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Title	Cause of a Lower - Tropospheric High - Ozone Layer in Spring Over Hanoi
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Citation	Journal of Geophysical Research: Atmospheres, 127(10), e2021JD035727 https://doi.org/10.1029/2021JD035727
Issue Date	2022-04-29
Doc URL	https://hdl.handle.net/2115/90912
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Type	journal article
File Information	JGR Atmospheres - 2022 - Ogino - Cause of a Lower - Tropospheric High - Ozone Layer in Spring Over Hanoi.pdf



JGR Atmospheres

RESEARCH ARTICLE

10.1029/2021JD035727

Key Points:

- Ozonesonde observations in Hanoi, Vietnam, for about 14 years since 2004 display an ozone enhancement at ~3 km altitude in March–April
- Enhanced ozone air at ~3 km originates from burning areas west of Hanoi whereas low-ozone near-surface air is marine from the east
- Model experiments show biomass fires in the Indochina Peninsula are the cause of Hanoi's enhanced ozone

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Citation:

Ogino, S.-Y., Miyazaki, K., Fujiwara, M., Nodzu, M. I., Shiotani, M., Hasebe, F., et al. (2022). Cause of a lower-tropospheric high-ozone layer in spring over Hanoi. *Journal of Geophysical Research: Atmospheres*, 127, e2021JD035727. <https://doi.org/10.1029/2021JD035727>

Received 27 AUG 2021

Accepted 23 APR 2022

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Cause of a Lower-Tropospheric High-Ozone Layer in Spring Over Hanoi

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Abstract The ozonesonde observations in Hanoi, Vietnam, over 14 years since 2004, have confirmed an enhancement in lower tropospheric ozone concentration at about 3 km altitude in the spring season. We investigated the evolution of the ozone enhancement from analysis of meteorological data, backward trajectories, and model sensitivity experiments. In spring, air masses over Hanoi exhibit strong height dependence. At 3 km, the high-ozone air masses originate from the land area to the west of Hanoi, while low-ozone air masses below about 1.5 km are from the oceanic area to the east. Above 4 km, the air masses are mostly traced back to the farther west area. The chemical transport model simulations revealed that precursor emissions from biomass burning in the inland Indochina Peninsula have the largest contribution to the lower tropospheric ozone enhancement, which is transported upward and eastward and overhangs the clean air intrusion from the ocean to the east of Hanoi. At this height level, the polluted air has the horizontal extent of about 20° in longitude and latitude. The polluted air observed in Hanoi is transported further east and widely spread over the northern Pacific Ocean.

Plain Language Summary Ozone in the lower atmosphere acts as an air pollutant and is one of the strong greenhouse gases. Understanding chemical and transport processes that control ozone variations provide important implications for the origin of air pollution and for climate change. In the past 20 years, the global tropospheric ozone amount has increased in the tropics and subtropics, with particularly large increases in Southeast Asia. We analyze a seasonal enhancement in ozone over Hanoi, Vietnam, using the 14-year record of upper air ozone measurements that began in 2004. The measurements reveal high levels of ozone particularly during the spring that peak at around 3 km. Numerical model simulations show that agricultural burning in the inland Indochina Peninsula is the major contributor to these high ozone amounts that are transported upward and eastward and overhang the clean air from the ocean to the east of Hanoi. Over time, the high-ozone air observed in Hanoi is transported further east and widely spread over the northern Pacific Ocean. The identified source of the ozone enhancements and its three-dimensional distribution provide new insights into the impact of agricultural burning over Southeast Asia on large-scale ozone distributions and are relevant to global air quality and climate studies.

1. Introduction

Tropospheric ozone is closely related with air quality by producing hydroxyl radicals, which control the oxidizing capacity of the atmosphere and is itself an air pollutant and one of the strong greenhouse gases (Brasseur et al., 2003; IPCC, 2013). It is important to describe and understand the mechanism determining the three-dimensional (3D) distribution of ozone and its temporal variation in order to trace the origin of air pollution and to understand climate variability and change.

Because of the strong ozone production efficiency, changes in precursor emissions associated with economic and natural activity can lead to substantial changes in ozone in the tropics and subtropics (Cooper et al., 2020; Gaudel et al., 2020; Y. Zhang et al., 2021). While chemistry–climate models predict substantial changes in tropospheric ozone over these regions in the future climate scenarios (e.g., Morgenstern et al., 2017; Revell et al., 2018; Young

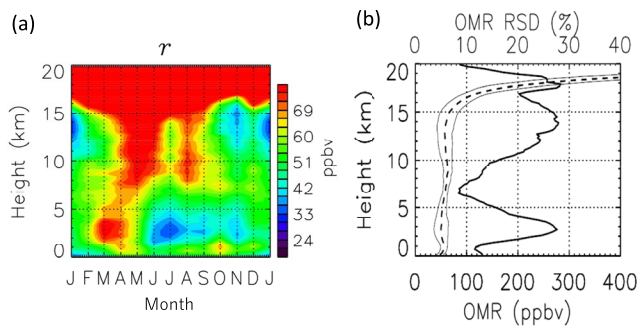


Figure 1. (a) Fourteen-year-averaged seasonal variation in ozone mixing ratio (OMR) obtained by calculating monthly mean values at each height to 20 km from September 2004 to June 2018. Each monthly mean value is plotted at each vertical dotted line. (b) Vertical profiles of annual-mean OMR (dashed line), standard deviation of fourteen-year-averaged seasonal variation in OMR (thin solid line), and standard deviation normalized by the annual-mean OMR at each height (thick solid line). These panels update Figure 1 in Ogino et al. (2013) by adding 7 years data.

Since September 2004, we have conducted regular ozonesonde observations once or twice a month in Hanoi (21.02°N, 105.80°E), Vietnam, on the Indochina Peninsula (Ogino et al., 2013), as part of the activities of Soundings of Ozone and Water in the Equatorial Region (SOWER; Hasebe et al., 2013) and Southern Hemisphere ADDitional OZonesondes (SHADOZ; Thompson et al., 2012). Hanoi lies at the southeastern edge of the Asian summer monsoon circulation, which is a region of exchange of mass and chemical species between the tropical troposphere and the extratropical stratosphere in the Eastern Hemisphere. Southeast Asia is one of the important regions, where tropospheric ozone shows a long-term increasing trend, suggesting a significant contribution to global ozone change (Gaudel et al., 2018; Y. Zhang et al., 2016). In this regard, chemical and meteorological observations in Hanoi can provide valuable information about regional pollution and transport patterns.

Figure 1a shows the seasonal variation in the ozone mixing ratio (OMR) observed in Hanoi averaged over about 14 years from September 2004 to June 2018. There are two height ranges with a large amplitude of seasonal variation. One is in the upper troposphere and lower stratosphere (UTLS) region (10–20 km), where ozone increases in spring (April to June) and summer (August), and decreases in winter, owing to the seasonal change of transport in the UTLS region. There is a northward transport of low-ozone air in winter due to the Rossby wave response to convective heating over the Maritime Continent and the western Pacific. This is followed by a southward transport of high-ozone air in summer due to the upper-tropospheric anticyclonic monsoon circulation associated with the South Asian High, as shown by Ogino et al. (2013). The other height range for large seasonal variability appears in the lower troposphere, with a peak OMR at around 3 km in the pre-monsoon season (March–April). This study aims to identify the origin of this peak OMR and propose a mechanism for the seasonal ozone enhancement.

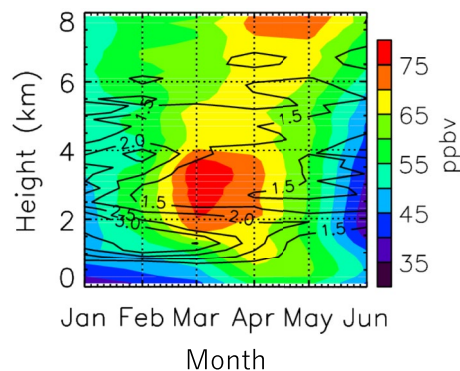


Figure 2. Fourteen-year-averaged seasonal variation in ozone mixing ratio obtained by calculating monthly mean values at each height to 8 km from January to June from 2004 to 2018. Black contours show fourteen-year-averaged seasonal variation of the Brunt-Väisälä number ($\text{rad}^2 \text{s}^{-2} \times 10^4$). Each monthly mean value is plotted at each vertical dotted line.

et al., 2013), the multi model discrepancies remain large in predicted ozone and its radiative forcing (Kuai et al., 2020). Ozone in these regions is also important for human health associated with the high population density and rapid economic development. However, the current in-situ observing network in these regions, for instance from Tropospheric Ozone Assessment Report (TOAR), is clearly insufficient for detailed evaluation of ozone changes for various applications including air pollutant human health impacts (Fleming et al., 2018). Studies using numerical models and limited observations have also illustrated the overall picture of non-local and local atmospheric pollution changes including ozone over South and Southeast Asia, such as wintertime northeasterly transports at low levels and summertime uplifting associated with Asian summer monsoon together with strong impacts of biomass burning (Lawrence & Lelieveld, 2010). Nevertheless, complete 3D structure of ozone in these regions and its driving factors remain unclear mainly due to a lack of detailed observational information. While long-term polar orbiting and recent geostationary satellite measurements, such as Geostationary Environment Monitoring Spectrometer (GEMS; Kim et al., 2020), can provide improved understanding of tropospheric ozone changes, vertical profile information from regular ozonesonde records is essential to study detailed chemical and physical processes.

