Effects of the January 2005 GLE/SEP events on minor atmospheric components

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Abstract: It is known from long ago that solar energetic charged particles, driven by the geomagnetic field, are able to produce ionization at different altitudes of the terrestrial atmosphere. Moreover, they can initiate catalytic cycles for the ozone depletion, involving NO\textsubscript{x} (N, NO, NO\textsubscript{2}) and HO\textsubscript{x} (H, OH, HO\textsubscript{2}) components. Nevertheless, only in recent years it was possible to compare chemical models involving atmospheric minor components with satellite data. In this work we looked for effects of the GLE/SEP events occurred during January 2005 on the OH and HNO\textsubscript{3} species of the atmosphere. Results show that there is a response on the minor atmospheric components, which is different in the winter and summer terrestrial hemispheres.

Introduction

It is widely known that minor atmospheric constituents are strongly dependent on the geomagnetic and solar activity. In particular, several induced effects by cosmic rays (CRs) can be singled out at diverse altitudes. For example, in the troposphere the CR flux seems to be connected to the cloudiness formation (see [8] for an early work and [9] for a recent analysis); in the lower stratosphere CRs are an important source of nitric oxide (NO) molecules, by the ionization and dissociation of the molecular nitrogen (N\textsubscript{2}). At Polar Regions and during the winter time, when the lack of sun light prevents the formation of NO via the reaction of nitrogen protoxide (N\textsubscript{2}O) with atomic oxygen, the CR flux can become the main source of nitric oxide [6]. Besides, in stratosphere and mesosphere, solar energetic particle (SEP) events are able to trigger catalytic cycles of O\textsubscript{3} destruction [1]; O\textsubscript{3} is the most important atmospheric component for the Earth’s lives.

During January 2005, after a low/moderate solar activity interval, a big injection of solar energetic particles in the terrestrial atmosphere occurred (\sim 16-22 January). Two Ground-Level Enhancements were identified (GLE68/SEP: 17 January, start \sim 10 UT and GLE69/SEP: 20 January, start \sim 07 UT) in the world-wide network of cosmic ray detectors. This paper focuses on the January 2005 GLE/SEP’s effects on the polar atmospheric chemistry.

Used Data

The employed atmospheric data come from the Micro-wave Limb Sounder (MLS) instrument on the AURA satellite. The NASA EOS (Earth Observing System) MLS is one of the four instruments of AURA launched on 15 July 2004 to sun-synchronous near polar orbit. The MLS is a limb emission instrument that scans the Earth’s limb viewing the microwave emission at different spectral regions. The measured chemical components are: O\textsubscript{3}, H\textsubscript{2}O, BrO, ClO, HCl, HOCl, OH, HO\textsubscript{2}, HCN, CO, HNO\textsubscript{3}, N\textsubscript{2}O, and SO\textsubscript{2} mixing ratios.

In this work we used EOS MLS Version 1.5 Level 2 Data (http://mls.jpl.nasa.gov/data/), from which O\textsubscript{3}, HNO\textsubscript{3}, and OH values were taken. The proton flux was retrieved from GOES 11 files (http://www.ngdc.noaa.gov/stp/GOES/).
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SEP events are able to produce ionization at different altitudes of the terrestrial environment and can initiate catalytic cycles for the ozone depletion, involving NO$_x$ (NO, NO$_2$) and HO$_x$ (OH, HO$_2$) components. Such effects generally refer to the Polar Cap regions (geomagnetic latitudes above 60°). Nevertheless, [3] showed that January 2005 GLE/SEP events produced also two weak and very short (< 12 h) ozone depletions at the external boundary of the Southern Polar Cap.

