>LIM limonene</td>
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<tr>
<td>TOL toluene,</td>
<td>ethylbenzene</td>
<td></td>
<td>iso-propylbenzene, styrene</td>
<td>benzene</td>
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<tr>
<td>XYL p,m-xylene, o-xylene</td>
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<td></td>
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<tr>
<td>HCHO HCHO</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>ALD acetaldehyde</td>
<td></td>
<td></td>
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<tr>
<td>KET acetone, MVK, MEK</td>
<td></td>
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<tr>
<td>MACR methacrolein</td>
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</tbody>
</table>
Fig. 1. Averaged diurnal variations in (a) O₃, (b) CO, (c) NO, (d) NO₂, and (e) NOₓ concentrations, (f) J(O¹D), (g) total NMHC concentrations (not including oxygenated VOCs), and (h) summed reactivity of CO, CH₄, H₂, SO₂, and NMHCs toward OH for high-O₃ days (solid circles) and low-O₃ days (gray filled squares). In (h), open circles and gray open squares are the reactivity when the contribution from oxygenated VOCs is added for high- and low-O₃ days, respectively. Open triangles and gray open triangles are the reactivity of CO, CH₄, H₂, and SO₂ for high- and low-O₃ days, respectively (shown for reference).

3 Model calculations

The photochemical box model we used is based on the Regional Atmospheric Chemistry Mechanism (RACM) designed by Stockwell et al. (1997), employing 237 reactions and 77 chemical species. The model has been outlined elsewhere (Kanaya et al., 2007). The observed concentrations of O₃, CO, H₂O, NO, NO₂, hydrocarbons, HCHO, CH₃CHO, CH₃OH, MVK+MACR, and MEK, temperature, and J values were averaged or interpolated with a time resolution of 10 min and used as constraints of the model. Data gaps were basically filled by time interpolation. In the base run, a NO concentration of 0.05 ppb was assumed when the observation was below the detection limit. This applied to ∼30% of the midday 6-h periods (09:00–15:00 CST) of the studied period (3–30 June). Sensitivity runs assuming 0.02 or 0.1 ppb of NO instead of 0.05 ppb did not change the conclusion of the study significantly (see Sect. 4.2).

Table 1 shows the categorization of the observed hydrocarbons and oxygenated VOCs into lumped species used for the
RACM. The concentrations of major anthropogenic hydrocarbons (25 species, second column in Table 1) measured by GC-FID and GC-MS were taken into account using regression equations for CO, for which continuous observations were available. Two regression equations, one for the first half of the campaign period (3–15 June) and the other for the latter half (16–30 June), were produced and applied for the periods. The \( R^2 \) values for the correlations between CO and hydrocarbon concentrations ranged between 0.01 and 0.88. Some hydrocarbons had different slopes for their concentration against CO concentration in the two periods; it is likely that intense biomass burning of crop residues affected the concentration ratios in the first half period (Suthawaree et al., 2009).

Other minor hydrocarbons (21 species, third column in Table 1) detected by GC-FID and GC-MS were taken as having constant concentrations, calculated as mean values over the campaign period. For HCHO, CH\(_3\)OH, acetaldehyde (ALD), acetone, MVK+MACR, and MEK, hourly data from the PTR-MS instrument were used after 15:00 CST on 12 June, when the measurements began. Regression equations with CO concentrations were used for each of the oxygenated species to estimate their concentrations before 12 June.

For benzene, PTR-MS data were used after the measurements began (15:00 CST on 12 June). The benzene concentrations before this time were estimated from the relationship between CO and benzene (GC-FID) observed before 15 June. Concentrations of isoprene and monoterpenes were based on hourly data from PTR-MS after 15:00 CST on 12 June. In the case of monoterpenes, averaged fractions of \( \alpha \)-pinene, \( \beta \)-pinene, camphene, and limonene over the campaign were used to calculate species-specific concentrations. Averaged diurnal variations in isoprene and monoterpene concentrations from PTR-MS were applied to estimate their concentrations for the period prior to 12 June. For benzene, isoprene, and monoterpenes, the PTR-MS data were scaled to the concentrations determined by GC-FID using the slopes of their regression lines.