The other height range for large seasonal variability appears in the lower troposphere, with a peak OMR at around 3 km in the pre-monsoon season (March–April). This study aims to identify the origin of this peak OMR and propose a mechanism for the seasonal ozone enhancement.

Figure 2 shows the seasonal variation in the OMR up to 8 km height in the first half of the year, and the atmospheric stability, measured by the Brunt-Väisälä number. It shows two stable layers at ~1.5 and ~5 km in March, as reported by Nodzu et al. (2006). There is an ozone maximum between those layers that implies a distinct and separate feature.

Over the Indochina Peninsula, March is a transition period from the dry season to the rainy season. Intermittent rainfall begins in April (Kiguchi & Matsumoto, 2005; Matsumoto, 1997), and the summer monsoon season starts typically in May (He et al., 1987; Hsu et al., 1999; Nguyen-Le et al., 2015; Yanai et al., 1992), the earliest onset in the Asian monsoon region.

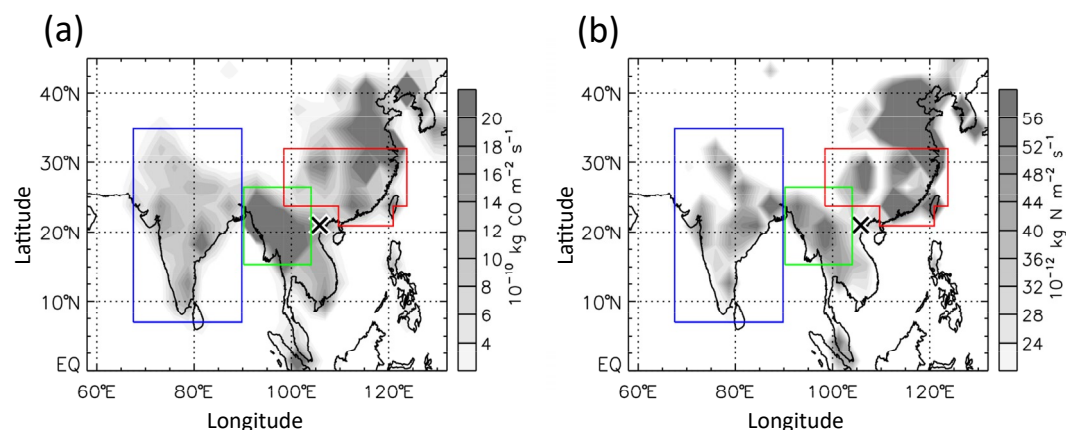


Figure 3. The rectangles show the regions where the ozone precursor emissions were eliminated in the sensitivity experiments: (Blue) the Indian Subcontinent, (Green) the northern Indochina Peninsula, and (Red) southern China. The gray shade shows the (a) carbon monoxide and (b) nitrogen oxide emissions from the surface in March averaged for 10 years from 2005 to 2014 adopted in the control run of the numerical experiment. A cross mark shows the location of Hanoi, Vietnam.

A similar but smaller ozone increase at the similar height and in the same season was observed in Hong Kong (22.31°N, 114.17°E; L. Y. Chan et al., 1998; Ogino et al., 2013). Several authors showed that such ozone increases could be attributed to the transport of air polluted by biomass burning over Southeast Asia. Their analysis was based on backward trajectories that tend to pass over the active region of biomass burning, such as the Indian subcontinent and the northern Indochina Peninsula (C. Y. Chan, Chan, Chang, et al., 2003; C. Y. Chan, Chan, Harris, et al., 2003; C. Y. Chan & Chan, 2000; Liao et al., 2021). Liu et al. (2002) further confirmed that Southeast Asian biomass fires are responsible for the ozone enhancement observed in Hong Kong based on model simulations in which biomass burning emissions were eliminated. Recently, Liao et al. (2021) showed that the spring time ozone concentration above Hong Kong has an increasing trend possibly due to the biomass burning increase over the upwind region.

Burning activities are maximized in March over the northern part of the Indochina Peninsula (Huang et al., 2016), leading to enhanced surface ozone (Pochanart et al., 2001) and tropospheric column ozone (Sonkaew & Macatangay, 2015) in Thailand, an inland region of the Indochina Peninsula. These studies suggest that air polluted by biomass burning is a plausible cause of the ozone increase at about 3 km above Hanoi in March.

Most of the previous studies on the source of the ozone enhancement relied on backward trajectory analyses over Southeast Asia. However, traditional trajectory analysis does not provide any information on which emissions within Southeast Asia contribute most to the ozone enhancement quantitatively. It is also unclear why the ozone increase in March appears at about 3 km height above the surface. C. Y. Chan, Chan, Harris, et al. (2003) showed that the increase was accompanied by a stable layer at the bottom of the layer with the ozone increase and mentioned that the inversion isolated the ozone rich layer from the influence of the local boundary layer. Nonetheless, the formation process and spatial distribution of the stable layer have not yet been fully described, and therefore, its relation to the high ozone layer is not clearly understood. The 3D structure of the ozone enhancement should be described by relating it to meteorological phenomena.

In this paper, we carry out a comprehensive study on the ozone enhancement with thermodynamic properties and the detailed analysis of chemical model sensitivity experiments. We first describe the variabilities of the ozone enhancement in March together with background meteorological properties, in which we identify three regimes in terms of ozone variability separated by the stable layers at about 1.5 km and about 4 km.

Next, we investigate the source region of ozone precursors that contributes the ozone enhancement above Hanoi by conducting sensitivity experiments using a chemical transport model with the observationally constrained top-down emissions. We further reveal the 3D structure of the ozone enhancement and investigate the mechanisms controlling the ozone enhancement that appears at about 3 km above Hanoi.

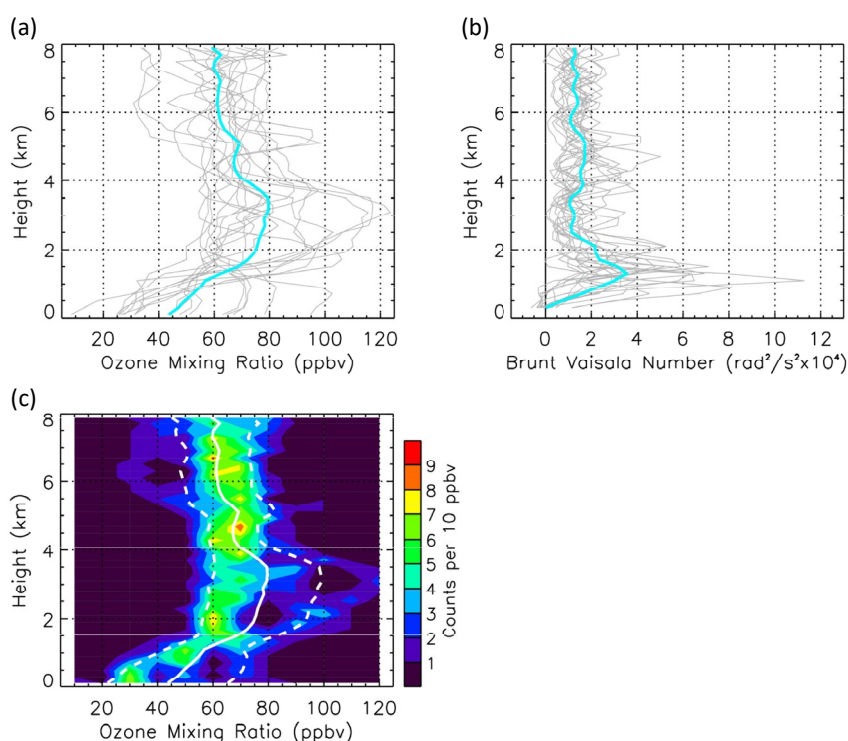


Figure 4. Vertical profiles of (a) ozone mixing ratio and (b) Brunt-Väisälä number observed by the ozonesondes in March for the whole time period (2005–2018). Blue line shows the mean of all data in each panel. (c) Frequency distribution (shade), the mean (white solid line) and one standard deviation on the either side of the mean (white dashed line) of ozone mixing ratio.

