Prediction of Ozone Concentrations over the Sea of Japan Coastal Area Using WRF/Chem Model

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Abstract: The fully coupled WRF/Chem (Weather Research and Forecasting/Chemistry) model is used to simulate air quality over the Sea of Japan coastal area. Anthropogenic surface emissions database used as input for this model are mainly based on Global hourly emissions data (dust, sea salt, biomass burning), RETRO (REAnalysis of the TROpospheric chemical composition), GEIA (Global Emissions Inventory Activity) and POET (Precursors of ozone and their Effects in the Troposphere). Climatologic concentrations of particulate matters derived from Regional acid Deposition Model (RADM2) chemical mechanism and Secondary Organic Aerosol Model (MADE/SORGAM) with aqueous reaction were used to deduce the corresponding aerosols fluxes for input to the WRF/Chem. The model was firstly integrated for 48 hours continuously starting from 00:00 UTC of 14 March 2008 to evaluate ozone concentrations and other precursor pollutants were analyzed. WPS meteorological data were used for the simulation of WRF/Chem model in this study. Despite the low resolution of the area global emissions and the weak density of the local point emissions, it has been found that WRF/Chem simulates quite well with the diurnal variation of the chemical species concentrations over the Sea of Japan coastal area. The simulations conducted in this study showed that due to the geographical and climatologically characteristics, it is still environmentally friendly by the transported pollutants in this region.

Key words: Air quality; Modeling; WRF/Chem; Ozone concentrations

INTRODUCTION

At the present, more than 60 million people are added to cities each year worldwide, about 325 cities have a population of more than one million, compared to 270 in 1990 [1]. In recent decades, air quality has deteriorated remarkably in the large cities of the developing countries. The export of air pollutants from urban to regional and global environments is of major concern because of wide-ranging potential consequences for human health and for cultivated and natural ecosystems, visibility degradation, weather modification, radiative forcing and changes in tropospheric oxidation (self-cleaning) capacity. Rapidly increasing urbanization will be a major environmental driving force in the 21st century, affecting air quality on all scales-local, regional and global [1]. Large numbers of people are likely to be exposed to unhealthy ozone concentrations when ground-level ozone accumulates in urban metropolitan areas under certain weather conditions [2] and quantitative atmospheric dispersion models will be of great help to provide effective decision support systems for planners and administrators.

The main pollutants emitted into the atmosphere in urban area are sulfur oxides (SO\textsubscript{x}), nitrogen oxides (NO\textsubscript{x}), carbon monoxide (CO), volatile organic compounds (VOCs), metal oxides and particular matter (PM/aerosols) mostly consisting of black carbon, sulfates, nitrates and organic matter. The oxidizations of CO, VOCs and NO\textsubscript{x} produce O\textsubscript{3} in the PBL (Planetary Boundary Layer) which has important impact on human’s health in urban area.

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The recent economical developments of Korea and Japan are very rapid due to rapid growth of heavy industrial operations. The rapid growing urbanization will cause wide-ranging potential consequences for environmental problems. These areas have been suffering server air pollution problems, such as high particular matter (PM) concentrations and poor visibility [3-6]. As the increase in industrial activity and number of automobiles, the emission of VOCs (volatile organic carbons) and NOx (NO + NO2) will be significantly increased. Both VOCs and NOx play critical roles in O3 formation in the troposphere [7, 8] and the variations in their concentrations and the ensuing effects on O3 production rate can be characterized as either NOx-sensitive or VOC-sensitive [7, 9-12]. Thus, better understanding the relationship between O3 precursors (VOCs, NOx) and O3 formation in East Asia is one of the critical pre-required information to develop effective O3 control strategies [13-15].