The uncertainty in the NMHC concentrations determined in these ways was roughly estimated to be a factor of 2. The benzene concentrations estimated by the method mentioned above showed similar variation patterns to those directly observed by PTR-MS after 12 June (see Fig. S1 of the supplement material: http://www.atmos-chem-phys.net/9/7711/2009/acp-9-7711-2009-supplement.pdf) and the central 80% of the ratio between the two concentrations was in a range of 0.80–1.67. Additionally, we found that 84% of the concentrations of important NMHCs (having large reactivity toward OH) estimated by the method above deviated from the directly measured concentrations by less than a factor of 2. The influence of the uncertainty on the estimation of the ozone production rate will be discussed in Sect. 4.3.

SO\(_2\) concentrations were estimated from the equation [SO\(_2\) (ppb)]=0.018×[CO (ppb)]+4.8, as observed at Lin’an in March 2001 (Carmichael et al., 2003b). CH\(_4\) and H\(_2\) concentrations were assumed to be constant as 1850 and 550 ppbv, respectively.

The model calculation was made for each day within the period of 3–30 June (28 days). The time of 00:00 CST was regarded as the initial time on each day, and integration over 24 h was conducted by 10-min binning of the data. This integration was conducted ten times in series to stabilize the concentrations of unconstrained species (e.g., unmeasured carbonyl and peroxide species). The results for the last 24 h were used as output for each day. The H\(_2\)O\(_2\) concentrations did not stabilize even by the 10th day of calculation, but the effect on the radical concentrations (and thus on the ozone production rate) was small.

Figure 1 shows averaged diurnal variations in OH, HO\(_2\), and RO\(_2\) concentrations estimated by a photochemical box model for high-O\(_3\) days (circles) and low-O\(_3\) days (gray squares).
Table 2. Major production and loss processes of OH, HO$_2$, and radical (OH+HO$_2$+RO$_2$) and their fractions as daytime averages (09:00–15:00 CST)$^a$.

<table>
<thead>
<tr>
<th>OH production</th>
<th>OH loss</th>
<th>HO$_2$ production</th>
<th>HO$_2$ loss</th>
<th>Radical production</th>
<th>Radical loss</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.3 × 10$^7$ cm$^{-3}$ s$^{-1}$</td>
<td>4.3 × 10$^7$ cm$^{-3}$ s$^{-1}$</td>
<td>3.8 × 10$^7$ cm$^{-3}$ s$^{-1}$</td>
<td>3.9 × 10$^7$ cm$^{-3}$ s$^{-1}$</td>
<td>2.1 × 10$^7$ cm$^{-3}$ s$^{-1}$</td>
<td>2.1 × 10$^7$ cm$^{-3}$ s$^{-1}$</td>
</tr>
<tr>
<td>HO$_2$+NO 0.59</td>
<td>OH+CO 0.27</td>
<td>OH+CO 0.30</td>
<td>HO$_2$+NO 0.65</td>
<td>O$_2$+D+H$_2$O 0.43</td>
<td>HC$_3$P+HO$_2$ 0.14</td>
</tr>
<tr>
<td>O$_3$+D+HO$_2$ 0.22</td>
<td>OH+ALD 0.12</td>
<td>CH$_3$O$_2$+NO 0.19</td>
<td>HO$_2$+HO$_2$ 0.07</td>
<td>HCHO+hv 0.09</td>
<td>HO$_2$+RO$_2$ 0.13</td>
</tr>
<tr>
<td>O$_3$+HO$_2$ 0.05</td>
<td>OH+OP2 0.08</td>
<td>HCHO+hv 0.05</td>
<td>HO$_2$+HO$_2$+H$_2$O 0.07</td>
<td>H$_2$O+hv 0.09</td>
<td>HO$_2$+RO$_2$+H$_2$O 0.12</td>
</tr>
<tr>
<td>H$_2$O$_2$+hv 0.04</td>
<td>OH+HC3 0.06</td>
<td>HC$_3$P+NO 0.05</td>
<td>O$_3$+HO$_2$ 0.06</td>
<td>MGLY+hv 0.08</td>
<td>CH$_3$O$_2$+HO$_2$ 0.10</td>
</tr>
<tr>
<td>OP2+OH 0.04</td>
<td>OH+ISO 0.05</td>
<td>HCHO+OH 0.05</td>
<td>HC$_3$P+HO$_2$ 0.04</td>
<td>OP2+hv 0.06</td>
<td>ISOP+HO$_2$ 0.06</td>
</tr>
<tr>
<td>OP2+hv 0.01</td>
<td>OH+MACR 0.04</td>
<td>ISOP+NO 0.03</td>
<td>CH$_3$O$_2$+HO$_2$ 0.03</td>
<td>MACR+OH 0.04</td>
<td>XO$_2$+HO$_2$ 0.06</td>
</tr>
<tr>
<td>LIM+O$_3$ 0.01</td>
<td>OH+HCHO 0.04</td>
<td>OH+SO$_2$ 0.03</td>
<td>XO$_2$+HO$_2$ 0.02</td>
<td>LIM+O$_3$ 0.04</td>
<td>XO$_2$+NO 0.05</td>
</tr>
<tr>
<td>OPI+OH 0.01</td>
<td>OH+H$_2$O$_2$ 0.03</td>
<td>H$_2$O$_2$+OH 0.03</td>
<td>ISOP+HO$_2$ 0.01</td>
<td>ALD+hv 0.03</td>
<td>OH+NO$_2$ 0.05</td>
</tr>
<tr>
<td>API+O$_3$ 0.01</td>
<td>OH+LIM 0.03</td>
<td>HC$_3$+OH 0.03</td>
<td>LIMP+HO$_2$ 0.01</td>
<td>API+O$_3$ 0.02</td>
<td>LIMP+HO$_2$ 0.04</td>
</tr>
<tr>
<td>OPI+hv 0.00</td>
<td>OH+SO$_2$ 0.03</td>
<td>others 0.24</td>
<td>others 0.05</td>
<td>MACR+hv 0.02</td>
<td>ACO$_3$+HO$_2$ 0.04</td>
</tr>
<tr>
<td>others 0.01</td>
<td>OH+NO$_2$ 0.02</td>
<td>others 0.23</td>
<td>others 0.11</td>
<td>OH+HO$_2$ 0.03</td>
<td>others 0.20</td>
</tr>
</tbody>
</table>