2. Data and Methods

We used ozonesonde data collected at the Aero-Meteorological Observatory (21.02°N, 105.80°E), Hanoi, once or twice in each month from September 2004 to June 2018. The observations are described in detail in Ogino et al. (2013). The data were re-processed in 2017 and 2018 (Witte et al., 2017, 2018) to obtain the homogenized data (SHADOZ Version 6 data sets). Total ozone from the soundings is in good agreement with satellite overpasses (e.g., from OMI and OMPS; Thompson et al., 2017; Witte et al., 2018) as well as from the Brewer spectrometer co-located at the Aero-Meteorological Observatory launch site. We used operational radiosonde data taken at the Observatory in Hanoi and the Japanese 55-year Reanalysis (JRA-55) objective analysis data (Kobayashi et al., 2015) to investigate meteorological fields.

The JRA-55 data were also used in back-trajectory calculations. Isentropic (without considering diabatic vertical motion) back-trajectories were calculated every 30 min using linearly interpolated JRA-55 meteorological reanalysis data (Kobayashi et al., 2015) and the second-order Runge-Kutta method for time integration, as in Hasebe et al. (2007) and Ogino et al. (2013). In each trajectory calculation, an initial point was set at every 200 m height from 100 m to the upper troposphere above Hanoi at the ozonesonde launch time in Hanoi.

To examine the relative importance of different emission source regions on the ozone enhancement above Hanoi, we performed sensitivity experiments using the global chemical-transport model, CHASER (Sudo et al., 2002) with T42 horizontal resolution (approximately 2.8° longitude $\times$ 2.8° latitude) and 32 vertical layers from the surface up to 10 hPa in sigma coordinate. The 2-hourly model outputs interpolated onto the constant pressure levels at 1,000, 990, 970, 930, 870, 790, 700, 610, 530, 460, 400, 350, 300, 260, 230, 200, 176, 153, 133, 116, 100, 88, 76, 66, 58, 50, 43, 36, 30, 22, 14, and 10 hPa were used in this study. Note that the updated model, MIROC-Chem (Miyazaki et al., 2017; Watanabe et al., 2011), includes more detailed chemical processes for both troposphere and stratosphere. Nevertheless, CHASER already includes the most important chemical processes in the NO_x -CO-Ozone reactions and can be used to evaluate the impact of NO_x emissions on ozone productions.

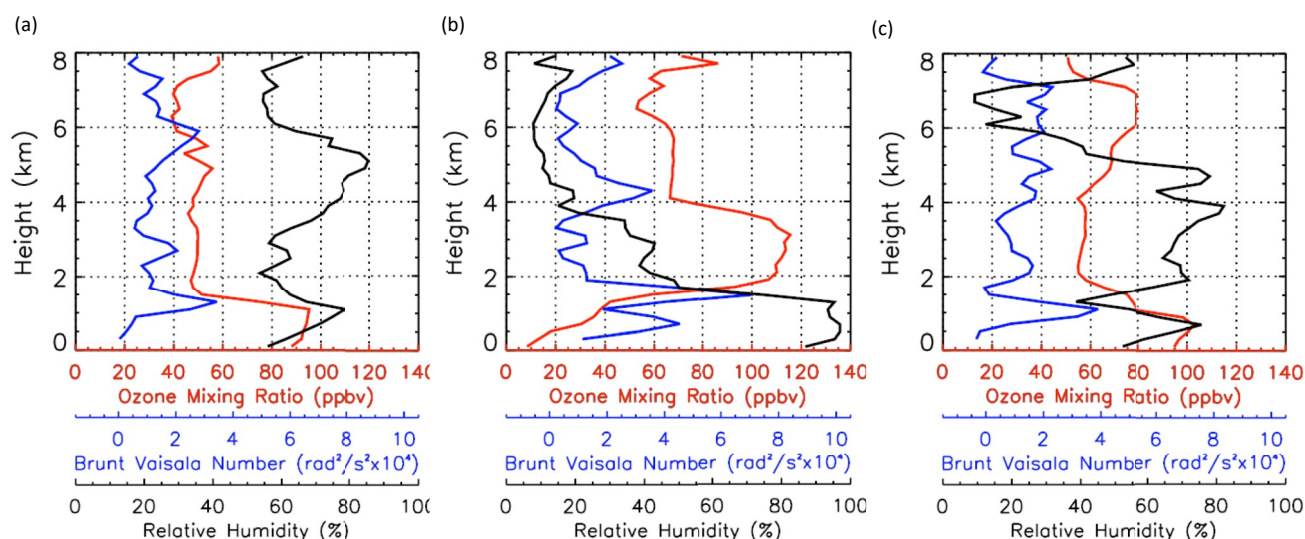


Figure 5. Vertical profiles of the ozone mixing ratio (red), the Brunt-Väisälä number (blue), and relative humidity (black) on (a) 26 April 2011, (b) 12 March 2007, and (c) 27 March 2009.

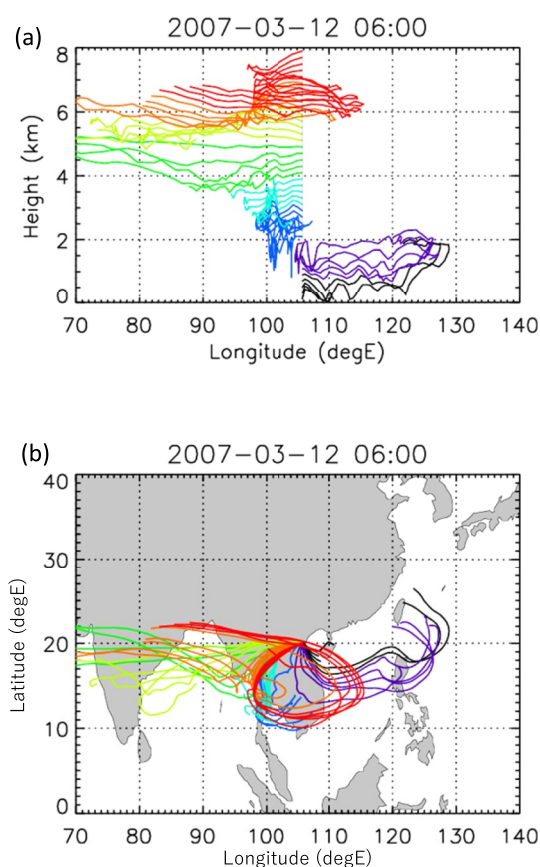


Figure 6. Backward trajectories initialized at 06:00 UT on 12 March 2007 plotted on (a) longitude-height section and (b) longitude-latitude section. Color of each trajectory differs depending on its starting height as found in $\circ$.

In addition, the simulated ozone performance and ozone response to NO_x emissions are comparable between CHASER and MIROC-Chem (Miyazaki et al., 2020). Thus, the results should not be sensitive to the choice of model.

The surface emission of major ozone precursors, such as carbon monoxide (CO), nitrogen oxide (NO_x), and nonmethane hydrocarbons, were included in the model based on the published emission inventories [the Emission Database for Global Atmospheric Research (EDGAR) version 4.2 (EC-JRC/PBL, 2011), the monthly Global Fire Emissions Database (GFED) version 3.1 (van der Werf et al., 2010), and monthly mean Global Emissions Inventory Activity (GEIA; Graedel et al., 1993)]. We employed daily NO_x and CO emissions that were optimized using the assimilation of satellite NO₂ and CO measurements, where the a priori emissions were constructed based upon bottom-up emission inventories (Miyazaki et al., 2015, 2017). These emissions, including both anthropogenic and biomass burning components, were obtained from the Tropospheric Chemistry Reanalysis version 1 (TCR-1, Miyazaki et al., 2015) and enabled us to evaluate the emission impacts for individual sources.

