SMILES measurements of diurnal variations of hydroperoxyl radical (HO$_2$) in the stratosphere and mesosphere

Nao Suzuki$^1$*, Hideo Sagawa$^2$, Yukio Nakano$^1$, Nori Mizuno$^2$, Yasuko Kasai$^2$

$^1$Tokyo Gakugei University, $^2$NICT

1 Introduction

HO$_2$ radical is among the most important oxidants for atmospheric chemical compositions in the upper atmosphere. The accurate measurement of HO$_2$ turned out to be an extremely difficult problem because HO$_2$ volume mixing ratios are about a few parts per billion by volume. No significant measurement was reported so far to evaluate the atmospheric chemistry of HO$_2$ such as its diurnal variations and behaviors in stratosphere and mesosphere.

In this study, we report the first wide altitude range observations from stratospheric to mesospheric HO$_2$ diurnal variations, the measurements of which were previously considered to be difficult, by Superconducting Submillimeter-Wave Limb-Emission Sounder (SMILES). We discuss hydrogen chemistry in the stratosphere and mesosphere.

SMILES is a part of Japanese Experiment Module (JEM) onboard International Space Station (ISS).

2 Dataset and Analysis of SMILES HO$_2$ measurements

ISS which has SMILES platform orbits at an altitude of 320-340 km above the surface and completes a single earth orbit in about 90 minutes. SMILES observed about 1630 points per day over the Earth during September 2009-April 2010, and produce a global map everyday for each composition. The latitudinal range of the observation covers from 65° N to 38° S.

SMILES detects thermal emission from chemical substances contained in the atmosphere and obtains the emission signal from atmospheric compositions at the several height in the atmosphere. HO$_2$ transition frequency is 649.70 GHz.

We have used HO$_2$ data from SMILES NICT products. We selected the data into three parts of latitude. These parts are equator (20° N-20° S), mid-latitude (20° N-50° N) and north polar (50° N-65° N), respectively. And we selected HO$_2$ vertical profiles at 20-95 km at night and in the daytime and HO$_2$ diurnal variations as a function of solar zenith angle (SZA) at 10° intervals in the stratosphere (29.0-49.0 km) and mesosphere (53.0-74.5 km), in each region of latitude.

3 Results and Discussions

HO$_2$ vertical profiles at 20-95 km are obtained. HO$_2$ volume mixing ratios generally increase with the altitude above 20 km and the appearance of the maximum (peak) volume mixing ratio in the profile is evident in the mesosphere, near 79.5 km at about 3ppbv at night and near 74.5 km at about 5.5ppbv in the daytime. We confirmed that the altitude of daytime peak in the shape of HO$_2$ profiles is lower than that of nighttime peak in all parts of latitude.

The diurnal variation of HO$_2$ through in the stratosphere and mesosphere was obtained for the first time (Figure 1). HO$_2$ volume mixing ratios at the all altitude are enhanced during the daytime due to photochemical reaction. The largest source of HOx (= H + OH + HO$_2$) radicals in the stratosphere are provided from reaction O(1D) atoms, which are generated predominantly from O$_3$ photolysis, with H$_2$O.

The increase of HO$_2$ mixing ratios in the stratosphere with altitude can largely be attributed to an increased formation rate due to the large increase in the abundance of O(1D) at high altitude in the stratosphere. On the other hand, the production processes for mesospheric HOx during the daytime involve the photodissociation reaction of water vapor, which plays an important role in HO$_2$ production additionally in the mesosphere.

Acknowledgements: Data processing and other research works in the present study was performed with the NICT Science Cloud at National Institute of Information and Communications Technology (NICT) as a collaborative research project.

Keywords: oxidant, HO$_2$, hydroperoxyl radical, diurnal variation, stratosphere, mesosphere
Figure 1. HO\textsubscript{2} diurnal variations from SMILES observations for equatorial region in the stratosphere (left panel) and mesosphere (right panel) at the altitude from 29.0 km up to 74.5 km.
Impacts of the Arctic ozone depletion on Japan observed with FTIR between 2009 and 2011

Yuki Hashimoto¹*, Isao Murata¹, Hideaki Nakajima², Isamu Morino²

¹Tohoku University, ²National Institute for Environmental Studies

The ozone depletion is one of the environmental problems. In 2011, ozone depletion which was comparable to the Antarctic ozone hole occurred in the Arctic.

The ozone depletion has occurred inside the polar vortex. The airmass from inside the polar vortex is spread to mid-latitude in spring after its breakup. The purpose of this study is to quantify the impact of polar ozone depletion on mid-latitude by comparing the amounts of ozone in mid-latitude airmasses before and after the breakup of the polar vortex.

Vertical profiles of ozone and hydrogen fluoride (HF) have been retrieved from infrared spectra observed with a Fourier transform infrared spectrometer (FTIR) at Tsukuba using the SFIT2 spectral fitting program developed by Rinsland et al. [1998].

HF can be used as a tracer of the transport, because HF is a remarkably stable species in the stratosphere. Ozone and HF usually show a very high correlation in the lower stratosphere because both species are stable. But the correlation will be changed when ozone is chemically perturbed. Therefore, we examined the correlations of mixing ratios between ozone and HF in the mid-latitude airmasses before and after the breakup.

Goto et al. [2010] compared the ozone-HF mixing ratio correlations in the mid-latitude airmasses before and after the breakups of the polar vortex between 2005 and 2008 and indicated that ozone was decreased by 0.2 to 0.4 ppmv at around 19km altitude in 2007 and 2008. Preliminary analysis in 2009 shows no impact of the ozone depletion at Tsukuba. We report the impacts of the Arctic ozone depletion on Japan between 2009 and 2011.
Simulation study of synergetic retrieval for tropospheric ozone with UV, TIR, and MW measurements

Takao M. Sato1, Hideo Sagawa2, Hitoshi Irie3, Katsuyuki Noguchi4, Naoko Saitoh1, Ryoichi Imasu5, Yoji Hayashi5, YASUKO KASAI2

1Institute of Space and Astronautical Science, Japan Aerospace Exploration Agency, 2National Institute of Information and Communications Technology, 3Center for Environmental Remote Sensing, Chiba University, 4Nara Women’s University, 5CCSR, The University of Tokyo

Tropospheric ozone, one of short-lived climate pollutants (SLCPs), plays an important role on climate change, atmospheric chemistry, and air quality. First, ozone acts a greenhouse gas, especially in the upper troposphere (UT) and its effect is the strongest near the tropopause where the climate system is more sensitive. Second, ozone in the planetary boundary layer (PBL) is known to be a major component of photochemical smog and causes severe damages the health of both plants and animals. Finally, ozone has an oxidizing capability to remove many pollutants (e.g., methane and carbon monoxide) from the atmosphere. In order to understand these processes of tropospheric ozone with the aid of numerical model calculations, it is essential to monitor the global distribution of tropospheric ozone with a fine vertical resolution. However, current available remote sensing instruments, which measure backscattered solar UV radiance (e.g., Aura/OMI) and thermal infrared emission (e.g., Aura/TES), cannot resolve tropospheric ozone alone.

For the purpose of improving the current capability of tropospheric ozone retrieval, we have proposed a new atmospheric remote sensing equipment named Air Pollution Observation (APOLLO) to be carried aboard the Japanese Experiment Module (JEM) on the International Space Station (ISS). APOLLO is planned to carry two nadir-viewing passive instruments (UV+VIS and TIR) and one limb-viewing microwave passive instrument (MW), which are dedicated to measure tropospheric ozone and its precursors relating to air quality with high spatial resolution (~2 km) and high vertical resolution (the requirement is to divide troposphere into three layers: UT, LT (lower troposphere), and PBL).

In this study, we investigate synergetic effect of combination of UV, TIR, and MW measurements on retrieval sensitivity of tropospheric ozone. We evaluate the retrieval sensitivity to tropospheric ozone profile with Optimal Estimation Method (OEM) [Rodgers, 2000]. Twenty atmospheric scenarios generated from a global-regional chemical transport model system are used as true profiles for this aim. The combination of UV and TIR measurements improves values of the degree of freedom for signal (DOFS) in PBL. MW limb measurements provide significant information on UT and the stratosphere. We find that adding MW limb measurements to UV+TIR measurements significantly improve values of DOFS in PBL, although MW limb measurements alone are not sensitive to PBL. We conclude that this is because UV+TIR measurements become more sensitive to PBL by adding MW limb measurements which can reduce the uncertainty of ozone concentration in UT compared with that from UV+TIR measurements.