$^a$ Intercorversion between HO$_2$ and HO$_2$NO and between RO$_2$ and PANs is omitted. See Table 1 for LIM, API, HC$_3$, ALD, ISO, MACR, HC$_3$P, ISOP, and LIMP are peroxy radicals formed from HC$_3$, ISO, and LIM, respectively. XO$_2$ and ACO$_3$ are peroxy radicals that account for additional NO to NO$_2$ conversion and saturated acyl peroxy radicals (acetyl peroxy and higher). OPI, OP2, and MGLY are CH$_3$OOH, organic peroxide higher than CH$_3$OOH, and methylglyoxal and other α-carbonyl aldehydes.

4 Results and discussions

4.1 Estimated radical concentrations

Figure 2a–c shows diurnal variations in the calculated concentrations of OH, HO$_2$, and RO$_2$ radicals. They were calculated for individual days and then averaged for the high-O$_3$ and low-O$_3$ days, respectively. The OH concentrations for low-O$_3$ days were higher than those for high-O$_3$ days until 11:00 CST, because higher NO concentrations for low-O$_3$ days facilitated conversion from HO$_2$ to OH. In the afternoon, the OH concentrations were quite similar for the two cases. The midday peak concentrations were 5.4 and 5.7 × 10$^6$ cm$^{-3}$ for high-O$_3$ and low-O$_3$ days, respectively. The 24-h average concentrations were 1.7 and 1.9 × 10$^6$ cm$^{-3}$. The daytime HO$_2$ and RO$_2$ concentrations were always higher on the high-O$_3$ days; daytime peaks were 39 and 52 pptv in comparison with 29 and 34 pptv on the low-O$_3$ days.

Table 2 lists major production and loss processes of OH, HO$_2$ and radicals (the group of OH, HO$_2$, and RO$_2$ radicals) and their fractions to the total rates as daytime averages (09:00–15:00 CST). For OH production, the HO$_2$+NO reaction was more important than ozone photolysis (O$_3$+D+H$_2$O). OH loss was dominated by its reactions with CO, ALD, and OP2 (organic peroxy higher than CH$_3$OOH). HO$_2$ production was dominated by the OH+CO and CH$_3$O$_2$+NO reactions. HO$_2$ loss occurred mainly through its reactions with NO and HO$_2$ itself. The initial production of the radicals occurred by the photolysis of ozone, HCHO, and H$_2$O$_2$. The ultimate loss of the radicals was normally governed by the self and cross reactions of HO$_2$ and RO$_2$. It should be noted that conversion between the radicals and HNO$_4$, peroxyacetyl nitrate (PAN) and higher saturated PANs, and TPAN (unsaturated PANs) was not included in the production or loss of the radical to calculate values in Table 2.