In the sensitivity experiments, we eliminated the emissions of ozone precursors from the following three source regions: the Indian subcontinent, the northern Indochina Peninsula, and southern China (Figure 3). We conducted spin-up calculations with the optimized emissions for all regions (i.e., standard emissions) from January 1st to the end of February in each year for 10 years from 2005 to 2014. Then, we performed four types of experiments from March 1st to 21st: the control experiment with the standard emissions (Figure 3), and the three sensitivity experiments with elimination of emission from the above-mentioned three regions, namely the Indian subcontinent, the northern Indochina, the southern China experiments. Because of the non-linear chemistry, the cumulative response from the sensitivity calculations can be different from the total ozone response in the control simulation to some extent as shown by the HTAP modeling work (Turnock et al., 2018; Wild et al., 2012). Nevertheless, they provided important information on relative contributions of emission sources from different regions. The results of the

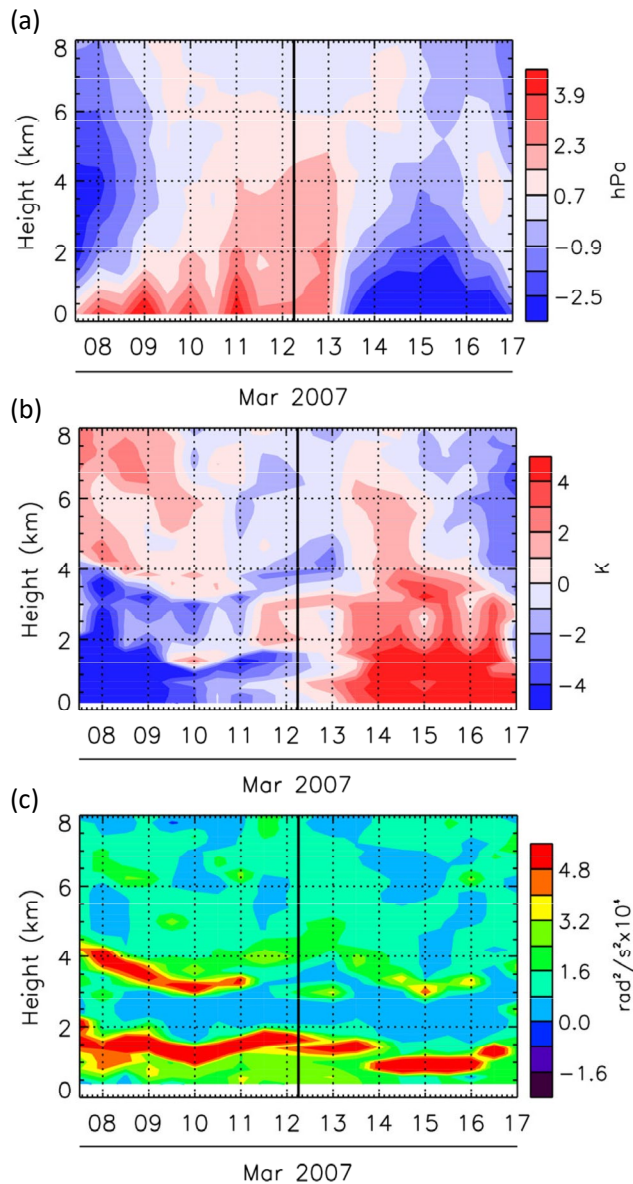


Figure 7. Time-height sections of (a) pressure anomaly, (b) temperature anomaly, and (c) Brunt-Väisälä number (N^2) drawn from operational radiosonde data collected in Hanoi from 7 to 17 March 2007. The anomalies are deviations from the 10-day mean at each height. Vertical solid line denotes the ozonesonde observation at 0600 UT on 12 March 2007.

sensitivity experiments will be compared with the control experiment to investigate the relative contributions of individual emission sources to the ozone enhancement over Hanoi. Note that although the period of the model experiment (10 years) is shorter than that of the observation (about 14 years), we consider that the 10-year model data are sufficient for our purpose to describe the climatological feature of the ozone distribution.

3. Characteristics of Ozone Profiles in March

Figure 4a shows all of the vertical profiles of the OMR observed in Hanoi in March for the whole time period from 2005 to 2018. Figure 4b shows the vertical profiles of the Brunt-Väisälä number for the same ozonesonde observations. The profiles show that the ozone variability is large in three height ranges—0–1.5, 1.5–5, and 5–8 km—which are separated by two stable layers at ~1.5 and ~5 km first shown by Nodzu et al. (2006). For example, at 1.5–5 km, where the mean OMR in March peaked, the OMR is 50–120 ppbv. At 0–1.5 km the OMR was 0–100 ppbv, and at 5–8 km it is 30–80 ppbv. On the other hand, the variability in OMR is relatively small near the two stable layers at ~1.5 and ~5 km, at 50–80 ppbv. The large variability at the three height ranges is confirmed by calculating the standard deviation at each height as shown in Figure 4c. Figure 4c also shows the frequency distribution of OMR at each height, in which we find different characteristics between the three height ranges. In 1.5–4 km, although the mode is about 60 ppbv, the distribution skews positively and the less-frequent large OMR values (>80 ppbv) contribute to the ozone enhancement with the large mean value (75–80 ppbv) at this height range. In the layer above 5 km, the distribution shows slight negative skewness, and the mean values are smaller than those in 1.5–4 km, although the modes are similar to or slightly larger than those in 1.5–4.0 km. In the layer below 1.5 km, the frequency distribution skews positively, and both mode and mean decrease as height decrease.

Figure 5 shows three typical profiles of the ozone increases at 0–1.5 km (a), 1.5–5 km (b), and 5–8 km (c). On 26 April 2011 (Figure 5a), the OMR increased below the lower stable layer. On 12 March 2007 (Figure 5b), the height profile is similar to that seen in the 14-year-averaged height profile of March (Figure 1b). This case also shows an ozone minimum at 0–1.5 km just below the lower stable layer. Note that there is also a sharp contrast in relative humidity between the levels below and above the lower stable layer: a humid (almost saturated) air mass below, and a relatively dry (about 40% RH) air mass above it. On 27 March 2009 (Figure 5c), the OMR increased above the upper stable layer at about 5 km.

The ozone increases in each of the three height ranges described above seem to be controlled by different mechanisms. Below, we investigate the mechanism of ozone variation in each layer. First, we investigate whether the ozone

density above Hanoi depended on the origin of the air mass by using backward trajectory analysis. Next, we interpret the obtained typical trajectories in terms of transport processes and meteorological conditions.

4. Analysis of a Typical Case

First, we examine backward trajectories and atmospheric fields for a typical ozone increase at 1.5–5 km and for a typical ozone decrease at 0–1.5 km observed on 12 March 2007 shown in Figure 5b. Figure 6 shows 5-day backward trajectories initialized at 0600 UT (launch time of the ozonesonde) on 12 March 2007 at every 200 m height above Hanoi. The backward trajectories show that the origins of the air masses differed between the height ranges above and below the lower stable layer at about 1.5 km. The air mass in the high-ozone layer at 1.5–4 km can be