In Japan, VOC, NOx and SOx categorized as a hazardous air pollutant (HAP) in 1996 and are on the list of Substances Requiring Priority Action published by the Central Environmental Council of Japan [16]. The Central Environmental Council made the second report on “Future direction of measures against hazardous air pollutants” in Oct. 1996. The second report also proposed that the voluntary action to reduce emission, as well as investigation of hazardousness, atmospheric concentrations and pollution source of those substances should be promoted. Although the industrial emission of VOC, NOx and SOx in Japan have been decreasing in recent years, primarily due to voluntary reductions from industrial sources, the risks of exposure to this pollutant have remained largely unknown. In this study, a rapid change of O3 and its precursors are simulated during the spring and summer time in 2008. The simulation result leads to better understanding O3 variability in the Sea of Japan coastal area [17]. A three dimensional chemical/dynamical regional model (WRF/Chem V-3.3) is applied to analyze the causes of the rapid changes of O3 during this period.

Ozone (O3) is not emitted directly into the air; instead it forms in the atmosphere as a result of a series of complex chemical reactions between oxides of nitrogen (NOx) and hydrocarbons, which together are precursors of ozone. Ozone precursors have both anthropogenic (man-made) and biogenic (natural) origins. Motor vehicle exhaust, industrial emissions, gasoline vapors and chemical solvents are some of the major sources of NOx and hydrocarbons. Many species of vegetation including trees and plants emit hydrocarbons and fertilized soils release NOx.

In the presence of ultraviolet radiation (hv), oxygen (O2) and nitrogen dioxide (NO2) react in the atmosphere to form ozone and nitric oxide (NO) [18, 19]. The resultant ozone reacts with NO to form nitrogen dioxide. A steady state is attained through these reactions:

\[
\text{NO}_2 + hv \rightarrow \text{NO} + \text{O} \\
\text{NO} + \text{NO}_2 + \text{H}_2\text{O} \rightarrow 2\text{HONO} \\
\text{HONO} + h \rightarrow \text{NO} + \text{OH} \\
\text{O} + \text{O}_2 \rightarrow \text{O}_3 \\
\text{CO} + 2\text{O}_2 \rightarrow \text{CO}_2 + \text{O}_3 \\
\text{O}_3 + \text{NO} \rightarrow \text{NO}_2 + \text{O}_2
\]

Even without anthropogenic emissions, these reactions normally result in a natural background ozone concentrations of 25 to 45 ppb [20, 21]. Ozone cannot accumulate further unless volatile organic compounds (VOCs), which include hydrocarbons, are present to consume or convert NO back to NO2 [19, 22, 23] as:

\[
\text{VOC} + \text{NO} \rightarrow \text{NO}_2 + \text{other products}
\]

This equation is a simplified version of many complex chemical reactions. As NO is consumed by this process, it is no longer available to react with ozone. When additional VOCs are added to the atmosphere, a greater proportion of the NO is oxidized to NO2, resulting in greater ozone formation. Anthropogenic sources of NO lead to higher levels of NO2 in the atmosphere, which will be available for photolysis. The formation and increase in ozone concentrations occur over a period of a few hours. Shortly after sunrise, NO and VOCs react in sunlight to form ozone. Throughout the morning, ozone concentrations increase while NO and VOCs are depleted. Eventually, either the lack of sunlight, NO, or VOCs limit the production of ozone. This diurnal cycle varies greatly depending on site location, emission sources and weather conditions. Precursor emissions of NO and VOC are necessary for ozone to form in the troposphere. Understanding the nature of when and where ozone precursors originate may help forecasters to factor day-to-day emissions changes. For example, if a region’s emissions are dominated by mobile sources, emissions and hence the ozone that forms may depend on the day-of-week commute patterns. The dominant NOx producers are combustion processes, including industrial and electrical generation processes and mobile sources such as automobiles. Mobile sources also account for a large portion of VOC emissions. Industries such as the chemical industry or others that use solvents also account for a large portion of VOC emissions. Biogenic VOC emissions
The WRF model is a mesoscale numerical weather prediction system designed to serve both operational forecasting and atmospheric research needs. The effort to develop WRF has been a collaborative partnership, principally among the National Center for Atmospheric Research (NCAR), the National Oceanic and Atmospheric Administration (NOAA), the National Center for Environmental Prediction (NCEP), the Forecast Systems Laboratory (FSL), the Air Force Weather Agency (AFWA), the Naval Research Laboratory, Oklahoma University and the Federal Aviation Administration (FAA). The WRF model is a fully compressible and Euler nonhydrostatic model. It calculates winds (u, v and w), perturbation potential temperature, perturbation geopotential and perturbation surface pressure of dry air. It also can optionally output other variables, including turbulent kinetic energy, water vapor mixing ratio, rain/snow mixing ratio and cloud water/ice mixing ratio. The model physics include bulk schemes, mixed-phase physics for cloud-resolving modeling, multi-layer land surface models ranging from a simple thermal model to full vegetation and soil moisture models, including snow cover and sea ice, turbulent kinetic energy prediction or non-local K schemes for planetary boundary layer calculation and longwave and shortwave schemes with multiple spectral bands and a simple shortwave scheme. A detailed description of the WRF model can be found on the WRF web-site http://www.wrf-model.org/index.php.