The production and loss rates of the radicals were in balance. The initial radical production rate was 2.1 × 10$^7$ cm$^{-3}$ s$^{-1}$ as a daytime average, similar to an average (2.2 × 10$^7$ cm$^{-3}$ s$^{-1}$) for summer in the urban center of
Tokyo (as an average in the period 09:00–15:00 local time) (Kanaya et al., 2007).

4.2 Photochemical production rate of ozone and its breakdown

It is known that photochemistry results in both the formation and destruction of ozone in the troposphere and thus the difference between the rates of gross formation \(F(O_3)\) and loss \(D(O_3)\), \(F(O_3) - D(O_3)\) (hereafter simply termed \(F-D(O_3)\)), is the important quantity describing the net photochemical production rate. The terms are given by

\[
F(O_3) = k_1([HO_2][NO]) + \Sigma k_{2i}[RO_2][NO]\phi_i, \tag{1}
\]

\[
D(O_3) = k_3[O^1(D)][H_2O] + k_4([OH][O_3]) + k_5([HO_2][O_3]) + \Sigma k_{6ij}[O_3][\text{olefin}], \tag{2}
\]

\[
F - D(O_3) = F(O_3) - D(O_3), \tag{3}
\]

where \(k_1\)–\(k_5\) are the rate coefficients of the \(HO_2+NO\), \(RO_2+NO\), \(O^1(D)+H_2O\), \(OH+O_3\), \(HO_2+O_3\) and \(O_3+\text{olefin}\) reactions, respectively, and \(\phi\) is the yield of \(NO_2\) from the \(RO_2+NO\) reaction. \(F\), \(D\), and \(F-D\) are calculated on the basis of the calculated radical concentrations (Sect. 4.1) and then they are averaged to obtain composite diurnal variations of the rates for the high- and low-ozone days (Fig. 3a–c). \(F(O_3)\) was higher for the high-ozone days during the 10:00–18:00 CST period. This was because the peroxy radical concentrations were higher on the high-ozone days as seen in Fig. 2b. \(F(O_3)\) for the high-ozone days was slightly lower at 09:00 CST because the NO concentrations were lower than they were for the low-ozone days (Fig. 1c), compensating the higher peroxy radical concentrations when they are multiplied together.

The gross production of ozone per day was estimated to be 85±43 and 60±32 ppb for high- and low-ozone days, respectively, by integrating \(F(O_3)\) from morning (05:00 CST) to evening (20:00 CST). \(D(O_3)\) was larger for high-ozone days because it was proportional to the ozone concentration in the atmosphere. The integrated gross losses per day were 18±5 and 11±3 ppb for the high- and low-ozone days, respectively. Because of the larger compensation by \(D(O_3)\) during the high-ozone days, the difference in \(F-D(O_3)\) for the two periods was small. The integrated net production was estimated to be 67±41 and 49±32 ppb, respectively. As an average over the campaign period, 58±37 ppb of daily ozone production was expected. This value is larger than 32 ppb day\(^{-1}\), the photochemical production rate of ozone estimated in a model for the grid surrounding the Mount Tai station (surface area of 400×400 km with 0–2.5 km altitude range) in June (Li et al., 2007). This may be because the nighttime loss chemistry was taken into account and precursor concentrations were not constrained in Li et al.’s model calculation.

Our value is larger than the production rate (38 ppb day\(^{-1}\)) estimated from the slope of a regression line for the ozone concentration observed at the same location in 2004 and the residence time of the air mass over the Central East China region based on the backward trajectory analysis (Pochanart et al., 2009). The difference would be attributed to dilution and deposition, effective only for the rate estimated using the residence time.

Our results are relatively insensitive to the assumption that NO concentrations were set to 0.05 ppb when measurements were below the detection limit. The average daily net productions were estimated to be 53 and 66 ppb when assuming 0.02 and 0.1 ppb of NO instead of 0.05 ppb.

This analysis implies that 67 and 49 ppb of ozone were produced each day by in situ photochemistry on high- and low-ozone days, respectively. These amounts were larger than the daytime increases in the ozone concentration observed at the mountain top (34 and 15 ppbv), suggesting that the in situ photochemistry is active enough to explain the ozone buildup in both cases, while the transportation of polluted air from lower altitude, as recognized by the daytime buildup of primary pollutants, may also be important. It should also be noted that the ratio between the daily production amounts on
Fig. 4. Breakdown of $F(O_3)$ and $D(O_3)$ terms as averages over the campaign.

Fig. 5. Average diurnal variations in $F-D(O_3)$ for the base run (black) and sensitivity runs (colored).