Moreover, Damiani et al. [2], analyzing the interhemispheric differences, emphasized that: at the Southern high latitudes (day state) the O$_3$ decrease is weak while in the Northern (night state) ones it is strong and long lasting, particularly in the mesosphere. For this reason, Figure 1 (upper panel) exemplifies only the daily ozone profiles in the Northern Hemisphere (location: ∼ 75°-82° N), before and after the GLE/SEP events of January 2005. The elevated values of the mesospheric O$_3$ mixing ratio on the day before the event (blue profile) is related to the existence of a third ozone maximum during the winter time [5]. In fact, at elevated latitudes of the winter hemisphere the low Sun light leads to a minor efficiency of the UV flux on the water vapour photolysis; therefore, the reduced production of HO$_x$ components in “night” condition facilitates the increase of the ozone concentration. Mesospheric ozone depletion (more than 1 ppmv) is evident on 18 January (the day after the first GLE/SEP) and lasts for some days. In the stratosphere (above 1 hPa) the ozone decrease is less evident from the daily profiles; an analysis of NO$_x$ component should be performed. The MLS instrument cannot measure atmospheric NO$_x$ but it is able to retrieve nitric acid, a good proxy for NO$_x$ (reservoir nitrogen). During January 2005 SEPs, together with the stratospheric ozone depletion, an HNO$_3$ increase appears between 10 and 2 hPa (∼ 30-40 km), persisting till the end of the month. Figure 1 (lower panel) shows its temporal evolution (location: ∼ 75°-82° N) by using derived HNO$_3$ (mixing ratio) contours, from 15 to 31 January 2005. This increase of nitric acid can originate from two different processes.

The first process involves the most important HNO$_3$ production phenomenon:

Figure 1: Daily ozone profile of averaged values (volume mixing ratio) during several days of January 2005 (upper panel) and contours of averaged HNO$_3$ (volume mixing ratio) values during the second part of January 2005 (lower panel). Selected location: ∼ 75°-82° N.
NO₂ + OH → HNO₃ + M.

Therefore, the HNO₃ increase can be the result of the OH and NO₂ raise during SEPs; this reaction is fast enough to produce the amount of nitric acid similar to that observed during October-November 2003 GLE/SEP events [4].

The second process is related to the ion chemistry [7] and may be important since it is an active process also during night time (i.e. winter hemisphere at elevated latitudes). Through the reaction of water cluster ions with NO₃ the nitric acid is able to raise. Nevertheless, the elevated photolysis at the summer hemisphere prevents to single out the HNO₃ variability for the January GLE/SEPs.

During January 2005 GLE/SEP events it was possible also to highlight the increase of the OH molecules (proxy for HOₓ) associated with the solar protons transit in the atmosphere. Figure 2 shows the OH temporal evolution for ~ 75°-82° N (upper panel) and S (lower panel) for 15-24 January 2005. The sudden and intense OH enhancement evidences the goodness of expectations from available models. The OH concentration raises several hundreds of % at Northern latitudes, whereas it remains almost constant at South (only a weak modulation is present) where the intense solar illumination increases the OH background values and hides the induced effects by SEPs. To notice that the response of odd hydrogen species to SEPs is very fast, almost contemporary, either in summer or winter hemisphere.

Summary and conclusions

GLE/SEP induced effects on some minor components of the terrestrial atmosphere have been illustrated for the GLE/SEP events occurred during January 2005. Results for a Polar Cap location (~ 75°-82°) can be summarized as follows:

- A relevant increase of the HNO₃ from 19 to 28 January 2005 was identified from ~ 2.5 hPa to 5.5 hPa at the Northern hemisphere (Figure 1, lower panel).
- The OH increase during the GLE/SEP events presented a fairly good reproduction in the Northern hemisphere of the proton flux profile (Figure 2, upper panel).

Figure 2: Solar proton flux (particle energy E > 10 MeV) from GOES 11 data and contours of OH values (volume mixing ratio) from MLS/EOS data for ~ 75°-82° N (upper panel) and ~ 75°-82° S (lower panel).
Moreover, in the upper mesosphere (∼ 60-80 km) the OH variability seems to be triggered by a cut-off value in the proton flux (∼ 10 pfu for particle energy E > 10 MeV). In fact, the SEP flux arriving on the terrestrial environment on 16 January initiated the OH modulation, which is ended when the proton flux went below ∼ 10 pfu. This behaviour is less evident in the Southern hemisphere (Figure 2, upper panel). Moreover, in the lower mesosphere the scenario is more complex.

In conclusion, it is stressed that the high OH values under solar illumination make difficult to single out the OH rise induced by SEPs, since the background concentration of H-species depends on the H₂O concentration (more elevated in the summer mesosphere). On the contrary, the reduced solar illumination during the winter time facilitates to highlight the atmospheric chemistry changes. Moreover, in the winter hemisphere the OH changes are long lasting (if compared with the summer ones, because of the OH short life under solar light). Very probably, the different OH concentration and life during the extreme seasons is the main reason of the elevated and long lasting O₃ depletion in the Northern mesosphere and the short and feeble O₃ decrease at the Southern one.

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References