Keywords: International Space Station, Tropospheric ozone, Multispectral observations
Improvement of retrieval algorithm of CO2 and CH4 profiles from GOSAT/TANSO-FTS TIR band

Naoko Saitoh1*, IMASU, Ryoichi2

1Center for Environmental Remote Sensing, Chiba University, 2Atmosphere and Ocean Research Institute, The University of Tokyo

The Greenhouse Gases Observing Satellite (GOSAT), which was developed by the National Institute for Environmental Studies (NIES), the Ministry of the Environment (MOE), and the Japan Aerospace Exploration Agency (JAXA), has been observing greenhouse gases continuously for about four years since its launch on January 23th, 2009. GOSAT consists of four spectral bands: three of the four bands in shortwave infrared (SWIR) region and one in thermal infrared (TIR) region. GOSAT can simultaneously observe CO2 and CH4 column-averaged dry-air mole fractions and their profiles in the same field of view from the SWIR and TIR bands. TIR Version 00.01 Level 2 (L2) data have been released to the public. The V00.01 L2 data were processed with a previous version (V100.100) of Level 1B (L1B) data. Although a single TIR V00.01 L2 CH4 profile does not have enough quality for scientific use, the TIR column-averaged CH4 value agrees to aircraft CH4 data within 0.5% [Saitoh et al., 2012]. On the other hand, TIR V00.01 L2 CO2 data do not have enough quality; they show a relatively large variability and have a clear bias in mid-troposphere in low latitudes. This study improves an algorithm for retrieving CO2 and CH4 profiles from TIR band. The new algorithm simultaneously retrieves several parameters other than target gases such as CO2 and CH4. The TIR retrieval processing in this study uses the latest version of L1B data; therefore, L2 data bias attributed to L1B spectral bias would be expected to decrease.

Keywords: greenhouse gas, satellite remote sensing, retrieval algorithm
The temporal variation of vertical profile of methane at Poker Flat observed by Fourier transform spectrometer

Koichi Maruno\textsuperscript{1}, Isao Murata\textsuperscript{2*}, YASUKO KASAI\textsuperscript{3}, Kagwa Akiko\textsuperscript{4}, Yasumasa Kasaba\textsuperscript{1}

\textsuperscript{1}Graduate school of science, Tohoku University, \textsuperscript{2}Graduate school of environmental studies Tohoku University, \textsuperscript{3}National Institute of Information and Communications Technology, \textsuperscript{4}FUJITSU FIP CORPORATION

Methane is the second important greenhouse gas. In the arctic region, there are many sources of the atmospheric methane, i.e., wetlands, permafrosts and natural gas fields. It is important to investigate the trend of the methane in the arctic region.

In this study, we analyzed the trend of the vertical profile of methane at Poker Flat (Latitude: 65.11 N, Longitude: 147.42 W) observed with FTIR (Fourier Transform Infrared Spectrometer) by NICT from 2000 to 2010 (except for 2006). The vertical profiles of methane were derived with the SFIT2 spectrum fitting program using Rodgers’ Optimal Estimation Method (OEM).

We investigated temporal variation of the columns in the lower stratosphere (10-21 km), troposphere (0-10 km), and total (0-100 km) which calculated from the vertical profiles with digital fitting method [Nakazawa et al., 1997]. Lower stratospheric column shows no significant trend. Tropospheric and total columns show increase in 2008 then decrease in 2009. This trend in the troposphere is different from those observed in Germany, East Asia and Global average which don’t show decrease after 2008. The difference of the trend indicates that observations at various regions are important in order to understand the spacial and temporal variations of sources and sinks of methane.

Keywords: methane, FTIR (Fourier transform Infra-Red spectrometer)
Optimization of wavenumber regions for the retrieval of the vertical profiles of CH$_4$ from infrared spectra

Isao Murata$^1$, Hideaki Nakajima$^2$, Isamu Morino$^2$

$^1$Graduate School of Environmental Studies, Tohoku University, $^2$National Institute for Environmental Studies

CH$_4$ is the second important greenhouse gas but there is large variability of its increasing rate that may be due to variabilities of sources. Therefore, it is important to investigate the vertical profile of CH$_4$.

Solar infrared spectra have been observed with a Fourier transform spectrometer (FTS) at Tsukuba, Japan. FTS has advantages in its high-resolution and the wide wavenumber range. Vertical profiles of some species can be derived from the high-resolution spectra with the SFIT2 spectral fitting program developed by Rinsland et al. (1998). It needs to select appropriate wavenumber regions and the optimization of fitting parameters is also needed. Now we are investigating these wavenumber regions and parameters in the NDACC/IRWG group for the retrieval of the vertical profiles and column densities of CH$_4$. Sussmann et al. [2011] analyzed with some combinations of the following wavenumber regions: 1) 2613.7 - 2615.4 cm$^{-1}$, 2) 2650.6 - 2651.3 cm$^{-1}$, 3) 2835.5 - 2835.8 cm$^{-1}$, 4) 2903.6 - 2904.03 cm$^{-1}$, 5) 2921.0 - 2921.6 cm$^{-1}$, and reported the combination of 1), 3), and 5) is best. We also compared the results from some combinations of these wavenumber regions and found that the discrepancy become large in summertime. It may be due to HDO absorption lines existing in these wavenumber regions as an interfering species.

Keywords: FTIR, Trace Species, Methane
Temperature from GPS RO meas. correlative to satellite and airborne obs. for comparing those CH4 profiles

Takafumi Sugita\textsuperscript{1}\textsuperscript{*}, Naoko Saitoh\textsuperscript{2}, Sachiko Hayashida\textsuperscript{3}

\textsuperscript{1}NIES, \textsuperscript{2}Chiba Univ., \textsuperscript{3}Nara Women’s Univ.

Evaluations of data quality of CH\textsubscript{4} retrieved from satellite-borne nadir sensors, from which are ADEOS/IMG in 1996-97, MetOp-A/IASI from 2007, and so on, in the troposphere and stratosphere have been performed in recent years (e.g., Clerbaux et al., ACP, 2003; Xiong et al., Remote Sens., 2010; Wecht et al., ACP, 2012; Razavi et al., ACP, 2009). GOSAT/TANSO-FTS started its operation from 2009, expected contributions in this area. For retrieving CH\textsubscript{4} profiles, it is necessary to input several external parameters such as temperature profiles, surface temperature, emissivity, and so on. A purpose of this paper is to understand the effect of temperature profiles on the CH\textsubscript{4} retrieval in the thermal infrared band of TANSO-FTS. So that, we will prepare temperature profiles from GPS radio occultations (RO). We will focus on GOSAT observations in the northern high-latitudes, where aircraft observations have been done by National Institute for Environmental Studies with the aid of Russian Academy of Science. We extracted data from coincidences between two Russian aircraft sites and GOSAT since 2009. Then, the RO temperatures for those pairs are prepared and compared with some meteorological datasets.

Keywords: temperature, methane, GPS, GOSAT, aircraft
Shipboard measurements of atmospheric CH4, CO2 and CO mixing ratios during the MR12-E03 cruise of the R/V Mirai

Yasunori Tohjima1, SASANO, Daisuke2, ISHIDOYA, Shigeyuki3, KATSUMATA, Keiichi1, MATSUSHITA, Junji1, Kentaro Ishijima4, Prabir Patra4

1National Institute for Environmental Studies, 2Meteorological Research Institute, 3National Institute of Advanced Industrial Science and Technology, 4Japan Agency for Marine-Earth Science and Technology

In order to investigate the potential sources of methane (CH4) in the Arctic region, continuous measurements of the atmospheric CH4 were carried out during a R/V Mirai Arctic Ocean cruise from September 3 to October 17, 2012. A cavity ring-down spectroscopy (CRDS) analyzer was used for the shipboard measurements of the atmospheric CH4, carbon dioxide (CO2) and carbon monoxide (CO). The analytical precisions evaluated from the measurements of the standard gases at a 24-hour interval during the cruise were 0.02 ppm, 0.3 ppb, and 0.9 ppb for the 5-min averages of CO2, CH4, and CO mixing ratios, respectively. When the wind blew from the relative direction of 200 +/- 20 degrees (rear left of the vessel), the contamination caused by its own exhaust fumes affected the CO2 and CO mixing ratios with a tight correlation (Delta_CO/Delta_CO2=3.8 ppb/ppm), while there was no significant influence from the exhaust fumes on the CH4 mixing ratio. Such pollution events are easily distinguishable by the characteristics of the relative wind direction, the tight correlation of CO vs. CO2, and large short-term (a few second) variability. The observed CH4 mixing ratios showed larger variations with elevated peaks of several tens ppb in the Bering Strait, Chukchi Sea, and Arctic Ocean (65-75°N, 155-175°W) in comparison with in the western North Pacific. The largest CH4 peaks of about 50 ppb were observed off the northern Alaskan cost. Since these CH4 peaks were associated with similar CO2 peaks but not with CO peaks, it is unlikely that the combustion processes or ocean were the sources of the elevated CH4. The backward trajectory analysis suggests that the North Slope of Alaska is the most probable CH4 source region. The simulated CH4 variations based on an atmospheric transport model and given flux maps well capture the observed CH4 variations, also suggesting that the most of elevated CH4 were derived from the land sources.