The daytime 6-h averages of $F-D(O_3)$ and $F(O_3)$ were 6.4 and 7.9 ppb h$^{-1}$ for this campaign. The former value is in rough agreement with the in situ production rate ($\approx 5$ ppb h$^{-1}$) estimated in a three-dimensional regional scale modeling study (Li et al., 2008b). The latter value falls into a range of the rates estimated for the boundary layer over major cities in the United States; our value is slightly lower than those over Philadelphia (11.3 ppb h$^{-1}$ for July and August 1999) and Houston (11.3 ppb h$^{-1}$ for August and September 2000), but higher than those over Nashville (6.2 ppb h$^{-1}$ for June and July 1995), New York City (4.3 ppb h$^{-1}$ for July 1996), and Phoenix (3.5 ppb h$^{-1}$ for May and June 1998) (Kleinmann et al., 2002). The production rates for Mount Tai are significantly higher than the $F-D(O_3)$ values at other mountain stations: from $-0.05$ to $+0.4$ ppb h$^{-1}$ at Jungfraujoch (3590 m a.s.l.) for midday in winter, spring, and summer (Parker et al., 2009; Zanis et al., 2003), 2–3 ppb d$^{-1}$ for Mt. Cimone (2165 m a.s.l.) in June (Fischer et al., 2003), and from $-0.8$ to $-0.4$ ppb d$^{-1}$ for Mauna Loa (3.4 km a.s.l.) for four seasons (Cantrell et al., 1996). Forward trajectory analysis suggests that the air mass leaving the top of Mount Tai stays 22 h (as median) longer within the boundary layer over CEC (31–40° N, 114–121° E, 0–2000 m altitude), producing a large amount of ozone in this region in the daytime.

Figure 4 shows the contribution from each chemical process to the $F(O_3)$ and $D(O_3)$ terms as an average over the campaign period. The reaction of HO$_2$ with NO dominated the $F(O_3)$ term, followed by the reactions of MO$_2$ (CH$_3$O$_2$), ACO$_3$ (CH$_3$COO$_2$ and higher acyl peroxy radicals), and ISOP (isoprene peroxy radicals) with NO. The dominant term in $D(O_3)$ was the O($^1D$)+H$_2$O reaction.

4.3 Estimation of regimes

We performed sensitivity model runs, where NO$_x$ concentrations were artificially multiplied by factors of 0.5, 1.1, 2, or 4 while NMHCs concentrations were kept unchanged from the base run, and runs where NMHCs concentrations were conversely multiplied by factors of 0.5, 1.1, or 2 while NO$_x$ concentrations were unchanged. Together with the changes in NMHCs concentrations, CO and CH$_4$ concentrations were also scaled by the same factors but with consideration of their background concentrations. The background concentration of ethane was also taken into account. The results of the averaged diurnal variations in $F-D(O_3)$ from the sensitivity runs are summarized in Fig. 5a. The analysis suggested that $F-D(O_3)$ was much more sensitive to NO$_x$ concentrations than to NMHCs, clearly indicating that ozone production was NO$_x$-limited. The analysis also implied that the ozone production should have been more efficient when the air mass was fresh and the NO$_x$/NO$_y$ ratios (and thus the NO$_x$ concentration for a given NO$_x$ concentration for that air mass) was higher than that observed on the mountain ($\approx 0.18$ as
an average over the campaign period). Additionally, the importance of the NO\textsubscript{x}-limited regime indicates that the ozone pollution will be severer in the future owing to stronger NO\textsubscript{x} emission anticipated in CEC (Yamaji et al., 2008; Ohara et al., 2007). Our result suggesting a NO\textsubscript{x}-limited condition in June for this region is consistent with that of Martin et al. (2004) based on satellite observations. Because of the dominance of the NO\textsubscript{x}-limited condition, the uncertainty in the NMHC concentrations of a factor of 2 propagated to the uncertainty in the daily production of ozone (on the basis of $F - D(O_3)$) of a factor of only 1.15 (Fig. 5a).