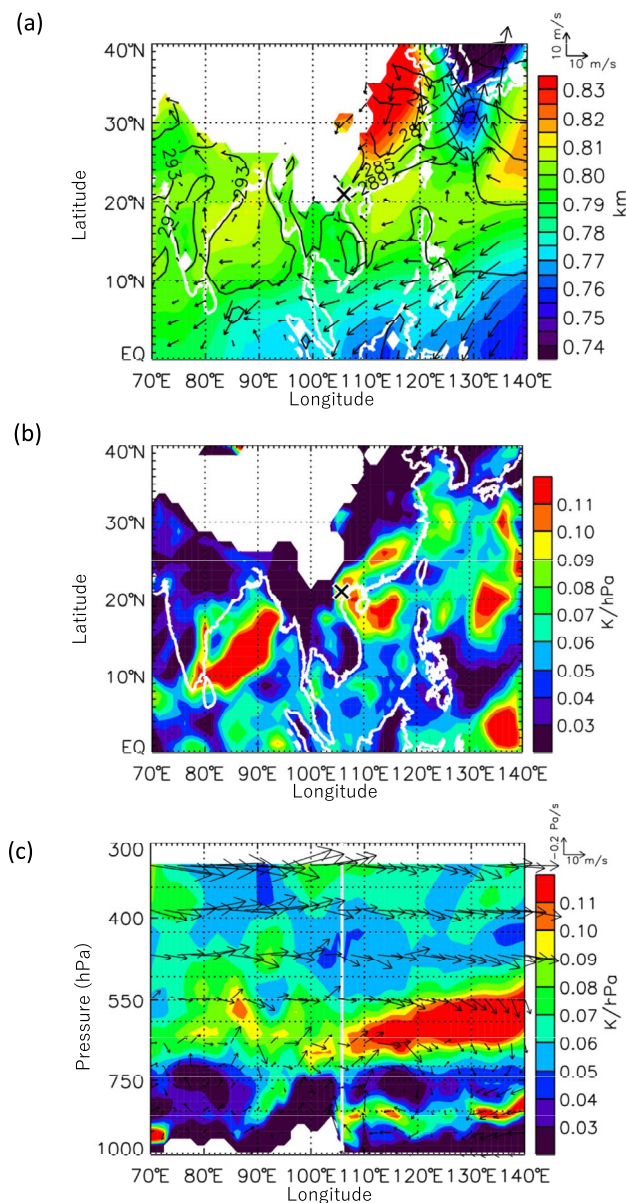


Figure 8. Horizontal distributions of (a) geopotential height (colors) and temperature (contours) at 925 hPa, (b) vertical stability (vertical gradient of potential temperature $-\partial\theta/\partial p$) at 862.5 hPa, and (c) longitude-pressure section of vertical stability at 20°N at 06:00 UT on 10 March 2007 drawn with JRA-55 data. The regions under the ground surface are masked with white color. Black cross marks in (a) and (b) denote the location of Hanoi. A white vertical line in (c) denotes the longitude of Hanoi.

traced back to the Indochina Peninsula and nearby to the west of Hanoi. The air masses in the low-ozone layer below 1.5 km, on the other hand, originated from the South China Sea and the western Pacific area to the east of Hanoi. This difference is consistent with the observations of humid air below 1.5 km and dry air at 1.5–4 km mentioned in Section 3. The change in ozone levels suggests that ozone concentration of the air mass from the Indochina Peninsula was enhanced by biomass burning, while that from the ocean areas was less polluted. The air masses above the upper stable layer (4–6 km) can be traced back to the farther west area than those at 1.5–4 km, suggesting that the mechanism that determines the ozone variation in this height range is different from those below about 4 km, and that the long-range transport, and various sources and source regions, such as air pollution far from Hanoi and stratospheric ozone inputs, can contribute the ozone variations in this height range. In the rest of the paper, we mainly investigate and discuss the ozone variations in the surface–1.5 and 1.5–4 km height ranges.

The pressure and temperature anomaly fields above Hanoi from 7–17 March 2007 (Figures 7a and 7b) reveal a high-pressure, cold-temperature anomaly in the first half of the period. The Brunt-Väisälä number distribution shows clear stable layers at ~ 1.5 and ~ 4 km (Figure 7c).

Figure 8a shows the horizontal distribution of the geopotential height and temperature of JRA-55 at 925 hPa on 10 March 2007, 2 days before the ozonesonde observation. The tongue-shaped high-pressure (Figure 8a, yellow to red), and its associated low-temperature represent an intrusion from eastern China toward Hanoi due to the anti-cyclonic circulation. This structure is characteristic of a cold surge typically seen in winter and spring over Southeast Asia (Compo et al., 1999). The stable layer just above the cold air intrusion was strengthened locally near Hanoi as seen in Figures 8b and 8c. Thus, the near-surface easterly flow bringing the low-ozone air mass from the ocean area shown by the backward trajectories in Figure 6 was associated with this cold surge event, which simultaneously transports the cold air to produce the stable layer at about 850 hPa (about 1.5 km). Figure 8c also shows the development of unstable layer from the surface to about 600 hPa (4–5 km) above the land regions of the Indian Subcontinent and the Indochina Peninsula. This is likely caused by the development of a deep mixed boundary layer (06:00UT corresponds to about noon time near the longitude region concerned) generated by the increased heating over the land surface through the seasonal transition from winter to summer (Nodzu et al., 2006; Ogino et al., 2010). This suggests that the polluted high-ozone air over the Indian Subcontinent and Indochina Peninsula can be lifted to 4–5 km level and be transported eastward to the ozone enhancement above Hanoi.

5. Model Experiment

Here, we use the chemical transport model simulations to further explore the relative contributions of individual emission source regions (cf., Section 2).

Figure 9 shows a time-height section of the 10-year mean of monthly mean OMR at the grid point near Hanoi in the control experiment using the standard optimized emissions constrained by satellite observations. The control experiments well reproduce the seasonal variations in OMR. For instance, the simulated OMR reproduces the observed relative ozone maxima in March at about 700 hPa (about 3 km in height, Figure 2), although the simulated peak value (65 ppbv) is smaller than the ozonesonde measured peak (85 ppbv) probably due to the coarser vertical resolution of the model and the model biases, such as the low stratospheric ozone input and the insufficient chemical production of ozone as suggested by Park et al. (2021). The model also well reproduces the

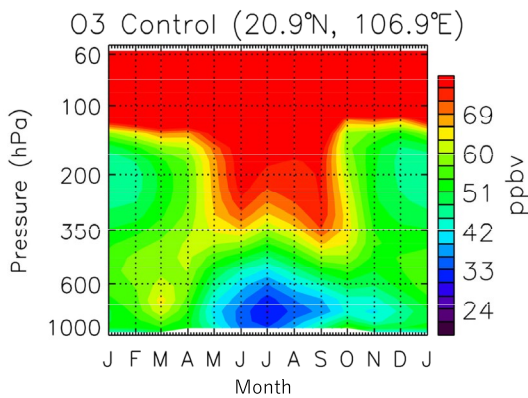


Figure 9. Time-height cross section of the 10-year mean of monthly mean OMR at the grid near Hanoi reproduced in the control experiment of the sensitivity experiments using the chemical transport model, CHASER. Each monthly mean value is plotted at each vertical dotted line. The regions under the ground surface are masked with white color.

observed relative timing and location of the seasonal and vertical distributions, such as the two maxima in May–June and September–October in the upper troposphere (400–200 hPa). About a 1-month delay in the seasonal in the upper troposphere simulated in the model relative to that observed with the ozonesondes implies systematic biases in the model chemical and physical processes, but it does not appear in the lower troposphere.

Temporal variability in ozone reproduced in the model shows similar features to the observation shown in Figure 4. Figures 10a and 10c show that the ozone variability is large at three height ranges—950–900 hPa (near-surface), 790–700 hPa (about 3 km, where the mean ozone maximum exists), and above 500 hPa—which are separated by the two stable layers at about 900 hPa and at 650–400 hPa. The frequency distribution (Figure 10c) shows positive skewness at 790–700 hPa, and the outlying large ozone values contribute to the mean value at this pressure level, which is the same feature with the observational results shown in Figure 4. Thus, the characteristic of ozone variability observed by the once- or twice-monthly ozonesondes was appropriately reproduced by the model experiment; conversely, the ozone-sonde observation has sampled ozone profiles without strong sampling biases.

At 700 hPa, the control experiment (Figure 11a) reveals high OMR (>62 ppbv) over each of the eastern parts of the two regions: the northeastern part of Indian subcontinent and northeastern part of the Indochina Peninsula. By eliminating the ozone precursor emission over the Indian subcontinent region, the ozone enhancement over northeast India disappears, whereas the one over the northeast Indochina Peninsula remains (Figure 11b). The ozone enhancement over the northeastern part of the Indochina Peninsula, including the Hanoi location, was associated with emissions over the northern Indochina region (Figure 11c). In the case of the southern China experiment, the Indochina signal remains (Figure 11d). The contribution from emissions over each region can

