Mesospheric nitric acid enhancements during energetic electron precipitation events simulated by WACCM-D

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Key Points:

- D-layer ion chemistry enhances production of mesospheric nitric acid through ion cluster recombination with a seasonal maximum in winter
- Medium-to-high energy electrons precipitating into the mesosphere in April-May 2010 further enhances the formation of nitric acid
- Occurring at a time of year with low background abundance, the enhancement brought nitric acid abundance close to its annual cycle maximum
Abstract

While observed mesospheric polar nitric acid enhancements have been attributed to energetic particle precipitation through ion cluster chemistry in the past, this phenomenon is not reproduced in current whole-atmosphere chemistry-climate models. We investigate such nitric acid enhancements resulting from energetic electron precipitation events using a recently developed variant of the Whole Atmosphere Community Climate Model (WACCM) that includes a sophisticated ion chemistry tailored for the D-layer of the ionosphere (50-90 km), namely WACCM-D. Using the specified-dynamics mode, i.e., nudging dynamics in the troposphere and stratosphere to meteorological re-analyses, we perform a one-year long simulation (July 2009-June 2010) and contrast WACCM-D with the standard WACCM. Both WACCM and WACCM-D simulations are performed with and without forcing from medium-to-high energy electron precipitation, allowing a better representation of the energetic electrons penetrating into the mesosphere. We demonstrate the effects of the strong particle precipitation events which occurred during April and May 2010 on nitric acid and on key ion cluster species, as well as other relevant species of the nitrogen family. The one-year-long simulation allows the event-related changes in neutral and ionic species to be placed in the context of their annual cycle. We especially highlight the role played by medium-to-high energy electrons in triggering ion cluster chemistry and ion-ionic recombinations in the mesosphere and lower thermosphere during the precipitation event, leading to enhanced production of nitric acid and raising its abundance by two orders of magnitude from $10^{-4}$ to a few $10^{-2}$ ppb.

1 Introduction

Nitric acid (HNO$_3$) is an important minor species in the middle atmosphere. Over the last decades, its distribution has been characterized by means of ground-based, aircraft and satellite observations at infrared and millimetre wavelengths [e.g., de Zafra and Smyshlaev, 2001; Santee et al., 2004; Orsolini et al., 2005, 2009; Stiller et al., 2005; Kinnison et al., 2008; Verronen et al., 2008, 2011; Urban et al., 2009; Damiani et al., 2016]. HNO$_3$ is most abundant in the polar lower stratosphere in winter, where its long lifetime during the dark conditions makes it a key reservoir of reactive nitrogen (NOx = NO+NO$_2$), which drives the main ozone-depleting cycle in the mid and upper stratosphere. Yet, satellite observations have revealed descent of HNO$_3$ from above the upper stratosphere, leading to the formation of a secondary maximum, well above the main lower stratospheric layer. These recurrent enhancements are observed in the polar regions of both hemispheres with a large degree of inter-annual variability, and they have been linked to energetic particle precipitation (EPP), the main source of NOx in the mesosphere-lower thermosphere (MLT). Satellite and ground-based observations are often limited to the stratosphere due to retrieval constraints, but occasional mesospheric HNO$_3$ enhancements have been documented during solar proton events (SPEs) [Verronen et al., 2011], when larger abundances expand vertically the validity range of the retrievals.

Both SPEs and energetic electron precipitation (EEP) lead first to the formation of primary ions such as O$^+$, O$_2$$^+$, N$^+$, and N$_2^+$ through dissociation and dissociative ionisation. These ions are then involved in fast ion-chemistry reactions, ultimately producing NOx and hydrogen oxides (HOx) (e.g., see Sinnhuber et al., 2012 for a review). The chemistry of ion clusters plays a key role in the production of the neutral nitrogen species. Ion clusters are groups of $m$ molecules tied to a positive or negative ion. For example, $m$ water molecules can form the hydrated water cluster $\text{H}^+(\text{H}_2\text{O})_m$, or else other water clusters like NO$_3^-$($\text{H}_2\text{O})_m$ or NO$^+(\text{H}_2\text{O})_m$. The number $m$, the order
of the cluster, indicates the attachment of $m$ molecules. Another important ion cluster group for
the production of HNO$_3$ which was identified in previous studies (Verronen et al., 2011;
Andersson et al., 2016) is the nitric acid cluster NO$_3^-(\text{HNO}_3)_m$, i.e., $m$ HNO$_3$ molecules attached
to the negative NO$_3^-$ ion.