Keywords: atmospheric CH4, the Arctic Ocean, cavity ring down spectroscopy analyzer (CRDS), shipboard measurements
Carbon monoxide and ozone measurements during summertime at the summit of Mt. Fuji

Shungo Kato¹*

¹Tokyo Metropolitan University

The top of Mt. Fuji is 3776 m and it located in free troposphere. Mt. Fuji weather station is a unique observatory for atmospheric measurements of free troposphere. But now the weather station is only open during summer. At the summit of Mt. Fuji, O₃ has been observed from 2007, and CO has been observed from 2008 during summer season.

CO and O₃ are monitored by Thermo Environmental Instrument Model 48C and 49i, respectively. Before and after the summertime intensive measurements, these instruments were calibrated by standard gas. Since CO analyzer is influenced by temperature and water vapor concentration, zero air produced by heated Pt catalyst was measured periodically.

Observed CO and O₃ concentrations showed large variation compared to other remote sites. Basically, CO and O₃ showed similar concentration change because clean air with low concentration and polluted air with high concentration are arrived to Mt. Fuji time to time. When only CO was high, polluted air experienced less photochemical activity was arrived. When only O₃ was high, air from upper troposphere with high O₃ was arrived. Scatter plot of CO and O₃ was categorized by water vapor. Clear trend of low water, high O₃ and low CO was observed.

Low water vapor but high CO was observed in some case. It is expected that polluted air was lifted up and transported to the observatory.

Diurnal variation of CO and O₃ were not observed clearly. The influence of mountain wind is not important at least for CO and O₃.

The average concentrations of CO and O₃ for each year have large difference year to year. Influence of clean ocean air mass and polluted continental air mass will be different for each year and it affected the average concentration of pollutants during summer.

Keywords: free troposphere, longrange transport, mountain site, high altitude
A decadal inversion of carbon dioxide using the Global Eulerian-Lagrangian Coupled Atmospheric model (GELCA)

Tomoko Shirai¹, Misa Ishizawa¹, Ruslan Zhuravlev², Alexander Ganshin², Tazu Saeki¹, Dmitry Belikov¹, Tomohiro Oda³, Makoto Saito⁴, Vinu Valsala⁵, Shamil Maksyutov¹

¹National Institute for Environmental Studies, ²Central Aerological Observatory, ³NIES, now at CSU/NOAA ESRL, ⁴NIES, now at LSCU, ⁵NIES, now at ITTM

A decadal estimate of global CO₂ flux distribution for the period of 2001-2010 was conducted using an atmospheric inversion modeling system called GELCA (Global Eulerian-Lagrangian Coupled Atmospheric model) with Kalman smoother inversion technique. The use of Lagrangian particle dispersion model (LPDM) to simulate the transport in the vicinity of the observation points enables us to avoid numerical diffusion from which Eulerian models suffer, and is suitable to represent observations at high spatial and temporal resolutions. An Eulerian model is run to generate the global background concentrations to be used as the boundary conditions for an LPDM that performs backward simulations from each receptor point (observation location). In GELCA, National Institute for Environmental Studies-Transport Model (NIES-TM) version 8.1i was used as an Eulerian global transport model coupled with FLEXPART version 8.0 as a LPDM. Two-day backward transport by FLEXPART was combined with the background CO₂ levels 2 days prior to the observations simulated by NIES-TM. The meteorological data for driving both models was taken from JMA Climate Data Assimilation System (JCDAS) with a spatial resolution of 1.25° x 1.25° and a temporal resolution of 6 hours. Our prior CO₂ fluxes consist of the following four types: daily terrestrial biospheric fluxes generated by the VISIT model (Vegetation Integrative Simulator for Trace gases); monthly oceanic fluxes generated by an ocean pCO₂ data assimilation system; monthly biomass burning fluxes taken from the Global Fire Emissions Database (GFED), version 3.1; and monthly fossil fuel emissions combining the high-resolution Open source Data Inventory of Anthropogenic CO₂ emission (ODIAC) version 3.0 dataset. We employed a Kalman Smoother inversion technique with fixed lag of 3 months, solving for 42 land and 22 ocean regions.

The purpose of the present study is to evaluate the performance of the GELCA inversion system with rather long period (10 years) CO₂ flux estimation and to examine the impact of observation network. We tested several different sets of observation datasets starting by using the NOAA flask network ground based observations as a control case. The sensitivity of the inversion to the choice of CO₂ observation dataset was discussed using the footprint of each observation dataset. The CO₂ flux estimate was examined in terms of observation network/coverage and also compared with previous studies.

Keywords: CO₂, sources/sinks, inverse modeling, coupled model
Trends and seasonal cycle of atmospheric radiocarbon in carbon dioxide observed at Hateruma Island

Yukio Terao\textsuperscript{1*}, Hitoshi Mukai\textsuperscript{1}

\textsuperscript{1}Center for Global Environmental Research, National Institute for Environmental Studies

We have been conducted monthly air samplings for measurements of atmospheric radiocarbon in carbon dioxide (\textsuperscript{14}CO\textsubscript{2}) at Hateruma Island (HAT, 24.05\textdegree N, 123.80\textdegree E, 47 m a.s.l.), Japan since 2004. We collected whole air samples using 2.0L glass flasks pressurized to 3 atm, and 5L air was used for radiocarbon analysis. The values of Delta\textsuperscript{14}C were measured using compact Carbon Accelerator Mass Spectrometry (CAMS, NEC 1.5SDH). Uncertainty in Delta\textsuperscript{14}C measured by CAMS is less than 2 per mil, which is based on the number of \textsuperscript{14}C counts and the scatter of \textsuperscript{14}C/\textsuperscript{12}C ratios during measurements. The reproducibility of CAMS measurements is +/-1.4 per mil (standard deviation of Delta\textsuperscript{14}C values in a reference air cylinder).

Here we show the Delta\textsuperscript{14}C values of background maritime air observed at HAT from 2004 to 2012. The seasonal cycle of Delta\textsuperscript{14}C was observed: minimum in winter-spring and maximum in summer, with amplitude of 10 per mil. Decreasing trends in Delta\textsuperscript{14}C were from -5 to -6 per mil/year, however, higher growth rates (less decreasing trends) of -2 per mil/year were observed in 2008-2009. The reason for the IAV in Delta\textsuperscript{14}C will be discussed.

Keywords: carbon cycle, carbon isotope measurements, accelerator mass spectrometry
Long-term measurements of black carbon concentrations in rainwater at a remote site in East Asia

Tatsuhiro Mori\textsuperscript{1*}, Sho Ohata\textsuperscript{1}, Yutaka Kondo\textsuperscript{1}, Nobuhiro Moteki\textsuperscript{1}, Hitoshi Matsui\textsuperscript{1}, Aya Iwasaki\textsuperscript{2}, Nobutaka Tomoyose\textsuperscript{2}, Hisashi Kadena\textsuperscript{2}

\textsuperscript{1}Graduate School of Science, University of Tokyo, \textsuperscript{2}Okinawa Prefectural Institute of Health and Environment

Black carbon (BC) particles are mainly emitted into atmosphere by incomplete combustion of fossil fuels and biomass. BC particles emitted from these sources are generally hydrophobic, but they are gradually coated by hygroscopic species during transport and become hydrophilic particles with higher cloud condensation nuclei (CCN) activity, and they are finally removed from the atmosphere through wet removal processes. Therefore, wet deposition of BC is one of the most important processes controlling BC concentrations and their distribution. In order to understand the importance of wet removal processes of BC, BC mass concentrations both in the surface air ($m_{air}$) and in rainwater ($m_{rain}$) were measured simultaneously for two years (April 2010 - March 2012) at Cape Hedo in Okinawa. This study is the first attempt to conduct long-term measurements of $m_{air}$ and $m_{rain}$ at a remote site in East Asia. Seasonal variations of both concentrations and the wet deposition flux of BC are presented.

A Continuous Soot Monitoring System (COSMOS) was used to measure $m_{air}$. Rainwater samples were collected on a daily basis and $m_{rain}$ was measured by an ultrasonic nebulizer (U-5000AT) and a Single Particle Soot Photometer (SP2).

The measured $m_{air}$ and $m_{rain}$ showed clear seasonal variations. The monthly mean $m_{air}$ and $m_{rain}$ were the highest during spring (March - May) with the values of 0.37 \text{ug m}^{-3} and 62.2 \text{ug L}^{-1}, and the lowest during summer (June - August) with the values of 0.07 \text{ug m}^{-3} and 5.82 \text{ug L}^{-1}. The correlation coefficient between the monthly mean $m_{air}$ and $m_{rain}$ was sufficiently high ($r^2 = 0.67$). The annual average amount of BC wet deposition, which was defined as the product of $m_{rain}$ and precipitation amount, was 39.6 \text{mg m}^{-2} during the observation periods. The contribution to the total amounts of BC wet deposition was found to be 74.4\% during spring. This is due to both higher $m_{rain}$ and larger precipitation amount in spring. The values of $m_{air}$ and $m_{rain}$ in winter (December - February) and spring are the highest because polluted air masses are frequently transported from the Asian continent by strong north-westerly wind, East Asian monsoon in winter and cold front passages in spring. On the other hand, these values are the lowest in summer because clean air masses are transported from south to the measurement site by the Pacific high.