In addition to a dynamical calculation, a chemical model (WRF/Chem) is fully (on-line) coupled with the WRF model. A detailed description of WRF/Chem is given by Grell et al. [26]. The version of the model (Version-3.3), as used in the present study, includes simultaneous calculation of dynamical parameters (winds, temperature, boundary layer, clouds, etc.), transport (advection, convective and diffusive), dry deposition [27], gas phase chemistry, radiation and photolysis rates [28, 29] and surface emissions, including online calculation of biogenic emission. Ozone chemistry is represented in the model by a modified Regional Acid Deposition Model, version 2 (RADM2) gas phase chemical mechanism [30] which includes 158 reactions among 36 species with Secondary Organic Aerosol Model (MADE/SORGAM) of aqueous reaction [31].

**Model Setting:**

A detailed description of WRF/Chem is given by Grell et al. [26]. The version of the model (Version-3.3), as used in the present study, includes simultaneous calculation of dynamical parameters (winds, temperature, boundary layer, clouds, etc.), transport (advection, convective and diffusive), dry deposition [27], gas phase chemistry, radiation and photolysis rates [28, 29] and surface emissions, including online calculation of biogenic emission. Ozone chemistry is represented in the model by a modified Regional Acid Deposition Model, version 2 (RADM2) gas phase chemical mechanism [30] which includes 158 reactions among 36 species with Secondary Organic Aerosol Model (MADE/SORGAM) of aqueous reaction [31].

**Method of Simulation**

**Model Setting:** The Sea of Japan coastal area considered in the present study were located (latitude 31.4 - 45.0 N and longitude 129.3 - 146.2 E), which is a marginal sea of the western Pacific Ocean, bordered by Japan, Korean peninsula and Russia and is referred to in South Korea as the East Sea [32, 33].

In this study, the model is used to simulate medium scale and regional circulations influenced by the complex terrain within and around the Sea of Japan coastal area during the period 00:00 UTC (Coordinated Universal Time) [34] of 14 March 2008 to 00.00 of 16 March 2008, whereas Japan Standard Time (JST) [35] is the standard time zone of Japan and is 9 hours ahead of UTC. One nesting domains was defined using Lambert projection, which can be seen in Figure 1. The domain covers the Sea of Japan coastal area with the center point of 38.5 N, 137.60 ºE, horizontal grids of 75 × 70 and grid spacing of 22 km. The domain settings and configuration options are shown in Table 1. The initial meteorological fields and boundary conditions are from NCEP global reanalysis data with a horizontal resolution of 0.5º by 0.5º and 35 vertical levels, non-uniformly spaced, from the surface to 20 mbar. The boundary conditions were forced every 6 hours. The ideal concentrations profiles [36] were used as initial and boundary conditions of chemical species.

**Emissions Data Acquisition and Pre-processing:** The linkages in the formation and transport of pollution between the local, regional and global scales are receiving increasing recognition and a variety of atmospheric chemistry models address these aspects. This puts new demand on the emission inventories that are used as input to these models. They need to be accurate at the local scale yet cover the entire globe with a consistent approach.

