ABSTRACT
During the NASA 2016 KORUS-AQ campaign, the ground based NASA GSFC ozone lidar and balloon borne instrumentation were deployed to the remote Taehwa Forest site (37.3 N, 127.3 E, 151 m AGL) to characterize the transport of pollution downwind of the Seoul metropolitan region. On most days from 02 May to 10 June 2016, continuous hours of lidar profiles of ozone were measured. Select days are shown to represent key ozone events that occurred at the rural site.

1. INTRODUCTION
In order to better assess pollution emission sources and plume evolution in Asia, there has been an international effort to launch the Geostationary Environmental Monitoring Spectrometer (GEMS) to provide hourly measurements of key pollutants (e.g. ozone; nitrogen dioxide (NO2); sulfur dioxide (SO2); particulate matter) over the Korean peninsula and the Asia-Pacific region. As a precursor to the Korean GEMS satellite launch and to further investigate the vertical distribution of pollutants impacting South Korea, the United States (U.S.) National Aeronautics and Space Administration (NASA) and the Korean Ministry of the Environment/Korean National Institute of Environmental Research (NIER) conducted an international cooperative field experiment, entitled the Korea-U.S. Air Quality (KORUS-AQ, https://espo.nasa.gov/home/korus-aq/content/KORUS-AQ) study. The work herein mainly utilizes the synergy of available data to focus on high ozone events at a downwind from Seoul (30 km southeast, Figure 1a) rural site (Taehwa Research Forest, TRF).

2. OBSERVATIONS
2.1 Ozonesondes
An unprecedented O3-sonde record, which were conditioned and prepared with current community standards [1], was recorded at TRF throughout the KORUS-AQ study from 10 May to 12 June. Afternoon soundings (13:30 to 16:30 KST) of O3 (top panel, Figure 2) and temperature (bottom panel, Figure 2) illustrate day-to-day variability at TRF. The two case studies (17 May and 09 June) are highlighted with black boxes.

There were large disparities in boundary layer O3 throughout the campaign period. From 10-16 May, concentrations were mostly between 70-80 ppb, which were associated with cooler temperatures and higher synoptic wind speeds. However by the early afternoon of 17 May, a
A stagnant high-pressure system located over the Yellow Sea ushered in warmer air, calmer winds, and clearer skies. For the remainder of the campaign until 03 June, a similar meteorological system was preserved, providing favorable conditions for rapid O₃ production of concentrations above 120 ppbv. On 04 and 06 June, mixed precipitation events occurred, limiting O₃ production in the surrounding days. However, by 09-10 June another high-pressure system approached the region, increasing temperatures, stagnating winds, and fostering O₃ production.

Figure 2: Afternoon soundings (13:30 to 16:30 KST) of O₃ (top panel) and temperature (bottom panel) illustrate day-to-day variability at TRF from 10 May to 12 June.

2.1 GSFC Ozone DIAL

Vertical profiles of O₃ were measured using the NASA Goddard Space Flight Center TROPospheric OZone Differential Absorption Lidar (GSFC TROPOZ DIAL or TROPOZ [2]) in support of the KORUS-AQ study. The TROPOZ system, a charter NASA instrument in the Tropospheric Ozone Lidar Network (TOLNet, http://www-air.larc.nasa.gov/missions/TOLNet/), derives O₃ concentrations to mostly within 10-15% as compared to nearby O₃-sonde profiles [3] and has been previously utilized to characterize O₃ episodes such as stratospheric-tropospheric exchange [4], terrain driven recirculation/transport events [5,6], and regional transport events. During the field study at TRF, the TROPOZ was operated nearly continuously in order to understand sub-hourly (15-min) variability of O₃ during the early morning, afternoon, and evening hours. Figure 3 includes several days where the lidar was operated continuously for 9-13 hrs and illustrates the key O₃ features systematically observed at TRF during the campaign. To add context with auxiliary measurements, each of the TROPOZ measurement days has the 1-min in situ GSFC O₃ monitor (Thermo 49i) data in the lowest bin, a dashed-line solar curve, and coincident ozone soundings previously reported in Figure 2.

3. RESULTS

During every morning at TRF between 07:00-08:00 KST, the surface ozone monitor observed low (<10 ppbv) surface O₃ concentrations. Low early morning values are associated with a stable surface layer, in which surface frictional effects and titration of O₃ occur. Additionally, TROPOZ measurements in six of the eight days in Figure 4 indicate residual layering of O₃ above the stable surface layer with concentrations above 75 ppbv between 08:00 and 10:00 KST. This O₃ resides between 500 and 1500 m ASL, and is related to the extent of vertical mixing during the previous day’s planetary boundary growth and decay. An example of residual layer height corresponding to the previous day’s ozone mixing height and concentrations is demonstrated in the 9-10 June observations. O₃ with concentrations near 100 ppbv are mixed to near 1700 m ASL by 19:00 KST and result in a residual layer residing at nearly the same height the following morning (as corroborated by the O₃ sonde on 10 June).

As solar radiation and convective mixing increase in the late morning hours (after 11:00 KST), surface concentrations of O₃ generally began to be more representative of concentrations measured aloft from the TROPOZ. During this time, boundary layer growth appears to entrain the residual layer(s) of pollutants helping to increase mid-day O₃ concentrations (e.g. 17-18 May, 9-10 June). Until 14:00 KST, boundary layer O₃ concentrations are mostly between 65-85 ppbv. However, as solar radiation decreases after 14:00 KST most days observe a large (10-40 ppbv) and generally well-mixed contribution of O₃ transported to the site. During all days, O₃ concentrations aloft on each day are above 90
limiting O₃ production in the surrounding days. June, mixed precipitation events occurred, concentrations above 120 ppbv. On 04 and 06 hourly (15-min) variability of O₃ during the early field study at TRF, the TROPOZ was operated.

as compared to nearby O₃-sonde profiles [3].

Figure 2: Afternoon soundings (13:30 to 16:30 KST), surface concentrations of O₃ generally increase in the late morning hours (after 11:00 KST). Surface concentrations of O₃ are a nearly the same height the following morning (as near midnight in the campaign period. The TROPOZ measurements in six of the campaign.

As solar radiation diminishes near 19:00 KST, 17 May, 18 May, and 09 June have observations that continue for several hours. During each of these measurement times after 19:00 KST, O₃ concentrations aloft are significantly larger (40-60 ppbv) than the coincident surface concentrations, indicating the rapid depletion and titration of surface ozone from local emissions. This process then results in the polluted aloft residual layer during nighttime hours that will be mixed down on the following morning to enhance the pollution budget.

4. CONCLUSIONS

Several days of nearly continuous ozone lidar observations have yielded a novel assessment of pollution reaching TRF and other rural sites in South Korea. Several case studies indicate rural sites may be experiencing long-term negative effects of O₃ (which are associated with much worse health and agricultural effects) and therefore more stringent domestic regulatory controls on O₃ precursors.

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REFERENCES