Keywords: Black Carbon, wet deposition
Number-size distribution of maritime aerosol particles over the Pacific Ocean

Sayako Ueda¹, Kazuhiko Miura¹, Ryou Kawata², Hiroshi Furutani², Mitsuo Uematsu²

¹Tokyo University of Science, ²AORI, University of Tokyo

Size and number concentration of atmospheric aerosol particles are the most fundamental parameters for estimating effects of aerosol on climate. Number-size distributions of aerosol in 10-500 nm diameters were observed on board the R/V Hakuho-Maru cruises over the Pacific Ocean during December 2011?March 2012. The KH-11-10 cruise started from Tokyo and reached Peru via Hawaii and the mid-latitudes eastern South Pacific Ocean. The KH-12-1 cruise started from Peru and reached Tokyo via the eastern equator and Hawaii. Number-size distribution of dried aerosol particles was measured using a Scanning Mobility Particle Sizer (3034, TSI Inc.) for diameters of 10-500 nm and a laser particle counter (LPC, KC01D; RION Co. Ltd.) for diameters greater than 300 nm. The obtained number-size distributions were analyzed to reveal their relationship with the condensation sink of precursor gases, air mass transport, meteorological condition, and chlorophyll concentration along trajectory.

Bimodal size distribution with mode peaks in 30-80 nm (Aitken mode) and 100-200 nm (accumulation mode) was frequently observed. Relatively large mode sizes were observed over the western equator. The 5 days backward trajectory shows that the air masses in equator are originated from the high chlorophyll area without experiencing precipitation. Over the mid-latitudes in the eastern South Pacific where chlorophyll concentration is low, new particle formation event was often observed in accordance with the low concentration of accumulation mode particles.

In this observation, new particle formation event was not observed under high chlorophyll condition over the equator. Condensation of precursor gases onto pre-existing particles strongly prevents nucleation of fresh particles. Because such condensation to pre-existing particles was effective enough in the equator under non-precipitating condition, new particle formation could not be observed over the equator. These results suggested that precipitation and biological productivity controlled the balances of formation and growth of aerosol particles, characterizing the number size distribution over each ocean areas.

Keywords: Maritime aerosol, Number-size distribution, New particle formation
Atmospheric pollutants originated Asian Continent included PM in Yakushima Island.

Osamu Nagafuchi1*, Kuriko Yokota2, Kenshi Tezuka3, Mayumi Jige4, Koyomi Nakazawa5

1The University of Shiga Prefecture, 2Toyohashi University of Technology, 3Yattane goyou team, 4Ootani University, 5Osaka University

Airborne particulate matter (PM) is a complex mixture of particles that are very different in size, chemical composition, physical state and morphology. Moreover, PM has a variety of emission sources which range from natural to anthropogenic and stationary to mobile. It also has a variety of physical and chemical properties. Therefore, not only the size distribution of particles but also information related to their chemical composition will play an important role in elucidation of the behavior and major emission sources of PM and their effect on human health and the ecosystem.

The samples of size-resolved PM were collected using a 3-stage NLAS impactor (Tokyo Dyrec Co., Ltd., particle cut size of stage is 10mm, 2.5 mm and 1.0mm for a flow rate of 3 L/min) with one day or three days sampling interval on the polycarbonate filter (25mmf) and a polycarbonate filter (back-up filter 47mmf). Sampling of the PM was conducted at Yakushima Island, from 13 to 26 Jan., 2013. Elemental compositions of these samples were determined by ICP/MS, and ionic species were analyzed by IC. In addition, we observed a morphology by SEM. Moreover ATR-FT-IR imaging measurements of individual particles were performed using a Perkin-Elmer Spectrum 100 FT-IR spectrometer interfaced to a Spectrum Spotlight 400 FT-IR microscope.

Keywords: particulate matter, long-range transport, major ion, heavy metal
Atmospheric Fe-containing particles over the North Pacific Ocean: the mixing states with water-soluble materials

Yusuke Miki\textsuperscript{1,}\textsuperscript{*}, Sayako Ueda\textsuperscript{1}, Kazuhiko Miura\textsuperscript{1}, Hiroshi Furutani\textsuperscript{2}, Mitsuo Uematsu\textsuperscript{2}

\textsuperscript{1}Tokyo University of Science, \textsuperscript{2}University of Tokyo

Fe is an essential element for marine phytoplankton growth. Long-range transportation of atmospheric aerosols from the continent and subsequent deposition is an important process to supply Fe to the ocean. The dry and wet depositions of aerosol particles depend on the particle size and the mixing states with water-soluble materials. In order to study the mixing states of Fe-containing particles with water-soluble materials, we collected aerosol particles on the ship over the mid-latitude western North Pacific Ocean during the KH-12-1 (EqPOS) Leg 2 cruise of the R/V Hakuho Maru, Atmosphere and Ocean Research Institute (AORI), University of Tokyo. The leg started from Hawaii on February 21, 2012, and arrived at Tokyo on March 7, 2012. We collected aerosol particles with a low pressure impactor. Collected particles were analyzed using a transmission electron microscopy (TEM) with a water dialysis method. Most of maritime aerosols are consisted of water-soluble materials such as sea-salt and sulfate particles. Water-insoluble materials such as minerals and industrial metals are main sources of Fe. This study focused on water-insoluble materials and performed an energy-dispersive X-ray (EDX) analysis.

We classified the origins of encountered air masses on the basis of backward air trajectory analysis and number concentration of aerosols measured by an optical particle counter (OPC). Particles larger than D = 0.5 um during dust events and background conditions (maritime or continental origins) were analyzed using the TEM. Number fractions of particles containing water-insoluble materials were 5-20\% (0.5-1.0 um in diameter) and 15-50\% (>1.0 um in diameter). Most of water-insoluble materials were mixed with water-soluble materials (mixed particles). Median values of the volume percent of the water-soluble materials in the mixed particles on a maritime sample were >90\% and those on the other samples were 60-80\%. Based on EDX analyses of water-insoluble materials, number fractions of Fe-containing particles were 2\% (maritime), 2\% (continental) and 8\% (dust event), respectively. Fe-containing water-insoluble materials were found with other mineral components (Si or Al), and mixed with water-soluble materials.

Since phytoplankton growth requires dissolved form of Fe, we performed EDX analyses for the same particles before and after extractions of water-soluble materials from particles to estimate fraction of water-soluble Fe. The amount of Fe after the extraction was smaller comparing with the amount of Fe before the extraction, indicating that the water-soluble Fe is presented with the water-insoluble Fe.

Keywords: Fe, water-soluble materials, water-insoluble materials, dust, aerosol
Evaluation of the method to measure black carbon particles suspended in rainwater and snow samples

Sho Ohata1*, MOTEKI, Nobuhiro1, SCHWARZ, Joshua P.2, FAHEY, David W.2, Yutaka Kondo1

1Department of Earth and Planetary Science, Graduate School of Science, The University of Tokyo, 2Earth System Research Laboratory, National Oceanic and Atmospheric Administration

The mass concentrations and size distributions of black carbon (BC) particles in rainwater and snow are important parameters for improved understanding of the wet deposition of BC. In this study, we have made a detailed evaluation of the method to measure these parameters. The method consists of an ultrasonic nebulizer (USN) and a Single Particle Soot Photometer (SP2). The USN converts sample water into micron-size droplets at a constant rate and then extracts BC particles to air by drying the water droplets. The mass of individual BC particles is measured by the SP2, based on the laser-induced incandescence technique. The loss of BC particles during the extraction process from liquid water to air depends on their sizes. We determined the size-dependent extraction efficiency using polystyrene latex spheres (PSLs) with twelve different diameters between 107-1025 nm. The PSL concentrations in water were measured by the light extinction at 532 nm. The extraction efficiency of the USN showed broad maximum of about 10% in the diameter range of 200-500 nm, and decreased substantially at larger sizes. Total uncertainty and reproducibility of the measured mass concentration of BC in sample water were +40% and +35%, respectively. Measured BC size distributions in rainwater and surface snow collected in Tokyo, Okinawa, and Sapporo showed negligible contribution of the BC particles larger than 600 nm to the total BC amounts. However, surface snow collected in Greenland sometimes contained significant amount of larger BC particles, beyond the upper detection limit of the present method.