We found that the morning of 10 June was an exceptional period, when $F - D(O_3)$ was sensitive to NMHC concentrations rather than to the NO\textsubscript{x} concentration (Fig. 5b). We calculated 10-day kinematic backward trajectories using the CGER-METEX program (http://db.cger.nies.go.jp/metex/) for 6-h intervals during the campaign period, using the National Centers for Environmental Prediction meteorological reanalysis field as input. The starting altitude of the trajectories was 1500 m. The analysis showed that the air mass on 10 June quickly intruded from the north to the Mt. Tai region and passed over Jinan (the capital of Shandong Province, with a population of 3 million and located ∼60 km north of the observatory) only 1 or 2 h before the arrival of the air mass at the observatory (Fig. 6). Consistently, the NO\textsubscript{x}/NO\textsubscript{y} ratio during the period (Fig. 7a) was exceptionally high, suggesting that fresh air pollution affected the observed air mass. The analysis suggests the possibility that the ozone production regime in the city tended to be VOC-limited and changed to being NO\textsubscript{x}-limited during the travel of the air mass, with conversion of NO\textsubscript{x} to NO\textsubscript{y} (HNO\textsubscript{3} and PANs) occurring more rapidly than the oxidation of NMHCs did. There were other periods on 8, 9, 11, 15, 23, and 24 June when ozone was produced under VOC-limited conditions, as marked by open squares in Fig. 7a. Here our definition of VOC-limited is that multiplying a factor of 1.1 to NMHC concentrations results in higher $F - D(O_3)$ values than multiplying the same factor to the NO\textsubscript{x} concentrations does. VOC-limited conditions mostly occurred during the morning and were interestingly associated with high NO\textsubscript{x}/NO\textsubscript{y} ratios in all cases, similar to the case for the morning of 10 June. It should be noted that the observed NO\textsubscript{x} concentrations used in this analysis were neither the input nor the output of the model calculations.

The VOC-limited periods were associated with lower NMHCs/NO\textsubscript{x} ratios (Fig. 7b), as past studies have found. From Fig. 7b, it is deduced that transition from the NO\textsubscript{x}-limited regime to the VOC-limited regime occurred at an NMHC/NO\textsubscript{x} ratio of ∼10 ppbC/ppb. Although not shown, if NMHC + oxygenated VOC concentrations are used for the vertical axis instead of the NMHC concentration, the (NMHC + oxygenated VOC)/NO\textsubscript{x} ratio of ∼25 ppbC/ppb corresponded to the transition. The critical NMHCs/NO\textsubscript{x} ratio falls into the range suggested by Sillman (1999) (10–20 ppbC/ppb) for summer at mid-latitude and is also roughly in agreement with the value for summertime Tokyo at ground
level (around 20 ppb/ppb) determined using the same tropospheric chemistry mechanism (Kanaya et al., 2008). We also conducted similar analysis using the sum reactivity of CO, CH₄, H₂, SO₂, NMHCs, and oxygenated VOCs toward OH (which propagate radical chain reactions) instead of using the concentrations. The results (Fig. 7c) suggest that the transition between the two regimes occurred around 2 s⁻¹ ppb⁻¹. This is slightly higher than 1 s⁻¹ ppb⁻¹ suggested by Kleinman et al. (2005) for the boundary layer over cities in the United States. The fraction of the radical + NOₓ reaction rates (L_N) to the total radical termination rate (L(radical)) was greater than 0.5 for the VOC-limited data points (Fig. 7d). In this case, the OH+NO₂+M and XO₂ (peroxy radicals that account for additional NO to NO₂ conversion)+NO reactions contributed largely to L_N. This is qualitatively consistent with the results of Kleinman et al. (2005), in which the contribution from the NOₓ-related processes to the radical termination increased under VOC-limited conditions.

Although the ozone production rates and regimes determined in this study are based on the point measurement at the top of Mt. Tai, they would be representative for a wider background region without strong influence of fresh pollution, assuming the concentrations of ozone precursors are relatively uniform and our data are representative. Considering that more than half of the 10-day-long backward trajectories arriving at the top of Mt. Tai for June 2006 did not travel over any Chinese large cities (91 cities with populations more than 500,000), the regional production of ozone under the NOₓ-limited condition and its accumulation over multiple days can be one of the major causes for the high O₃ levels at a regional scale. See Li et al. (2008a) for detailed analysis based on a 3-dimensional model that combines the regional-wide ozone production and vertical and horizontal transport for this campaign period.

4.4 Other sensitivity studies

We made additional sensitivity runs where (1) the CO concentrations were constrained to 90 ppb for all time, (2) the ALD concentrations were scaled down by a factor of 3, (3) concentrations of biogenic hydrocarbons (isoprene and monoterpenes) were set to zero, and (4) the heterogeneous loss of HO₂ radicals on aerosol surfaces was taken into account.