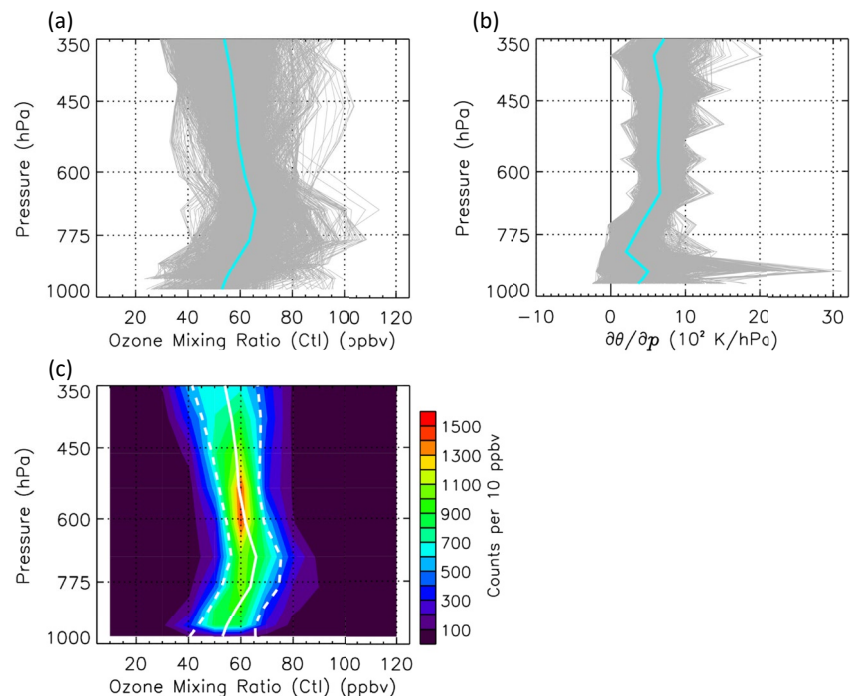


Figure 10. Vertical profiles of (a) ozone mixing ratio and (b) vertical gradient of potential temperature reproduced in the control experiment. All the 2-hourly profiles in March 1–21 for 10 years from 2005 to 2014 are plotted in gray lines, and the mean of each parameter is plotted in blue line in each panel. (c) Frequency distribution (shade), the mean (white solid line) and one standard deviation on the either side of the mean (white dashed line) of ozone mixing ratio.

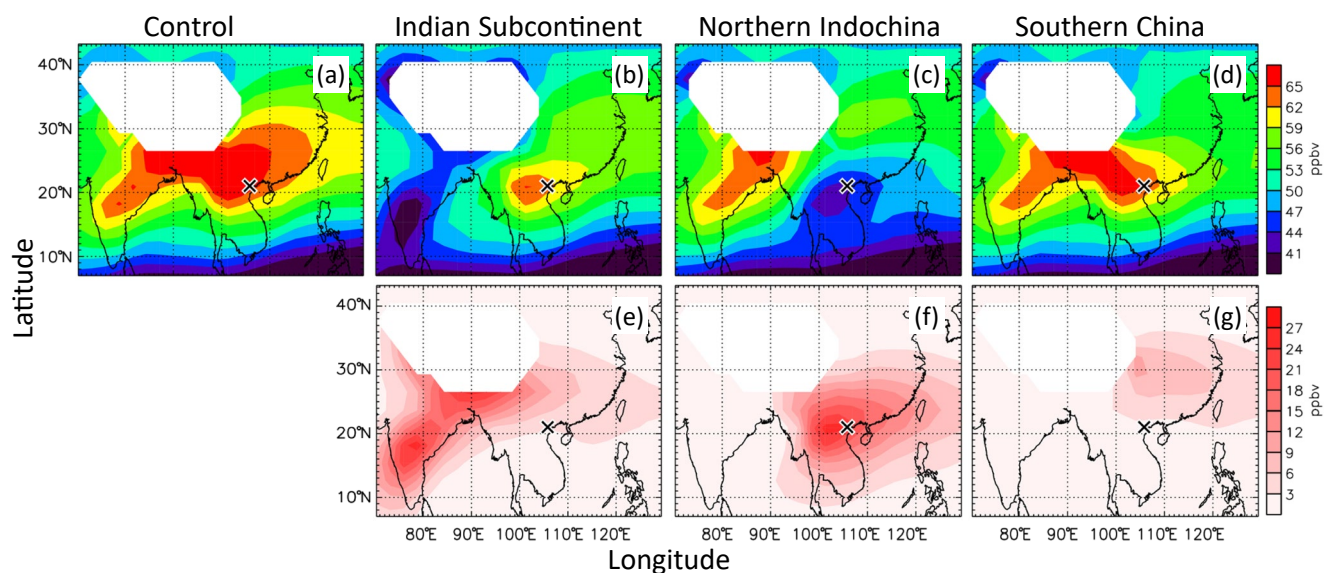


Figure 11. The mean horizontal ozone distributions at 700 hPa for the period from March 1st to 21st obtained from the 10-year sensitivity model experiments. (Upper panels) Ozone mixing ratio for (a) the control, (b) the Indian subcontinent, (c) the northern Indochina, and (d) the southern China experiments. (Lower panels) The ozone production due to the ozone precursors emitted from (e) the Indian subcontinent, (f) the northern Indochina, and (g) the southern China regions. A cross mark in each panel shows the location of Hanoi. The regions under the ground surface are masked with white color.

be more clearly seen by subtracting the result of each sensitivity experiment from that of the control experiment, as seen in Figures 11e–11g. We find that the Indochina emission produces the ozone enhancement above Hanoi and its impact at 700 hPa (about 3 km) spreads around Hanoi with horizontal extent of about 20° in latitude and longitude (Figure 11f).

In terms of vertical structure, the ozone enhancement above Hanoi originates from the northern Indochina emission (Figures 12c and 12f). Figure 12f shows that the ozone enhancement over Indochina (93°E–106°E) is tilting and overhanging toward the east. This is the morphological reason why the ozone enhancement above Hanoi

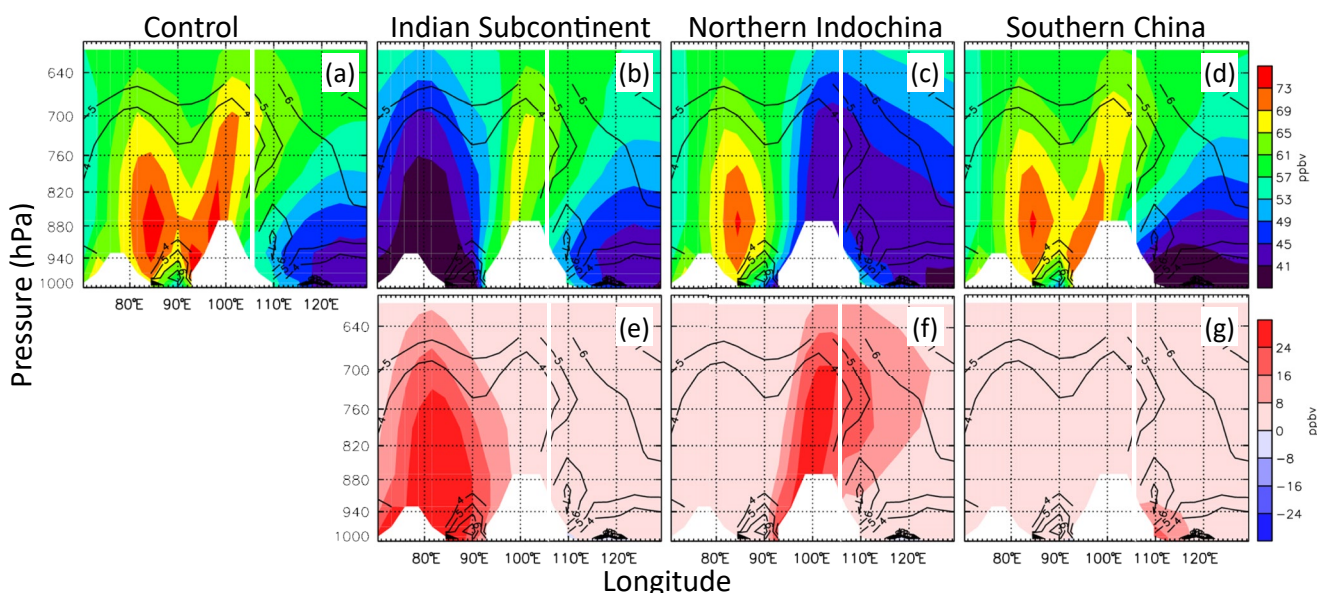


Figure 12. Same as Figure 11 but for zonal-vertical sections at 20.9°N. Contours show the vertical stability estimated by vertical gradient of potential temperature in unit of 10^2 K/hPa. A white vertical line in each panel shows the longitude of Hanoi.