Keywords: black carbon, wet deposition
Emission of biogenic VOCs from evergreen broadleaf tree: variation in composition of monoterpenes

Sou Matsunaga1*, TAKAGI, Masahiro2, KUSUMOTO, Dai3, HIURA, Tsutom1

1Tomakomai Research Station, Hokkaido University, 2Faculty of Agriculture, University of Miyazaki, 3The University of Tokyo Tanashi Forest, the University of Tokyo

Introduction

Biogenic volatile organic compound (biogenic VOC, BVOC) is known to have an important role on atmospheric chemistry in both regional and a global scale. BVOCs consist of many classes of organic compounds such as isoprene (C5H8), monoterpenes (MNTs: C10H16) and others. Although emission pattern of isoprene, which has the largest emission rate, is relatively well modeled as a result of many efforts, that of monoterpenes is still uncertain and seems to be much more complicated compares to isoprene. There should be a consistent reason for plants to synthesize and emit BVOCs because the emission of BVOCs requires considerable cost. However, there has not been any unified understanding on the factor controls BVOC emission, yet. To solve this mystery, emission of MNTs under "natural" environment should be one of the hopeful objects because MNTs are known to have variable effect to survive in the environment (e.g. avoidance of herbivory) and because MNTs consist of numerous isomers, it implies the MNT emissions may contain higher information than other single BVOC such as isoprene. MNTs are, in general, known to be emitted from coniferous trees. Some of evergreen broadleaf (EB) trees also emit MNTs. However, report about MNT emission from the EB tree is still quite limited. Only a few EB species emit the BVOCs while other EB species emits no BVOCs even they live in the same community. Although they are living in similar environmental condition, BVOC emission of the EB trees quite differ each other. We hypothesized that there might be a clue to reveal the fundamental principle which determines and controls the BVOC emission. In this presentation, we report a preliminary result of BVOC measurement targeted on the dominant EB trees in Japan.

Experiment

Screening samples were collected for seven EB tree species using a blanch enclosure at the university of Tokyo Tanashi Forest in July 2012. The intensive sample collection has been conducted in Tano Forest Science Station of Miyazaki University (31°51′8″N 131°18′23″E) from 25th through 28th September 2012. The BVOC samples were collected from twenty Castanopsis sieboldii leaves into a glass tube filled with adsorbents using a leaf cuvette on the top of 15 m canopy tower in EB tree community. The samples were analyzed employing a gas chromatograph (GC-FID) coupled with a cryo-focus and thermal desorption system.

Result and discussion

As a result of the screening, only Castanopsis sieboldii was found to be MNT emitter among investigated EB trees. The MNT emission rates ranged from 0.04 - 30 (alfa-pinene) micro gC g⁻¹ h⁻¹. The averaged total MNT emission rate was 19 micro gC g⁻¹ h⁻¹, it is comparable to or exceeds that of Pinus trees which is commonly known as major MNT emitter. Light dependence on the emission was not clearly observed. The emission rates were normalized to obtain basal emission rate, which can be regarded as "emission activity", based on G93 temperature dependence model with empirical coefficient beta of 0.10. Figure 1 shows composition of MNT basal emission rates. A clear contrast of the composition among the individuals can be observed while total emission rates were relatively close each other. Individual 1 emitted sabine as the most abundant MNT while ocimene was one of minor MNTs. On the contrast, individual 2 and 3 emitted ocimene as 2nd or 3rd most abundant MNT while sabinene was minor MNT. Although these individuals are same tree species and are almost same ages, in addition, growing in quite similar environment (within only 1-2 m of distance each other), the composition of the MNTs showed a clear difference. It can be hypothesized that the C. sieboldii trees emitted different types of MNTs as responses to stresses they are exposed to. In other words, spatial distribution of BVOC emitting / non-emitting trees and/or the distribution of BVOC types in the community probably contain a clue to the mystery of fundamental factor of the BVOC emissions.

Keywords: Biogenic Volatile Organic Compound, Atmospheric Chemistry, Biosphere Atmosphere Interaction, Biogeochemistry, Material Cycle, Evergreen Broadleaf Tree
Figure 1 Basal emission rates of monoterpenes emitted from Castanopsis alatoides trees obtained by leaf level BVOC measurement.
Comparison of concentrations and photoreactivities of oxalic acid and metal-oxalate complexes in aerosols

Yoshiaki Yamakawa\textsuperscript{1}, Kohei Sakata\textsuperscript{2}, Aya Sakaguchi\textsuperscript{2}, Yoshio Takahashi\textsuperscript{2}

\textsuperscript{1}Faculty of Science, Hiroshima University, \textsuperscript{2}Graduate School of Science, Hiroshima University

Aerosols have cooling effect to the earth, which is divided into direct and indirect effects. The direct effect is reflection of sunlight directly by aerosols, whereas the indirect effect is the reflection by clouds formed by the aid of aerosols working as cloud condensation nuclei (CCN). Oxalic acid is a main component of secondary organic aerosols (SOA) and abundant in the atmosphere, which is formed by degradation of organic components with longer carbon chain such as malonic acid. Oxalic acid is hygroscopic, which works as CCN with indirect cooling effect. It has been estimated that the degree of cooling effect by the aerosols are equal to that of the warming effect of carbon dioxide. However, there is large uncertainty in the estimation. In fact, if oxalic acid forms insoluble metal-oxalate complexes in the atmosphere, it is possible that the indirect effect can be smaller than the estimation. In addition, in the presence of metal-complex, it is also important to understand the formation processes. Therefore, this study was aimed (i) to decide the ratio of insoluble metal-oxalate complexes to oxalic acid in the aerosols by X-ray absorption fine structure (XAFS) spectroscopy to contribute to precise prediction of global warming and (ii) to evaluate stabilities of oxalic acid and metal-oxalate complex in the atmosphere during the photoreactions. The latter factor can be related to the dominance of metal-oxalate complexes in the atmosphere, if we can show that the photodegradation reactivity of oxalic acid is much larger than that of metal-oxalate complex by ultraviolet light.

We collected size-fractionated aerosols at Higashi-Hiroshima to determine chemical species of oxalic acid. As a result, (i) zinc (Zn) oxalate complex was found in fine particles ($< 2.1$ micron) and (ii) calcium (Ca) complex was present in all the particle sizes except for the 4.2 to 10.2 micron particles. Oxalic acid is SOA that is formed by degradation of organic matters, whereas oxalic acid can be distributed to the particle size from 0.5 to 1.0 micron known as droplet mode, where metal-oxalate complexes can be formed.

To estimate the stabilities of oxalic acid and its metal-oxalate complex with Mg\textsuperscript{2+}, their photodegradations by ultraviolet lights in water were determined by measuring their concentrations at various elapsed time using total organic carbon analyzer. As a result, there was no appreciable difference between the two systems up to 8 hours, but the concentration of the complex was kept larger than that of oxalic acid after 8 hours. Therefore, it is possible that oxalic acid actually exists as metal-oxalate complex in the atmosphere possibly by preferential photodegradation of oxalic acid.

We will also analyze chemical species employing XAFS analysis on other metal ions such as lead and copper to measure precisely the ratio of metal-oxalate complex to total oxalate species. We will also conduct photodegradation experiments by ultraviolet light not only for Mg\textsuperscript{2+} complex but also for Zn\textsuperscript{2+} and Cu\textsuperscript{2+} complexes to confirm the difference of photodegradation caused by chemical species.

Keywords: aerosol, metal-oxalate complex, indirect cooling effect, photoreactivity, X-ray absorption fine structure
Vertical profiles of aerosol size distributions near the surface boundary layer

Masanori Yabuki¹*, Kenshi Takahashi¹, Makoto Matsuda¹, Toshitaka Tsuda¹

¹Research Institute for Sustainable Humanosphere, Kyoto University

Knowledge of the properties of the atmospheric minor constituents is essential in studies on climate change and its effects on human health. The concentrations of ambient trace gases and aerosols, which are emitted by both natural and anthropogenic sources, are influenced by diffusion due to the thermodynamic processes during the air-mass transportation along the prevailing wind flow. Their chemical and physical properties vary both temporally and spatially as a result of various atmospheric processes such as scavenging, nucleation, evaporation, and condensation. Therefore, a comprehensive approach that takes into consideration atmospheric chemistry as well as dynamics and thermodynamics is required for a thorough understanding of air quality.

For elucidating the properties of the minor constituents in the surface boundary layer, we have carried out AEROsol and GAses Profiling (AEROGAP) experiments using a combination of in situ and remote sensing measurements at the Kyoto University Middle and Upper Radar site (34.9 N, 136.1 E) in Shiga Prefecture, Japan, during the summers of 2011 and 2012. In this study, we focus on the temporal variations in the vertical profile of nano- and submicron particles observed by a condensation particle counter and an optical particle counter fixed to tethered balloon platforms. We also discuss the properties derived from aerosol lidar observations compared with those observed using in situ instruments.