4.4.1 CO

One of the significant features of atmospheric composition over China is that the CO concentration is very high (the monthly mean was 560 ppb on average during the campaign). Therefore, we examined the role of CO in the photochemistry. Constraining the CO concentration to 90 ppb (minimum concentration observed during the campaign) for all time resulted in a daily production of ozone of 58 ppb, which was the same as the value for the base run (58 ppb). This run suggests that the other hydrocarbons were present at sufficiently high concentrations to cycle OH to RO₂ (HO₂), and thus CO itself was not necessarily important. This analysis implies that the reduction in CO emission does not readily result in a reduction in ozone concentrations if not accompanied by a reduction in emission of other precursors.

4.4.2 ALD

PTR-MS showed that the CH₃CHO concentrations were normally higher than the HCHO concentrations during the campaign period (1.8 and 3.2 ppb respectively for HCHO and CH₃CHO concentrations as midday 6-h averages (09:00–15:00 CST)). This magnitude relationship has rarely been reported, and thus it could be another feature specific to the atmospheric composition over China. However, we cannot dismiss the possibility that the ALD signal at m/z 45 in the PTR-MS was affected by other compounds. Thus, we conducted an additional sensitivity run with the ALD concentration reduced by a factor of 3. The daily production of ozone reduced to 52 ppb in this run from 58 ppb in the base run. This reduction would be because 1) the initial radical production rate from the photolysis of ALD was reduced, 2) the OH+ALD reaction, forming a relatively efficient catalytic cycle of ozone production, slowed, and because 3) PAN formation decreased, making ozone formation inefficient.

4.4.3 Biogenic hydrocarbons

The average isoprene and Σ-monoterpene concentrations observed at midday are 0.28 and 0.19 ppb. The observed air mass might have been affected by the emission of biogenic hydrocarbon from nearby trees or the local biosphere on the mountain slopes, over which the air masses have traveled as the daytime upslope wind. The CO₂ concentrations measured at the site has a regular daytime drop (nighttime average (21:00–03:00 CST) – daytime average (09:00–15:00 CST)=3.8 ppb (Komazaki et al., unpublished data, 2006)), suggesting the possibility that the air mass is influenced by the vegetation. Thus we made an additional sensitivity run in which isoprene and monoterpane concentrations were set to zero to calculate the representative ozone production rate for the altitude level (1.5 km) without influence from vegetation. In this case, the daily production of ozone decreased to 49 ppb from the 58 ppb of the base run. The analysis suggests that biogenic hydrocarbons make some but limited contribution to ozone production. Thus, we conclude that the determined ozone production rate was mostly representative for the altitude in the middle of CEC, even when the biogenic hydrocarbon concentrations had some variability.
4.4.4 Heterogeneous loss of HO\(_2\)

Although the heterogeneous loss of HO\(_2\) on aerosol surfaces has been suggested in recent studies (e.g., Morita et al., 2004; Taketani et al., 2008), it is rarely incorporated in a model exploring ozone production. The loss rate of HO\(_2\) (\(k_{v}\)) was calculated using the following Fuchs–Sutugin equation and incorporated in our box model.

\[
k_{v} = \int \frac{4 \pi D r}{1 + K_n \left( \frac{1.333 + 0.71 K_n^{-1}}{1 + K_n^{-1}} + \frac{4(1 - \gamma)}{3 \gamma} \right)} N d r, \tag{4}
\]

where \(K_n\) is the Knudsen number, \(r\) is the radius of the aerosol, and \(N\) is the number density of the particles. The gas phase diffusion coefficient \(D\) of HO\(_2\) was assumed to be 0.247 cm\(^2\) s\(^{-1}\) (Mozurkewich et al., 1987). The uptake coefficient \(\gamma\) employed in this run was the central value in the range of our laboratory measurements, 0.25±0.09 (\(n=10\)), which were determined using aerosol flow tube-chemical conversion-laser-induced fluorescence (Taketani et al., 2008). In the experiment, aerosol particles were generated in an aerosol flow tube in the laboratory by atomizing the water extract of total suspended particles collected on quartz filters (10 samples) using a high-volume sampler during the field campaign. The relative humidity during the laboratory experiment was 75±2%, close to the campaign average (68%). Details of the experiments will be described elsewhere (Taketani et al., 2009).