The GEIA (Global Emissions Inventory Activity) / ACCENT (European Network of Excellence on Atmospheric Composition Change) data portal is a cooperative effort providing surface emissions data
Table 1: WRF/Chem domain setting and configuration options

<table>
<thead>
<tr>
<th>Option</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Horizontal grid (x,y)</td>
<td>75, 70</td>
</tr>
<tr>
<td>Grid spacing</td>
<td>22 km</td>
</tr>
<tr>
<td>Meteorological time step</td>
<td>180 s</td>
</tr>
<tr>
<td>Chemical time step</td>
<td>180 s</td>
</tr>
<tr>
<td>Microphysics</td>
<td>WSM3-class simple ice scheme [59]</td>
</tr>
<tr>
<td>Advection scheme</td>
<td>5th horizontal/3rd vertical [60, 61]</td>
</tr>
<tr>
<td>Long-wave radiation</td>
<td>RRTM [62]</td>
</tr>
<tr>
<td>Short-wave radiation</td>
<td>GODDARD [63, 64]</td>
</tr>
<tr>
<td>Surface layer</td>
<td>Moni-Obukhov (Janic Eta) [65]</td>
</tr>
<tr>
<td>Land-surface model</td>
<td>NOAH [66]</td>
</tr>
<tr>
<td>Boundary layer</td>
<td>Mellor-Yamada-Janic TKE [67, 68]</td>
</tr>
<tr>
<td>Cumulus parameterization</td>
<td>Grell-Devenyi ensemble scheme [69]</td>
</tr>
<tr>
<td>Chemistry option</td>
<td>RADM2 [70]</td>
</tr>
<tr>
<td>Dry deposition</td>
<td>Wesley, 1989 [71]</td>
</tr>
<tr>
<td>Biogenic emissions</td>
<td>Guenther scheme [72, 73]</td>
</tr>
<tr>
<td>Photolysis option</td>
<td>Madronich, 1987 [74]</td>
</tr>
<tr>
<td>Aerosol option</td>
<td>MADE/SORGAM [75]</td>
</tr>
</tbody>
</table>

Table 2: Global inventories offered by the GEIA/ACCENT data portal

<table>
<thead>
<tr>
<th>Inventory</th>
<th>Categories</th>
<th>Spatial resolution</th>
<th>Temporal resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>POET [76, 77]</td>
<td>Anthropogenic biomass burning (natural)</td>
<td>1° x 1°</td>
<td>Annual (anthro.) monthly (biom. burn.) monthly (natural)</td>
</tr>
<tr>
<td>RETRO [78]</td>
<td>Anthropogenic biomass burning</td>
<td>0.5° x 0.5°</td>
<td>Monthly</td>
</tr>
<tr>
<td>EDGAR</td>
<td>Anthropogenic biomass burning</td>
<td>1° x 1°</td>
<td>Annual</td>
</tr>
<tr>
<td>GFED v2 [82, 83]</td>
<td>Biomass burning</td>
<td>1° x 1°</td>
<td>Monthly 8 day (available:<a href="http://daac.ornl.gov/VEGETATION/">http://daac.ornl.gov/VEGETATION/</a> guides/global_fire_emissions_v2.1.html )</td>
</tr>
<tr>
<td>CO₂ [84]</td>
<td>Anthropogenic</td>
<td>1° x 1°</td>
<td>Annual</td>
</tr>
<tr>
<td>GEIA v.1 [80]</td>
<td>Anthropogenic biomass burning natural</td>
<td>1° x 1°</td>
<td>Annual + monthly for NOx, SO₂ and nat.VOC</td>
</tr>
</tbody>
</table>