Keywords: Aerosol, Size distribution, Tethered balloon, Lidar
Measurement of organic nitrates in the atmosphere using thermal dissociation/cavity attenuated phase shift spectroscopy

Yasuhiro Sadanaga\textsuperscript{1,*}, Ryo Takaji\textsuperscript{1}, Kazuo Nakajima\textsuperscript{1}, Kazunari Suzuki\textsuperscript{1}, Atsushi Matsuki\textsuperscript{2}, Keiichi Sato\textsuperscript{3}, Hiroshi Bandow\textsuperscript{1}

\textsuperscript{1}Osaka Prefecture University, \textsuperscript{2}Kanazawa University, \textsuperscript{3}Asia Center for Air Pollution Research

Peroxyacyl nitrates (PANs) and alkyl nitrates (ANs) act as one of the reservoirs of nitrogen oxides (NO\textsubscript{x}) in the atmosphere. Since their lifetime is longer than that of NO\textsubscript{x}, they can be transported over a long-distance and would be important as transboundary pollutants. PANs and ANs are generally observed using GC/ECD (gas chromatograph / electron capture detector) or GC/NICI-MS (gas chromatograph / negative ion chemical ionization-mass spectrometry) method. While these GC methods can measure concentrations of each constituent in PANs and ANs, they have low time resolution. Measurements by GC/ECD which has radioisotope are very difficult in Japan because of laws and regulations. On the other hand, frequent maintenances are required for measurements using GC/NICI-MS. Thus observation data of PANs and ANs in Japan are quite low.

In this research, continuous measurement system of total PANs and ANs in the troposphere has been developed by using a thermal dissociation / cavity attenuated phase shift spectroscopy (TD/CAPS) method. Both PANs and ANs are thermally decomposed to produce NO\textsubscript{2} and then NO\textsubscript{2} is measured by CAPS method. This system can observe PANs and ANs with high time resolution while this system cannot separate constituents of PANs and ANs.

This system has three lines; (1) NO\textsubscript{2} line, (2) PANs line, and (3) ANs line. The NO\textsubscript{2} line consists of a quartz tube without heating. The PANs and ANs lines have quartz tubes heated at 433 K and 633 K, respectively. Concentrations of NO\textsubscript{2}, NO\textsubscript{2} + PANs and NO\textsubscript{2} + PANs + ANs can be obtained from the NO\textsubscript{2}, PANs and ANs lines, respectively. These concentrations are sequentially measured by switching solenoid valves and then NO\textsubscript{2}, PANs and ANs concentrations are obtained. Since a part of HNO\textsubscript{3} is pyrolyzed in the ANs line, annular denuder coated with NaCl to remove HNO\textsubscript{3} is set before the heated quartz tube in the ANs line.

The decomposition efficiencies of PANs and ANs were investigated and obtained to be 100 and 90\%, respectively, for all kinds of PANs and ANs examined.

Continuous field observations of PANs and ANs concentrations have been being carried out at NOTOGRO supersite at Suzu, Noto Peninsula, since November 2012. In this presentation, the observational results and preliminary analyses of PANs and ANs concentration variations with NO\textsubscript{x}, NO\textsubscript{y}, total inorganic nitrate, O\textsubscript{3} and CO concentrations simultaneously observed are introduced.

Keywords: total odd nitrogen species, organic nitrates, cavity attenuated phase shift spectroscopy
Spheroidal carbonaceous particles (SCPs) are produced by the high-temperature combustion of fossil fuels (e.g., in thermal power station) and are emitted to the atmosphere. They are unambiguous indicators of atmospheric deposition from industrial fossil fuel combustion. We examined elemental composition of the particles in surface sediments nearby industrial cities in East Asia to clarify the elemental characteristic of the particles emitted from each country. Then we also analysed the particles in surface sediments collected in islands in Sea of Japan and the sites along the sea to clarify their source area. We collected 9 samples in Japan, 6 samples in China, 6 samples in Korea, and 5 samples in Taiwan as samples of industrial cities, and 4 samples in remote islands in Japan Sea, and 3 samples at sites along the sea as samples of remote area. We used EDS (energy dispersive spectroscopy) to quantitatively analyze the concentrations of Na, Mg, Al, Si, P, S, Cl, K, Ca, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, and Zn in the particles. As a results, the compositions of SCPs differ among Japanese, Chinese, Korean and Taiwanese cities, especially in terms of Si, S and Ti; particles in Japan and Korea are enriched in Si and S, particles in China are enriched in Si and Ti, and particles in Taiwan are enriched in Si, S and Ti. Results of liner discriminant analysis indicate the particles in each country are discriminated with high reliability of approximately 90%, based on their chemical composition. Based on these discriminant functions, the particles in Oki Islands, Iki Islands and Goto Islands were classified to three types of Japan-Korea type, China Type and Taiwan type. Over 30% of china type was recognized in each sample in these Islands, that are probably derived from China.

Keywords: spheroidal carbonaceous particles, fossil-fuel combustion, chemical composition, East Asia, long transportation
Simulation of aerosol-cloud interactions in spring over East Asia using WRF-chem model: Comparison with aircraft obs.

Ray Takatani¹*, Makoto Koike¹, Hitoshi MATSUMI¹

¹University of Tokyo

1. Introduction

Aerosols act as cloud condensation nuclei and increase/decrease cloud droplet number concentrations. They play an important role for cloud microphysics, dynamics, and radiative forcing. Recently, various observational and modeling studies have been conducted to understand aerosol-cloud interactions in the lower stratus over the Eastern Pacific (e.g., California, Chile). However, there are still few studies focused on aerosol-cloud interactions over East Asia, although aerosol concentrations are considerably high and large aerosol-cloud interactions are expected over the regions.

In this study, we calculated aerosol and cloud droplet number concentrations using a regional three-dimensional model, WRF-Chem, which explicitly expresses the impact of aerosols on cloud microphysics processes. The purpose of this study is to validate model-calculated aerosol and cloud droplet concentrations with the observation during the A-FORCE aircraft campaign.

2. A-FORCE aircraft observation

Our target is low-level stratus/stratocumulus clouds (<2 km in altitude) without precipitation. During the A-FORCE campaign, aerosol number concentrations below the cloud base and cloud droplet number concentrations just above the cloud base were observed 9 times over the Yellow Sea and the East China Sea in March and April 2009 [Koike et al., 2012].

3. WRF-Chem model calculation

Model simulations were conducted over East Asia using the WRF-Chem v3.4. The simulation periods are from 21 March to 26 April 2009. We used 3 domains with the horizontal grid resolutions of 108 km, 36 km, and 12 km, respectively. Vertical resolutions are 46 layers from the surface to 100 hPa (the lowest layer is about 30 m). Both anthropogenic and biomass burning emissions are considered in the simulation. Chemical and microphysical processes of aerosols were calculated by the MOSAIC module with 8 size bins. The Morrison double moment scheme was used for cloud microphysics.

4. Results

Mean observed and calculated aerosol number concentrations (>130 nm in diameter) have good agreement within 30% (underestimation by 27% by model). They also have a positive correlation (r²=0.32), suggesting that spatial and temporal variations of aerosol number concentrations (the transport of anthropogenic aerosols from the Asian continent) were generally reproduced by the model. On the other hand, cloud droplet number concentrations were overestimated by 90%, while a good correlation (r²=0.83) was found between measurements and model simulations.

To quantify the impact of aerosols on cloud microphysics, we compared the relationship between aerosol (>130 nm in diameter) and cloud droplet number concentrations. Positive correlations are found for both measurements and simulations. However, the ratio of cloud/aerosol number concentrations is by overestimated by 170% by the model. The potential reasons of this overestimation are 1) the underestimation of entrainment and 2) the overestimation of aerosol activation to cloud droplets in the model.

Keywords: aerosol, cloud, indirect effect, numerical simulation
Model inter-comparison for evaluation on source sensitivities of atmospheric pollutants over East Asia

Kazuyo Yamaji1*, Kohei Ikeda1, Masayuki Takigawa1, Tatsuya Nagashima2, Yugo Kanaya1

1Japan Agency for Marine-Earth Science and Technology, 2National Institute for Environmental Studies

Atmospheric pollutants were simulated by using two regional CTMs, WRF-Chem and WRF/CMAQ and a global CTM, CHASER over East Asia for the year 2005. Simulated surface O3 over Japan by WRF/CMAQ was higher than that by CHASER especially in summer, and overestimated observed O3 at EANET monitoring sites. Contributions from 5 source regions; north China(CHN), central China(CHC), and south China(CHS), Korea(KRE), and Japan(JPN) on 6 areas on these regions were evaluated based on sensitivity simulations with 20% reduction in anthropogenic emissions. These models resulted that the 20% emission reductions on CHC would gain 0.8% decrease of surface O3 over Central Japan in spring, and that was comparable to the O3 decrease of 0.8-0.9% over Central Japan by the 20% emission reductions on JPN. As for the summer case by using CHASER, the O3 decreases of 0.6% and 1.6% over Central Japan by 20% emission reductions over CHC and JPN, respectively. On the other hand, the regional CTMs, WRF-Chem and WRF/CMAQ, resulted 0.9% (CHC emission reduction) and 3.1% (JPN) O3 decreases and 0.7% (CHC) and 2.4% (JPN) O3 decreases, respectively, and that were 1.2-1.9 times higher than those by CHASER.