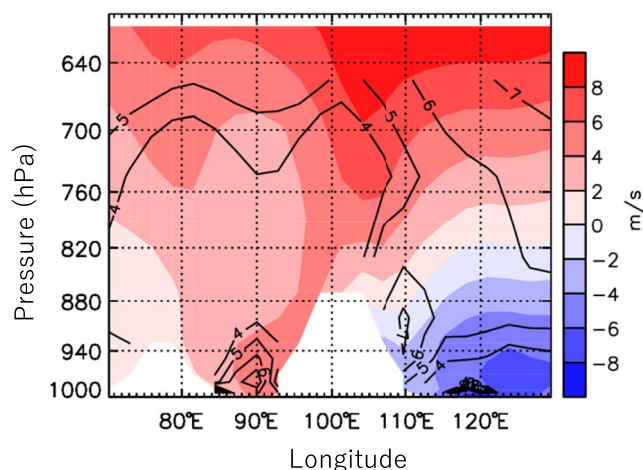


Figure 13. Zonal-vertical section of the zonal wind at 20.9°N reproduced in the control experiment. Contours show the vertical stability estimated by the vertical gradient of potential temperature in unit of 10^2 K/hPa. A white vertical line shows the longitude of Hanoi. The regions under the ground surface are masked with white color.

appears at about 3 km height above the surface. Thus, the overhanging ozone enhancement is formed by the polluted high-ozone air transport from the inland Indochina Peninsula over the clean oceanic air intrusion from the east near the surface. This picture is consistent with the backward trajectories for the typical case shown in the previous section (Figure 6) and the zonal wind distribution shown in Figure 13. Meanwhile, the distribution of carbon monoxide (CO; Figure 14) almost coincides with that of ozone. Because of its direct emissions and relatively long lifetime in the atmosphere, enhanced CO in this region can be regarded as an indicator of air originated from biomass burning and less affected by stratospheric sources unlike ozone. The simultaneous enhancement of ozone and CO indicates that the ozone enhancement occurs due to the biomass burning associated with slash-and-burn agriculture and agro-residue burning.

Figure 15a shows the vertical profiles of OMR above Hanoi obtained in the control and sensitivity experiments. The ozone enhancement appears at 700 hPa with the value of 66 ppbv in the control experiment. The OMR at the peak altitude (700 hPa) reduces to 63, 44, and 64 ppbv in the Indian Subcontinent, the northern Indochina, and the southern China experiments, respectively. The contributions of the ozone precursors emitted over the Indian Subcontinent, the northern Indochina, and the southern China regions on ozone at 700 hPa over Hanoi were estimated from the sensitivity experiments to be 3 ppbv (the Indian Subcontinent experiment), 21 ppbv (the northern Indochina experiment), and 2 ppbv (the southern China experiment;

Figure 15b), accounting for 5% (the Indian Subcontinent experiment), 33% (the northern Indochina experiment), and 3% (the southern China experiment) changes (Figure 15c), with additional 59% contributions from other regions. Although these values from the sensitivity experiment results involve ambiguity to some extent owing to the non-linear chemistry as mentioned in Section 2, they clearly highlight the dominant contribution from the northern Indochina region to the lower tropospheric ozone enhancement above Hanoi in spring.

Near the surface below the lower stable layer at about 1.5 km, the contribution from the southern China precursors is predominant (account for about 20% of ozone production), with almost negligible contributions from the Indian Subcontinent and the northern Indochina precursors (Figure 15). The winter monsoon northeasterly could play a role in bringing the polluted air from the southern China region. This may seem inconsistent with the explanation that the intrusion of less polluted, oceanic air masses associated with the winter monsoon flow is responsible for the low mean ozone concentration the surface as mentioned in Section 4, but it can be reasonably

interpreted as follows. Although the mean flow in March is almost easterly, the daily wind can fluctuate depending on the short time scale disturbances, such as cold surges, which resulted in the mixture of the dominant transport of the clean oceanic air from the east and of the intermittent intrusion of the polluted air from the south China region to the north and the northeast of Hanoi. Such an intermittent intrusion of high ozone event associated with the monsoon activities is a new insight for the near-surface ozone variation, details of which should be studied in a further investigation.

Finally, let us look at the global spread of ozone produced by the Northern Indochina emission. Figure 16 shows the ozone amount generated from the northern Indochina emission during March 1–20 in the middle troposphere (610 hPa). The chemical transport model suggests that ozone originating from northern Indochina can be transported widely eastward in the middle troposphere across the Pacific Ocean, and reach North America and the North Atlantic Ocean, while the signals are also accumulated over the southern mid-latitudes in the middle troposphere. This feature has been reported by several studies based on the airplane experiments, such as Pacific Exploratory Mission-West (PEM-West; Hoell et al., 1997) and Transport and Chem-

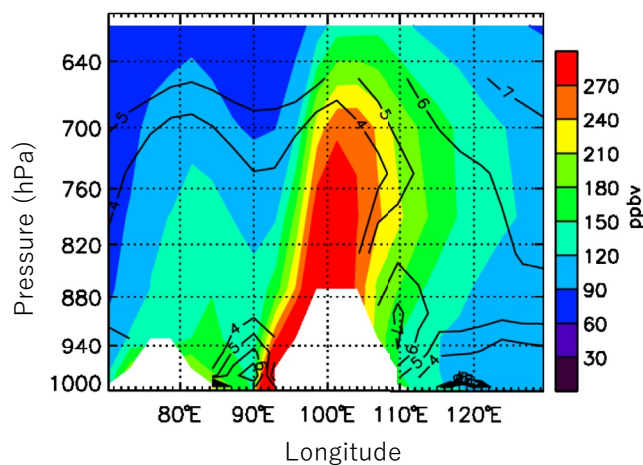


Figure 14. As in Figure 13, but for the CO mixing ratio in the control experiment.

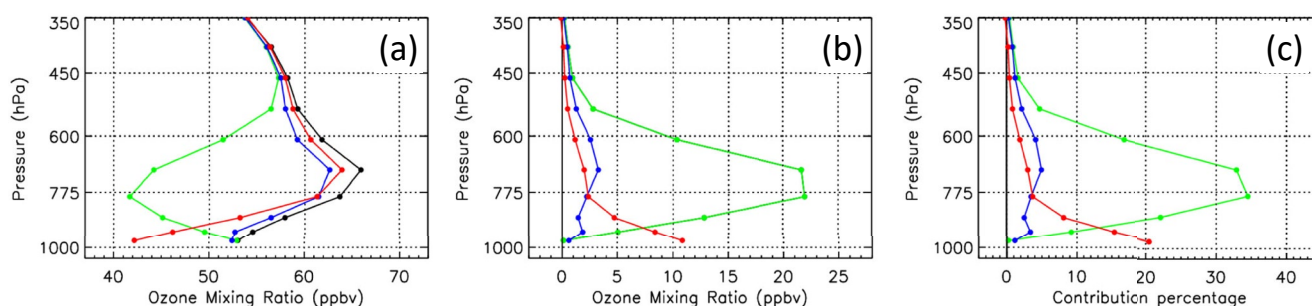


Figure 15. Vertical profiles of (a) ozone mixing ratio obtained from the sensitivity experiment over the model gridpoint nearest to Hanoi. Each line represents a result from (black) the control, (blue) the Indian Subcontinent, (green) the Northern Indochina, and (red) the Southern China experiment. (b) Its difference between the control experiment and each of the sensitivity experiment, which means ozone mixing ratio contribution from each emission region, and (c) contribution percentage due to the ozone precursor emission in each sensitivity experiment (obtained by dividing the difference values show in (b) by the amount obtained from the control experiment shown by the black line in (a)) are also shown.

ical Evolution Over the Pacific (TRACE-P; M. Zhang et al., 2003). The present study has revealed the detailed and 3D structure near the source region and also over remote regions, which would provide important insights into the impact of biomass burning over Southeast Asia on global ozone distributions and the radiation balance of the atmosphere. Further studies using the presented model simulations along with detailed validation using ozonesonde measurements would benefit global air quality and chemistry-climate studies, for instance, under the Hemispheric Transport of Air Pollution (HTAP) project.

6. Discussion

On the basis of the analyses of backward trajectories, meteorological fields, and the numerical experiments, we propose a mechanism, illustrated in Figure 17, for the increase of ozone frequently observed at ~3 km above Hanoi in March. During this month, the atmosphere is relatively unstable, allowing shallow convection to develop below the upper stable layer at ~5 km over the inland Indochina Peninsula and the Indian Subcontinent. It is well known that the occurrence of biomass burning in these regions increases during February to April (Huang et al., 2016). The near-surface polluted air is well mixed by shallow convection up to the height of the upper stable layer. This well-mixed air is advected by the westerly flow; but the air below the lower stable layer is blocked by the near-surface easterly flow with the oceanic, less-polluted airmass, and only the air above the lower stable layer can reach the eastern region of the Indochina Peninsula. The main source of the enhanced ozone above

Hanoi originates from the ozone precursors emitted over the northern part of the Indochina Peninsula. The polluted high-ozone air at the peak height covers the region with a horizontal scale of about 20° in longitude and latitude over the northeastern part of the Indochina Peninsula around Hanoi. The airmasses are transported further eastward and upward and spread over the northern Pacific Ocean in the middle troposphere.