Observation of Ozone Concentrations over Japan: Annual and monthly pollutant concentrations observed at monitoring stations operated by Japanese, local governments and Japan Meteorological Agency [37, 38] (JMA) are available on the website [39]. It contains the monitoring data since 1970, which are valuable to evaluate the long-term or short-term trends of pollutant concentrations in Japan. The increasing trend of surface ozone in Japan has been reported annually by the Japanese government based on this database [40]. Observation data at 1045 monitoring stations which continued monitoring for photo-chemical oxidants during 1996-2005 was notated in this study to know the scenario of ozone concentrations in March and to compare with simulation result. Most of the monitoring stations are located in coastal populated areas in Japan as shown in the left map of Figure 1 because they are operated mainly to evaluate the attainment of environmental quality standard (EQSs). This database has compiled surface ozone concentrations as a daytime average and maximum concentrations of photo-chemical oxidants. Daytime corresponds to thirteen hours from 05:00 a.m. to 06:00 p.m. Photo-chemical oxidants include other trace oxidants like H₂O₂ and PAN, but instruments which detect only ozone...
have been officially approved by the Japanese government because differences in concentrations of photochemical oxidants and ozone are expected to be small [41]. Therefore, differences in concentrations between ozone and photochemical oxidants can be ignored for monthly variation of observed concentrations of surface ozone. Additionally, the Acid Deposition Monitoring Network in East Asia (EANET) operates monitoring stations located mainly in remote areas. Observed concentrations of surface ozone at ten EANET monitoring stations in Japan were used to validate the performance of the simulation in Japanese background areas during 2000-2005. The locations of EANET monitoring stations are shown in the middle map of Figure 2. They are scattered in remote areas throughout Japan.

The concentrations of surface ozone at EANET monitoring stations in Japan shows a seasonal variation with low values in summer (August) and higher value in Spring (April). The average ozone concentrations in March in different EANET monitoring stations were found about 37 ppb to 55 ppb and about 33.5 ppb over Japanese populated areas. Figure 2 shows monthly variations of concentrations of observed surface ozone, averaged during 2000-2005 at each of ten EANET monitoring stations. The red line in Figure 2 shows monthly variations of observed concentrations of surface ozone averaged during 1996-2005 over Japanese populated areas.

Modeling Results and Discussions

Ozone Concentration by WRF/Chem Simulation: Figure 3, which is the composite image from the modeling results of single nesting domains, illustrates the spatial distribution of surface O₃ concentrations averaged from 00:00 UTC of 14 March (9:00 a.m. of Japanese Standard time) to 00:00 UTC of 16 March of 2008. In order to display the time evolution of the simulated ozone concentrations, Figure 4 shows the structure of the ozone field from 03:00 UTC of March 14 to 00:00 UTC of 16 March of 2008 at 100 meter height above the surface level. This level was chosen in order to emphasize the transport processes across the Sea of Japan coastal area relative to surface processes. The two days simulation shows that Sea of Japan has low to moderate ozone concentrations. The high O₃ concentrations zone was calculated mainly over the Sea of Japan ocean regions during 14 March, whereas the ozone concentrations was calculated 40 to 45 ppb in the Figure 4. At 09:00 UTC of March 14, ozone starts building up in the boundary layer with concentrations of 44 ppb near the Sea of Korean peninsula. An anticyclone [42] moved the high-ozone region rotates anticlockwise from the Sea of Korean peninsula (09:00 UTC of March 14) to the southwest part of the Honshu island of Japan (15:00 UTC of March 14) and then Pacific ocean (06:00 UTC of March 15). Due to the longer life time of O₃ (about a week) in the troposphere, the distribution of O₃ concentrations are greatly influenced by airflow of the anticyclone. Figure 5 shows the clockwise movement of airflow in an anticyclonic rotation at latitude 41.6N, longitude 138.4E at 15:00 UTC of March 14, which moved to the east at 00:00 UTC of March 15 at latitude 42N, longitude 140E. After 00:00 UTC of March 15, maximum ozone concentrations (50 ppb) were found in the south part of the Honshu island of Japan. Another small area of high ozone concentrations built up near the coastal area of the east part of the Honshu island of Japan (00:00 UTC of March 15).