Keywords: air quality models, East Asia, atmospheric pollutants, ozone, PM2.5, inter-comparison
Dynamics of particulate matter in the atmosphere

keisuke ikeda\textsuperscript{1,∗}, Osamu Nagafuchi\textsuperscript{1}, Ken’ichi Osaka\textsuperscript{1}

\textsuperscript{1}Graduate School of Environmental Science, the University of Shiga Prefecture

In recent years, due to the rapid economic growth and industrial development, many air pollutants from East Asian regions are discharged into the atmosphere. This trend is seen to continue, transboundary pollution to Japan may become more serious. In this study, we collected particulate matter at two points. One is the surface layer which we live, and the other is the atmospheric boundary layer whose air is easily transported. Our purpose is the elucidation of particulate matter dynamics and of the effect of transboundary pollution from East Asia.

Collecting particulate matter is conducted on period between April, 11, 2012 and August, 3, 2012 at the University of Shiga Prefecture as the surface layer and on period between October, 17, 2012 and November 22, 2012 at the Mt. Ibuki in Shiga prefecture as the atmospheric boundary layer. Sample we collect is measured ion component by ion chromatography and heavy metal by ICP-MS. Analysis of backward trajectories used HYSPLIT model provided by NOAA.

At the surface layer, we observed $\text{NO}_3^−$, $\text{NH}_4^+$, nss $\text{SO}_4^{2−}$, nss $\text{Ca}^{2+}$ at high concentrations, when we observed yellow sand in spring. At this time, the air mass comes from the continent. On the other hand, same trend was not seen in summer. At this time, the air mass comes from the ocean. Forms of deposition are mainly $\text{NH}_4\text{NO}_3$, $(\text{NH}_4)_2\text{SO}_4$, $\text{Ca(NO}_3)_2$, $\text{CaSO}_4$ during observing period.

At the atmospheric boundary layer, we didn’t observe same trend as the surface layer. Forms of transport are mainly $\text{NaNO}_3$, $\text{Na}_2\text{SO}_4$ during observing period. This indicates that $\text{NaCl}$ which derived from sea salt is altered by $\text{HNO}_3$ and $\text{SO}_2$.

Keywords: particulate matter, transboundary pollution, yellow sand
Calculated mercury deposition in north of Lake Biwa

Naoko Hishida1*, NAGAFUCHI, Osamu2, OSAKA, Ken’ichi2, MIYAKE, Takayuki2

1Environmental Science Graduate School, the University of Shiga Prefecture, 2the University of Shiga Prefecture

Gaseous elemental mercury (GEM) account for above 95% out of the atmosphere of mercury, have the property that is hard to dissolve in water and circulates through the whole earth with the atmosphere. GEM becomes reactive gaseous mercury (RGM) that is easy to dissolve in water by the oxidation, and it is removes by deposition from the atmosphere with particle-mercury (p-Hg). Therefore mercury concentration in precipitation may elevate at the place apart from the mercury emission source. Furthermore, mercury removed from all over the atmosphere by deposition is in aquatic area and bioaccumulates in the process of the food chain. In this study, it was intended that calculated mercury deposition from mercury concentration in precipitation in north of Lake Biwa.

Sampling site is Surumi in north of Shiga prefecture there is annual mean precipitation 2800mm and classified in the heavy snowfall area. Two forms samplers, one is automatic rain sampler for researched a change of the mercury concentration in precipitation in one rain, and one is bulk deposit sampler for calculated mercury deposition in here. Automatic rain sampler developed so that every fixed quantity collected precipitation (as for every 5mm). Precipitation sampling was conducted from June to November, 2011 and from March to November, 2012. Sampling was not able to observe it for the snow in the winter season (from December to February). From July, 2012, samples were filtered by PTFE filter, and measured by alkali reduction cold vapor atomic fluorescence spectrometry.

It was 29 rain that samples were collected by September, 2012, and the total mercury concentration in precipitation fluctuated from 1.09 to 25.9 ng/L, and mean was 7.15 ± 5.31 ng/L. Pattern that became least concentration to begin to fall was frequent, but total mercury concentrations change in one rain did not showed constant pattern. By the precipitation with the typhoon of September, 2011, total mercury concentration in precipitation gradually rose. When rain cloud arrived at Surumi through mercury emission areas, mercury concentration rises. It is thought that the change of the mercury concentration in precipitation is related to the passage course of the rain cloud. In addition, mercury deposition that calculated from bulk deposit samples became 19.7 g/km⁻²/year (from September, 2011 to August, 2012).

Keywords: mercury, deposition, Lake Biwa
Atmospheric mercury in the free troposphere

Yuki Nishida¹*, Osamu Nagafuchi², Ken’ichi Osaka², Takayuki Miyake²

¹Environmental Science Graduate School, the University of Shiga Prefecture, ²School of Environmental Science, the University of Shiga Prefecture

In order to clarify the dynamics of atmospheric mercury in the free troposphere (FT), continuous observation of atmospheric mercury were carried out in Norikura Solar Observatory in Japan. The observation period is one week each of Oct 2011 and Oct 2012. Gaseous mercury (TGM) and particulate mercury (PHg) concentration in Norikura Oct 2011 were 0.72 - 1.23 ng m⁻³ and 17.0 - 121.0 pg m⁻³, respectively. TGM concentration in 2012 was 0.2 - 2.4 ng m⁻³, and the mean was 1.6 ng m⁻³. A diel variation of TGM was observed with daytime highs and nighttime lows in almost all of both periods. This diel pattern is considered to be due to upslope wind of boundary layer air resulting from a mountain surface that is warmed up with sunlight in daytime. Therefore such observation of atmosphere in the FT to use mountains, data is collect at night. In addition, the observation of 2012, sharp rise in the TGM concentration has been observed. Almost simultaneously with the concentration increasing of TGM, temperature has dropped drastically, and we saw a half inches of snow. This is thought to be due to cold air mass that flowed into Japan came from the continent, this suggested the possibility of long-distance transport of mercury.

Keywords: mercury, atmospheric chemistry, free troposphere, long - range transport
Measurement of fluorescent particles in Fukue Island

Fumikazu Taketani$^{1,*}$, Xiaole Pan$^1$, Yugo Kanaya$^1$

$^1$JAMSTEC

Among various optical techniques applied for the atmospheric particle detection, fluorescence is useful for detecting certain types of organic particles, especially those of biological origin. In this study, we employed a single-particle fluorescence sensor, WIBS-4, for the detection of fluorescent particles, to demonstrate the capability of the classification of organic particles in the ambient air.

We conducted ambient air measurements from Sep 16 to Dec 14, 2011 using the WIBS-4 instrument at Fukue Island (32.75N, 128.68E) in Japan. We detected 36,000,000 particles during the observation period, and the ratio of the fluorescent particles to the total varied in the range of 2 - 65%. In Oct.6, high number concentrations were observed, suggesting transportation from the continent by the trajectory analysis. During this period, the ratio of the fluorescent particles to the total was about 50%, suggesting that transported particles include fluorescent material. In the presentation, we will discuss the comparison of fluorescence pattern from individual particles.

Keywords: aerosol, fluorescence
Inhomogeneity of NO2 over Fukuoka, an urban site in Japan observed by MAX-DOAS

Hisahiro Takashima1,*, Yugo Kanaya2, Kodai Ito1

1Faculty of Science, Fukuoka University, 2JAMSTEC/RIGC

Since August 2012, continuous NO2 observations have been performed using ground-based Multi-Axis Differential Optical Absorption Spectroscopy (MAX-DOAS) at Fukuoka (33.55N, 130.36E), an urban site in Japan. MAX-DOAS is a passive remote sensing technique using scattered visible and ultraviolet solar radiation at several elevation angles. We investigate inhomogeneity of NO2 by observing at two azimuth angles, Tenjin (downtown area) direction and Itoshima (out of downtown area) direction. We discuss the observed inhomogeneity with a focus on the three factors: inhomogeneity of NO2 emissions, development of the boundary layer, horizontal transport associated with land/sea breeze.
A Study on Detection Methods for Atmospheric Small Particles Based on Lidar Techniques

Yutong Liu1*, Masanori Yabuki1, Toshitaka Tsuda1

1Research Institute for Sustainable Humanosphere

Knowledge of the aerosol size distribution in the accumulation mode (50-1000 nm) is essential in studies on human health because small particles could even penetrate into lung foam, thus increasing the risk of bronchitis or lung and heart diseases. Optical Remote sensing techniques such as a lidar are effective for monitoring aerosols with high temporal and spatial variations. Aerosol instruments using light of wavelengths 350-1500 nm have been put into practical use, and they are effective for detecting particles with sizes comparable to the wavelength. However, to estimate quantitatively the shape of the particle size distribution, more information of small particles with radii around 100nm is required.