This mechanism occurs from March to April, because land heating is not strong enough in January and shallow convection does not reach 4–5 km as a result (Nodzu et al., 2006). Also, it is before the rainy season that starts in May over the Indochina Peninsula (e.g., He et al., 1987; Hsu et al., 1999; Yanai et al., 1992). Thus, the ozone transport and distribution from March to April are characterized not only by the seasonality of biomass burning but also by the monsoon transition from the dry season to the rainy season.

Our results show that the ozone increase at ~3 km in March above Hanoi resembles that seen over Hong Kong. A polluted air mass from the same origin may sometimes pass over both Hanoi and Hong Kong. Such polluted air masses are considered to originate from the north of the Indochina Peninsula, which is closer to Hanoi than to Hong Kong. This is consistent with the

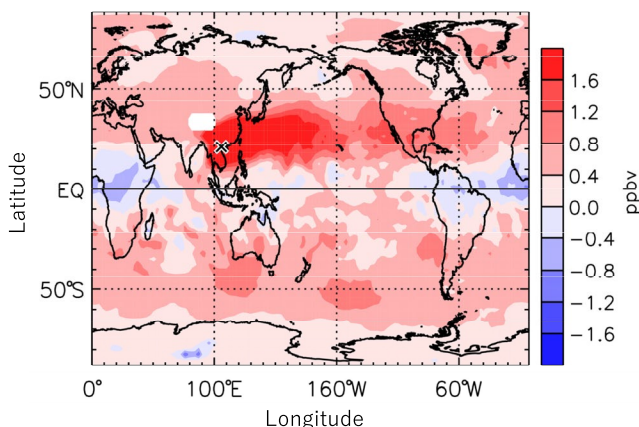


Figure 16. Global distribution of ozone mixing ratio at 610 hPa obtained from the difference between the control experiment and the northern Indochina experiment. A cross mark denotes the location of Hanoi.

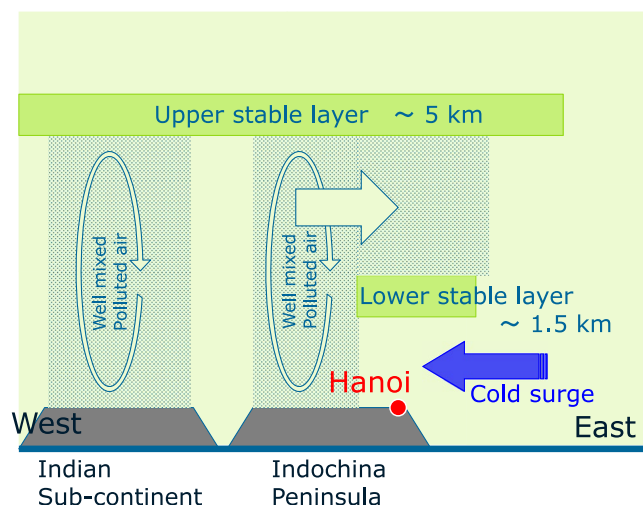


Figure 17. Schematic diagram of the proposed mechanism for the ozone increase at 3 km above Hanoi in March.

fact that the climatological signal of the ozone increase is clearer in Hanoi than in Hong Kong as seen in Figure 1 of Ogino et al. (2013).

We showed that a stable layer (at ~ 5 km) exists at the top of the ozone increase, adding to the finding of the stable layer at the bottom of the ozone increase by L. Y. Chan et al. (2000) and C. Y. Chan et al. (2003). The ozone increase appears between two stable layers at ~ 1.5 and ~ 5 km. The ozone density above the upper stable layer seems to be separated from that below it and is controlled by a different mechanism. L. Y. Chan et al. (2000) and C. Y. Chan et al. (2003) stated that the lower stable layer isolates the layer with the ozone increase at ~ 3 km from the influence of the local boundary layer. We add that the horizontal advection of clean, oceanic air associated with the cold surge passage is essential for the isolation.

Recent studies suggested that stratospheric ozone intrusion plays an important role in tropospheric ozone variations including the spring ozone enhancement in spring above Hong Kong (Liao et al., 2021; Zhao, Hu, et al., 2021; Zhao, Huang, et al., 2021). Although we cannot discuss quantitatively the stratospheric contribution to the spring ozone enhancement above Hanoi, because the experiment in this study tested the sensitivity to the surface emissions of ozone precursors, a certain amount of stratospheric ozone can contribute to the spring ozone enhancement above Hanoi. However, if we

look at the spatial distribution as seen in Figures 12 and 13, the observed ozone enhancement in the lower troposphere above Hanoi is clearly connected with the surface emission over the Indochina Peninsula. In any case, the geographical difference between Hong Kong and Hanoi can lead to different stratospheric contributions. The 3D transport process should be clarified in more detail in future investigations to understand the relative importance of stratospheric intrusion.

7. Conclusions

Regular ozonesonde observations for about 14 years above Hanoi revealed an ozone increase at ~ 3 km in March. The ozone densities in March showed large temporal variability at three height ranges that are separated by two stable layers at ~ 1.5 and ~ 5 km. Meteorological and backward trajectory analyses showed that a typical ozone enhancement at ~ 3 km originated from air polluted over the Indochina Peninsula and a typical decrease below 1.5 km was caused by the advection of the clean oceanic air associated with a cold surge event.

We conducted a sensitivity experiment using a chemical transport model, in which we eliminated the emissions of ozone precursors (CO and NO_2) in three regions (the Indian Subcontinent, the northern Indochina, and southern China). The ozone concentration above Hanoi was most effectively suppressed when emissions of ozone precursors over northern Indochina Peninsula are eliminated. This suggests that air pollution from northern Thailand (probably due to biomass burning) contributed most to the ozone increase in Hanoi. The model showed that the ozone increase originated from the northern Indochina Peninsula emission expanded in the lower troposphere over an area of about 20° in longitude and latitude. The model also reproduced the ozone decrease near the surface due to the clean air intrusion associated with the winter monsoon easterly wind. The polluted air was further transported eastward into the middle and upper troposphere across the Pacific Ocean, and some of it reached the west coast of the USA.

We propose that the ozone increase at 3 km over Hanoi in March is caused by the eastward advection of polluted, high-ozone air that was well mixed up to the stable layer at ~ 5 km over the land mass of the Indochina Peninsula to the west of Hanoi, and by the westward advection of less-polluted, low-ozone air from the oceanic area to the east of Hanoi associated with a cold surge event below the stable layer at ~ 1.5 km. Such a mechanism occurs only between March and April, after the development of active shallow convection and before the start of summer monsoon rainfall. We conclude that the ozone enhancement over Southeast Asia is caused not only by the biomass burning enhancement, but also by atmospheric circulation system formed in pre-monsoon season. The circulation system determines the detailed 3D structure of ozone distribution.

Data Availability Statement

The ozonesonde data used in this study are available in the SHADOZ archive <https://tropo.gsfc.nasa.gov/shadoz/> (SHADOZ, 2021). JRA-55 data can be obtained from the Japan Meteorological Agency http://jra.kishou.go.jp/JRA-55/index_en.html (Japan Meteorological Agency, 2018). The radiosonde data at the Hanoi station are available in the archive site of University of Wyoming <http://weather.uwyo.edu/upperair/sounding.html> (Wyoming University, 2018). The data of the numerical experiment presented in Section 5 are available at <https://doi.org/10.5281/zenodo.6459880> (Miyazaki, 2022).

Acknowledgments

This work was supported by JSPS KAKENHI Grant Nos. JP15204043, JP18204041, JP21244072, JP26220101, and JP16K00535. The SHADOZ sondes at Hanoi are supported by NASA's Upper Atmosphere Research Program through funding to Goddard Space Flight Center (Maryland USA) and the Ozone group at NOAA's Global Monitoring Lab (Colorado USA); special thanks to Patrick Cullis (CIRES at NOAA). Part of this work was conducted at the Jet Propulsion Laboratory, California Institute of Technology, under contract with the National Aeronautics and Space Administration (NASA).

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