![Fig. 1: Location maps of monitoring stations in populated area, and EANET monitoring stations in Japan which were used in this study [41].](image-url)
Fig. 2: Monthly variations of observed concentrations of surface ozone, averaged during 2000-2005 at each of ten EANET monitoring stations. The red line shows monthly variations of observed concentrations of surface ozone averaged during 1996-2005 over Japanese populated areas.

Fig. 3: Nesting domain setting of the model.

14) which stagnated for some hours, then moved to the Pacific ocean, the whole polluted air mass is pushed away eastward by a cold front coming from the northwest [43]. In addition, the level of primary pollutants, such as NOX, were relatively lower over the ocean, resulting in the less consumption of O3, thus the high concentrations of O3 are found in ocean. An important fact revealed by the simulated result, ozone levels did not reach extreme values (70-90 ppb) [44].

Comparison of Simulation Result with Eanet Monitoring

Data: Table 3 shows the accuracy of WRF/Chem simulation result comparing with EANET monitoring data. Though the finer resolution in WRF/Chem effectively titrates ambient ozone and moved the simulated concentrations of surface ozone closer to the observed value, WRF/Chem underestimates surface ozone concentrations in March over the most of the EANET monitoring stations in Japan. Besides WRF/Chem overestimates the surface ozone concentrations over the Populated area in Japan. Standard deviation was calculated about 6 in EANET monitoring stations comparing with WRF/Chem simulation result, which shows a reasonable agreement with the monitoring data with respect to predicting localized atmospheric ozone concentrations. Absolute values of ozone concentrations are comparable but seasonal variations are different between observed and simulated surface ozone. Therefore, it is complicated to compare the WRF/Chem simulation result of ozone concentration in March with observation data. Concentrations of observed surface ozone are the highest in spring and the lowest in summer at all stations. Second peaks in autumn were found at most of stations. In March the range of observed average ozone concentrations were found about 37.5 ppb to 55 ppb, whereas the average ozone concentrations calculated by WRF/Chem was 40 ppb to 48 ppb in EANET monitoring stations.
Fig. 4: Ozone concentrations (ppm) over the Sea of Japan coastal area from 00:03 UTC of March 14 to 00:00 UTC of March 16 of the year 2008, which is the simulation result of WRF/Chem model.
Fig. 5: Temperature distribution (°C) in 15:00 UTC of March 14 and 00:00 UTC of March 15.

Fig. 6: Vertical distribution of ozone in east west cross-section at fixed latitude 35.7 at 21.00 UTC of March 14 and 15.00 UTC of March 15.

Table 3: Comparison study of average zone concentrations in March between EANET monitoring data and WRF/Chem simulation result.

<table>
<thead>
<tr>
<th>EANET monitoring stations</th>
<th>EANET monitoring data (ppb)</th>
<th>WRF/Chem simulation result (ppb)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rishiri</td>
<td>50</td>
<td>42</td>
</tr>
<tr>
<td>Tappi</td>
<td>55</td>
<td>44</td>
</tr>
<tr>
<td>Ogasawara</td>
<td>46</td>
<td>40</td>
</tr>
<tr>
<td>Sado</td>
<td>50</td>
<td>48</td>
</tr>
<tr>
<td>Hppo</td>
<td>54</td>
<td>46</td>
</tr>
<tr>
<td>Ijirako</td>
<td>37.5</td>
<td>44</td>
</tr>
<tr>
<td>Oki</td>
<td>47.5</td>
<td>43</td>
</tr>
<tr>
<td>Banryuko</td>
<td>44</td>
<td>45</td>
</tr>
<tr>
<td>Yasuhara</td>
<td>42.5</td>
<td>43</td>
</tr>
<tr>
<td>Hedomisaki</td>
<td>50</td>
<td>40</td>
</tr>
<tr>
<td>Populated area</td>
<td>33.5</td>
<td>39</td>
</tr>
</tbody>
</table>