The particle size distribution was not observed during the campaign and thus it was roughly estimated from the summed observed mass concentrations of major chemical species. In detail, the estimation was made separately for 9 size bins segregated at 0.43, 0.65, 1.1, 2.1, 3.3, 4.7, 5.8, and 9.0 \(\mu\)m. First, the total mass concentration for each bin, as estimated by adding mass concentrations of water-soluble ions, organics, elemental carbon, and water (assumed as 20\% of dry mass), was converted to volume density by assuming a constant density of particles (1.4 g cm\(^{-3}\)). In this step, it was assumed that the size distributions of elemental carbon and organic carbon were similar to those of sulfate particles and that the mass concentrations of organic aerosol were 1.6 times the observed OC mass concentration. Next, the volume distribution was converted to a number distribution. For the smallest bin (less than 0.43 \(\mu\)m), a typical size distribution for the “background and aged urban plume case” (accumulation mode diameter (0.12 \(\mu\)m) with \(\sigma_e=1.84\) and coarse mode diameter (0.83 \(\mu\)m) with \(\sigma_e=2.12\)) reported by Whitby and Sverdrup (1980) was assumed for converting the volume to a number density distribution. For the other size bins, diameters representative of the bins (0.5, 0.8, 1.5, 3.0, 4.0, 5.0, 6.0, and 10 \(\mu\)m) were roughly estimated and used to convert volume to number density. The median number density of particles integrated over the full size range was 4.4\(\times\)10\(^3\) cm\(^{-3}\) for 30 filter samples covering the whole campaign period. The typical surface area concentration was 6.3\(\times\)10\(^5\) \(\mu\)m\(^2\) cm\(^{-3}\).

The median \(k_{v}\) value calculated from Eq. (4) using the number density size distributions was 0.014 s\(^{-1}\). Considering that the HO\(_2\) loss rate due to self reaction was 8.0\(\times\)10\(^{-3}\) s\(^{-1}\), the heterogeneous loss had a large influence on the HO\(_2\) concentration and hence on ozone production.

With this heterogeneous loss of HO\(_2\), the daytime maximum HO\(_2\) concentration averaged over the campaign period reduced from 34 pptv to 20 pptv (Fig. 8). The OH and RO\(_2\) maximum concentrations also indirectly reduced from 5.0 to 3.7\(\times\)10\(^6\) cm\(^{-3}\) and from 43 to 39 pptv, respectively. In general, the impact of heterogeneous loss would be large when the \(OH+NO_2+M\) reaction does not play a significant role in the radical termination reaction. The upper part of the boundary layer in CEC, where the number density of aerosol particles is still high and the role of the \(OH+NO_2+M\) reaction is limited (Table 2), is one of the regions where the impact of the heterogeneous loss is large.
The daily production of ozone decreased to 39 ppb from 58 ppb with the inclusion of the heterogeneous loss of HO₂. The daytime 6-h average of the net ozone production rate decreased to 4.3 ppb h⁻¹ from 6.4 ppb h⁻¹ in the base run. However, these values are still large enough to explain the observed buildup of ozone concentration.

Additional model sensitivity runs with the NOₓ or NMHC concentrations multiplied by a factor of 1.1 in the presence of heterogeneous loss suggested that the main conclusion of this paper (that production is normally NOₓ-limited) remains unchanged; that is, inclusion of the heterogeneous loss of HO₂ made the situation more NOₓ-limited because more NOₓ is needed to increase the radical termination rate of the OH+NO₂+M reaction to a level that can compete with the rate of heterogeneous loss of HO₂.

It should be noted that this sensitivity analysis is rough because the size distribution of particles was only estimated and because it is assumed that the effect of the ambient aerosol particles is similar to that of the regenerated particles in the laboratory. However, this is the first time to indicate that the heterogeneous loss of HO₂ can be very important to the ozone production rate from the kinetic experiments using aerosol particles collected in the ambient atmosphere.

5 Summary

The net photochemical production rate of ozone at the top of Mount Tai, China, was estimated using a photochemical box model on the basis of a tropospheric chemistry mechanism constrained by simultaneous measurements of ozone and its precursors. It is estimated that 58±37 ppb of ozone can be produced in one day, suggesting that the local photochemistry is active and an important factor controlling ozone concentration. Normally, NOₓ-limited ozone production is suggested from sensitivity model runs. Considering that the NOₓ emission is anticipated to increase in CEC at least until 2020 (Ohara et al., 2007), the regional ozone production will become severer in the future. Sensitivity runs indicated that reductions in the concentrations of CO (to 90 ppb), CH₃CHO (by a factor of 3), and biogenic hydrocarbons (to zero) resulted in 0%, 10%, and 16% reductions in the daily production of ozone. Heterogeneous loss of HO₂ is potentially important in that it possibly reduces production by 33%. However, the conclusion of there being a NOₓ-limited condition did not change with the inclusion of heterogeneous loss.

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