In this study, a lidar analysis method is proposed to derive the aerosol size distribution for a wide range of particle sizes, including 100 nm. The algorithm can be divided into two parts. The first part applies the conventional analysis using the wavelength dependence of aerosol physical parameters based on single scattering lidar techniques. The second part involves the use of multiple scattering via multiple-filed-of-view lidar signals for achieving a probing wavelength less than 350 nm; although such short wavelengths can be used to obtain information on small particles, they have not been used for aerosol measurement owing to the strong light absorption by atmospheric constituents such as ozone. The analysis is accomplished by directly fitting the observed lidar signals and the retrieved aerosol parameters to the theoretical values that are based on the look-up-table which is constructed on the basis of Mie scattering and multiple scattering calculations as a function of the log-normal size distribution.

The retrieved particle mode radii in a range of 50-200 nm agree with the originally assumed parameters in combination of single scattering and multiple scattering algorithms, even if noise is contained. In order to evaluate the noise effect, we simulated a sensitivity analysis. When random errors of 5%, 10%, 20%, and 50% were added to the prepared parameters theoretically calculated by use of the geometric mean radius of 100 nm for the aerosol size distribution, the retrieved mean radius (upper and lower limits) derived from the proposed method were 103 nm (83-128 nm), 100 nm (69-144 nm), 89 nm (48-163 nm), and 84 nm (38-186 nm), respectively.

Keywords: Lidar, Aerosol
Development of a scanning Raman lidar for observing the spatio-temporal distribution of water vapor

Makoto Matsuda, Masanori Yabuki, Toshitaka Tsuda, Kenshi Takahashi, Ken-ichi Yoshikawa

1Research Institute for Sustainable Humanosphere, Kyoto University

Water vapor and aerosol particles are important atmospheric constituents that play a key role in the atmospheric processes such as thermodynamics, radiative forcing, cloud physics, and chemistry. Atmospheric constituents near the surface are highly variable spatially and temporally, because of the complex turbulent flow over the surface. It is required to innovative techniques for observing the distributions of atmospheric constituents with good spatio-temporal resolution. We have newly developed a scanning Raman lidar to measure the spatio-temporal distributions of the water vapor and aerosol particles near the surface, which is useful to study the detailed behavior of meteorological phenomena as well as interactions of aerosol particles with water vapor.

Considering the eye-safe operation in urban districts, we employed the UV laser of 355 nm. We developed a scanning mirror system which comprises with highly reflective mirrors and a rotational stage. By use of the program-controlled rotational stage, vertical scan into any zenith direction can be operated with a maximum speed of 1.8 deg./s. Differences between the temporal variations of water vapor mixing ratio by the scanning Raman lidar and those by the conventional lidar for observing a vertical point are less than 2.5 %. It is indicated that the developed system can measure water vapor correctly though the scanning system is attached.

We have demonstrated the potential of the scanning Raman lidar in the forest region at the Shigaraki MU observatory in August and October, 2012. We performed a vertical scan in a zenith sector of 48 deg. with a constant step width of 1.5 deg. The temporal resolution of each pointing direction was 30 s. In this observation, we found that water vapor mixing ratio within the surface boundary layer varied in a range of 13.5 - 16.5 g/kg within the range of 400 m. It is suggested that the spatial variations are highly sensitive under the different topography. During the vertical plane measurements with the wide range scanning, it is found that the thickness of the atmospheric boundary layer changes into the range of about 200m according to the observed directions. During the high spatial and temporal resolution measurements with the continuous scanning at a constant speed, a single cross-sectional distribution can be acquired every 90 s; this indicates the possibility of understanding the cloud formation and modification processes via continuous observations. Scanning Raman lidar has an advantage of obtaining a detailed structure that is difficult to gather from other instruments.

Keywords: lidar, water vapor, aerosol
Parametric studies on temperature lidar with a multispectral detector

Kenichi Yoshikawa, Masanori Yabuki, Toshitaka Tsuda

1Research Institute for Sustainable Humanosphere, Kyoto University

Measurements of atmospheric temperature and water vapor in the troposphere are essential for studying atmospheric processes such as dynamics, thermodynamics, and cloud physics. Remote sensing techniques have obvious advantages for continuous observation of the spatial distributions of meteorological parameters. The Raman lidar is a laser-based remote sensing instrument used to quantify the distribution of water vapor mixing ratio and atmospheric temperature. We have developed several Raman lidar systems that measure water vapor by detecting the vibrational Raman scattering. Recent improvements in the performance of optical components have led to the development of better water vapor lidar systems that are portable and easy to use. More detailed spatio-temporal distributions of water vapor can be acquired by a lidar equipped with a scanning mirror system. On the other hand, the polychromator design for temperature lidar is much more complex than that for water vapor, because the temperature lidar method is based on the fact that the intensities of the lines within the rotational Raman band exhibit slightly different dependencies on temperature. Therefore, temperature lidar is used less widely than the water vapor system.

In this study, temperature lidar with a multispectral detector is proposed to construct a system that is compact, robust, and easy to align for the detection of rotational Raman signals. The multispectral detector enables 32-channel simultaneous photon counting acquisition and provides spectral and range-resolved data by applying lidar techniques. While conventional temperature lidar methods detect the ratio of two rotational Raman lidar signals of opposite temperature dependence in combination with several edge and interference filters, the multispectral lidar detector can grasp the shape of the rotational Raman spectrum. Therefore, the estimation of temperature can be accomplished by directly fitting the observed lidar signals to the shape of the theoretical values of a rotational Raman spectra that exhibit different dependencies on temperature. The use of the multispectral detector for detecting rotational Raman signals has several advantages. In particular, it can reduce uncertainties in the optical alignment of the polychromator and in the stability of the laser wavelength. Furthermore, the multispectral receiver system can be made more compact and less expensive than the conventional system with an interference-filter-based polychromator.

The accuracy of temperature derived from multispectral lidar signals depends on both the spectral resolution and spectral range of the multispectral detector. Therefore, the values of these parameters should be set appropriately to improve the accuracy of temperature estimation. In this study, we estimated the effect of both spectral resolution and spectral range on the accuracy of temperature estimation, and found the ideal combination of those optical parameters. Then, we calculated the accuracy of our proposed method for temperature estimation via a computer simulation of selected cases in combination with both the spectral resolution and spectral range of a multispectral detector.

Keywords: temperature lidar, multispectral detector
NMD Fractionation Estimated from SO Isotopologues Photolysis UV Spectra

Sebastian Danielache\textsuperscript{1,*}, Tomoya Suzuki\textsuperscript{1}, Shinkoh NANBU\textsuperscript{1}, Yuichiro Ueno\textsuperscript{1}

\textsuperscript{1}Department of Materials & Life Sciences, Faculty of Science & Technology, Sophia University, \textsuperscript{2}Department of Earth and Planetary Sciences, Tokyo Institute of Technology

Understanding the mechanism of sulfur isotopic fractionation phenomena has been used for some time as tool to the understanding of reducing atmospheres. Sulfur Non-Mass Dependent (NMD) fractionation signals reported for the Archean and Early Proterozoic (>2300 Ma) atmosphere where the photodissociation of sulfur bearing species play a significant role since the concentration of oxygen is estimated to be 10-5 times present atmospheric levels and therefore ultraviolet light permeates throughout the entire atmosphere, however the underlying mechanisms are not fully understood. In order to explain the NMD signal preserved in the geological record other than SO\textsubscript{2} photodissociation chemistry of sulfur compound should be taken into account. In this study we consider isotopic fractionation during photodissociation of SO. Experimental studies are difficult since SO is highly reactive and unstable under most experimental and atmospheric conditions. Consequently, theoretical studies are suitable to evaluate its spectroscopic properties. We report ultraviolet absorption cross sections for 32,33,34,36SO isotopologues. R-matrix theory was employed to compute absorption cross sections. Potential energy curves were calculated at multi-reference correlation interaction (MRCI) method with augmented correlation consistent polarized valence sextuple-z (aug-cc-pV6Z) basis set. The six lower-lying electronic states were explored. Because there are pseudo-crossings between electronic excited states, non-adiabatic effect has been taken into account. Isotopic fractionation constants were theoretically estimated at difference temperatures by assuming that the photolysis rates are proportional to the photoabsorption cross sections for each isotopologues. We make the one box model of the atmospheric chemistry where obtained photolysis rates are utilized, and discuss the results of this model.

Keywords: Sulfur Cycle, Archean Atmosphere, Stable Isotopes