Local Meteorological Characteristics: Large-scale winds give the general wind regime and local winds influence the convergence zones. As such, regional circulations play an important role in the dispersion of pollutants [45]. Many studies have showed that wind and weather conditions are one of the most important factors for formation of photochemical pollution. These factors include stable boundary layer, strong solar radiation, clear sky, weak wind, high temperature and low humidity, etc. Figure 5 shows the temporal variations of simulated meteorological conditions of the entire domain. Due to the winter season, the air temperatures are lower, with daily maximum temperatures are below 6 °C (Figure 5) on March and the daily minimum relative humidity was below 75% in general, in which the photochemical pollution was weak. Figure 5 (15:00 UTC of March 14) shows the strong wind in the nighttime in the coastal area of Japan Sea. This illustrates that photochemical reaction may play an important role on the studied episode. It should be noted that the wind speed were relatively strong in the north part of Japan and the temperature was lowered by the influence of the anticyclone, but the O₃ peaks were
little higher at the south part of Japan (Figure 4), this high $O_3$ peaks may be mainly attributed to the transport from west to east. Figure 5 shows the magnitude of wind speed and wind direction by WRF simulation. The simulated wind direction shows that Japan was under the control of the westerly or northwesterly winds from the continent [41].

**Vertical Distribution of $O_3$ Concentrations:** This section analysis the vertical distribution of $O_3$ concentrations in the PBL and discuss correlations between concentrations at the ground and elevated levels on the vertical sections at a fixed latitude 35.7 between east west cross-section of enter domain at 21.00 UTC of March 14 (6:00 am of March 15 of Japanese Standard Time) and 15.00 UTC of March 15 (12:00 am of March 16 of Japanese Standard Time) respectively. Figure 6 shows the vertical profiles of mean $O_3$ concentrations in the 1.5 km (350 Pa) layer of the atmosphere, based on the WRF/Chem simulation. The vertical concentrations of $O_3$ were higher in the morning time than that of midnight, while the vertical distribution of tropospheric ozone showed an increase in concentrations with height. It is evident that due to the influence of urbanization in Tokyo metropolitan (latitude 35.7 N, longitude 139.7 E) area the atmospheric boundary layer become quite unstable and thus it is difficult for $O_3$ to be transported upward and diffused, therefore $O_3$ was lower in the ground level.

In general, the surface ozone exhibits strong diurnal variation with a mid-afternoon maximum and an early morning minimum [46, 47]. In this study concentrations of ozone in daytime was much larger than those at nighttime (Figure 6). At nighttime, the presence of temperature inversion isolates the surface level ozone from that of upper levels, as there is little or no vertical mixing between the surface layer and levels above the nocturnal boundary layer (NBL). Surface ozone is partly removed by deposition and reaction with nitric oxide (NO). Since no ozone is produced in the absence of sunlight at night, ozone concentrations begin to decrease after sunset, reaching minimum in early morning before sunrise.

After sunrise, the height of the unstable boundary layer begins to increase as the surface is heated and the nocturnal inversion is destroyed. This results in downward mixing of ozone from aloft [48]. Surface ozone is also locally generated by reactions of nitrogen oxides with volatile organic compounds (VOCs) in the presence of sunlight [22, 49, 50]. In this study, ozone concentrations increase rapidly from early morning to about noon. By noon, the mixing height typically exceeds 1 Km and ozone is well mixed within the mixing layer. Strong photochemical production and strong convection in the mid-afternoon period cause ozone concentrations to reach peak values in the late afternoon. As a result, ozone production decreased with diminishing intensity of sunlight, resulting in a decrease of ozone concentrations with time. At the time of sunset, concentrations decreased substantially below the afternoon maximum value. As a new NBL begun to form in the evening, ozone concentrations near the surface continue to fall due to surface deposition and reaction with NO. Figure 6 shows the typical vertical profile at nighttime was unbalanced caused by dry deposition to the surface and reaction with NO and the typical daytime profile was stable, which reflecting the important roles of both the local production of ozone and greater convective mixing in the vertical.

**Tropospheric Nox Transportation:** Due to anthropogenic activities, the atmospheric trace gas composition has undergone significant changes during the past decades. In particular the tropospheric concentrations of many trace species (e.g. of $O_3$, CO, NO, NO$_2$, CH$_3$) has largely increased. Nitrogen oxides ($NO_x = NO + NO_2$) are key species in atmospheric chemistry and are heavily influenced by anthropogenic emissions. The availability of NOx limits photochemical ozone formation in rural and remote regions [22, 51] and particularly in the upper troposphere [52], where the impact of ozone on radiative forcing is strongest [53]. NOx also contributes to acid deposition from the atmosphere [54]. While the lifetime of NOx in the atmospheric boundary layer (about 1 day) is too short to allow transport over long distances, its lifetime in the upper troposphere is 5 of the order of 5-10 days [55] which is sufficient even for intercontinental transport [56]. However, due to removal processes, transport of reactive nitrogen from the surface, where the largest sources are located, to the upper troposphere is inefficient [57].

In this study, We simulated the evolution of the NOx plume with the atmospheric chemical transportation WRF/Chem model. The simulation results show a plume that traveled from East China (06:00 UTC March 14) to eastward over the East Sea, passed the south part of Tohoku region of Japan (00:00 UTC March 15) and then floated towards the Pacific Ocean (00:09 UTC March 15).
Figure 7 shows the NOx concentrations (g/m³) over the Sea of Japan coastal area. The pink color in the Figure 7 indicates high NOx concentrations (150 µg/m³) which was formed in East China and driven to eastward by the influence of strong wind. The average range of NOx concentrations in the entire domain was 70-120 µg/m³ simulated by WRF/Chem. Since, the photochemical lifetime of NOx depends strongly on presence of O₃ concentrations through NOx removal reactions, especially conversion to N₂O₅ via O⁺NO₂→NO₃+O₂ and NO₂+NO₃→N₂O₅[58], NOx concentration was about 50-60 µg/m³ in the purple area of the Figure 7.

CONCLUSIONS

By using the new generation of regional air quality model WRF/Chem V3.3, a two days heavy-pollution episode in March 2008 was investigated. A rapid O₃ changes from 00:00 UTC of March 14 to 00:00 UTC of March 16 was observed in the entire domain of the coastal area of the Sea of Japan. During this 2 days period, the daytime and nighttime concentrations of ozone was changed from 50 ppbv to 30 ppbv in the entire domain. In order to understand the processes in controlling this rapid change in O₃ concentrations, a three dimensional regional chemical/dynamical model (WRF/Chem) is applied to analyze the causes of this rapid O₃ changes with meteorological conditions. The result shows that the calculated trends of O₃ concentrations, O₃ precursors, are consistent with the measured trends. The calculated magnitudes and diurnal variations of O₃ are slightly underestimated to the measured results in most of the rural areas, while are overestimated at a urban populated areas, suggesting that the industrial emissions or emission from biomass burning at this sites are significantly underestimated in this study and needs to be improved. Because the model horizontal resolution is 1° × 1° in this study, the relatively coarse resolution also can lead to the overestimation in O₃ calculation. Therefore, the resolution of WRF/Chem in this study may not be enough to predict the ozone concentrations situation over Japanese populated areas. Besides, since the model is capable to calculate the rapid O₃ change, it is suitable to study the causes of the O₃ change trend. The analysis of model result suggests that weather conditions play important roles in controlling the surface O₃ in the coastal areas of the Sea of Japan. Under the winter weather system, the high O₃ concentrations, which were chemically formed in the inland areas, were rapidly transported to the downwind region of the coastal areas, resulting in low O₃ concentrations in the urbanized region of East Asia. This study illustrates that the WRF/Chem model is a useful tool to study the high variability of O₃ concentrations. The study of variability of O₃ has important implication for the prediction of O₃ concentrations and for the control strategies of high O₃ events in the coastal areas of the Sea of Japan.
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