With only a few exceptions, the EPP-related upper stratospheric/mesospheric source of HNO$_3$ is
not accounted for in middle atmosphere global models, as they do not incorporate the relevant
ion chemistry. Reddmann et al. (2010) included a parametrization of HNO$_3$ hydrolysis in a
(2012) used the same approach, along with an idealised background distribution of hydrated
water clusters, in simulations using the free-running Whole Atmosphere Community Climate
Model (WACCM). In addition to producing wintertime enhancements of HNO$_3$, the latter study
showed complex chemical-dynamical feedbacks, which extended the effect of the EPP-induced
winter increase of NOx into the warm season by weakening the polar vortex and enhancing the
poleward transport of NOx from lower latitudes. The vortex weakening (when the parametrized
ion chemistry reaction was included) arose from strengthened planetary waves due to zonal
asymmetries in ozone induced by enhanced NOx anomalies.

It is apparent that ion chemistry needs to be taken into account to correctly represent the
observed distribution of HNO$_3$, and also that this chemistry could exert a potential feedback on
ozone and dynamics. Hence, a more comprehensive ion chemistry is needed to account for the
effect of EPP on neutral species in models. Previous studies with comprehensive ion chemistry
were limited to one-dimensional (1-D) models, which at best only crudely represent transport.
For example, Verronen et al. (2011) investigated the chemistry involved in the HNO$_3$
enhancements during the SPEs of January 2005 and December 2006. They compared the results
from the comprehensive 1-D Sodankylä Ion and Neutral Chemistry (SIC) model with satellite
observations from the Microwave Limb Sounder (MLS) instrument at the highest sampled
latitudes, where horizontal mixing was assumed to be small. The same model was used to
analyse the chemical impact of EPP during pulsating auroras (Turunen et al., 2016).

Very recently, an extended ion chemistry based on a subset of reactions and species from the
comprehensive SIC model has been incorporated into the global chemistry-climate model
WACCM. This variant of WACCM called WACCM-D, where D stands for the D-layer of the
ionosphere (below the mesopause), is described in detail by Verronen et al. (2016). Andersson et
al. (2016) investigated the chemical effects of the SPE of 2005, in particular on HNO$_3$, and
showed that the production of HNO$_3$ increased by two orders of magnitude between 40 and 70
km, bringing WACCM-D closer to the MLS satellite observations.

The aim of this paper is to investigate, through a case study, the effect of EEP events on the
distribution of HNO$_3$ and relevant ion clusters in the new WACCM-D model in comparison to
the standard WACCM, in the presence of medium-to-high energy electron (MEEs) forcing. We
analyse a series of events which took place in April and May 2010, the first in a series of
moderate geomagnetic storms that perturbed the upper atmosphere from April to July 2010
(Kirkwood et al., 2015). Our focus is on the high-latitude mesosphere and lower thermosphere.
Hence, our study differs from the above-mentioned studies with WACCM-D in that we
investigate EEP events rather than a SPE. Our simulation also covers an entire year rather than
the weeks around the precipitation event, allowing us to examine the EEP events in the context of the seasonal variations of neutral and ionic species.

2. The WACCM-D simulations and ancillary data

We use the WACCM-D model jointly developed by researchers from the National Center for Atmospheric Research (NCAR) and the Finnish Meteorological Institute (FMI) (Verronen et al., 2016; Andersson et al., 2016). WACCM-D is a recent development of WACCM (version 4), a global circulation model with fully coupled chemistry and dynamics, which extends from the surface to ~145 km (with 88 pressure levels in total). The ion chemistry in WACCM-D, with 20 positive ions, 21 negative ions and 307 reactions, includes a selected subset of ionic species and reactions based on the analysis of Verronen and Lehmann (2013), and it has been shown to produce a representation of the D-region ionosphere very similar to the more comprehensive but 1-D SIC model (Verronen et al. 2016). The selected additional ionic species were added to 5 pre-existing ones in the standard WACCM (O\(^+\), NO\(^+\), O\(^{+2}\), N\(^{+2}\), N\(^+\)).

The EPP-related production of HOx and NOx which is parametrized in WACCM is, importantly, replaced by the ionic chemical reactions. The reader is referred to the two above-mentioned papers for an exhaustive description of WACCM-D, and to Marsh et al. (2007) for the MLT representation in the standard WACCM. Our simulations with WACCM-D or the standard WACCM are mostly inter-compared in terms of the ratio (WACCM-D divided by WACCM) in key minor constituents. Only in section 3.3 will the absolute differences be used. All simulations have been performed with enhanced eddy diffusion, using a Prandtl number of 2 (Garcia et al., 2014). Galactic cosmic rays are also accounted for in these simulations.

Note also that all the simulations described in the current study are in specified dynamics (SD-WACCM) mode (e.g., Marsh, 2011), i.e. nudged up to about 0.79 hPa toward the Modern-Era Retrospective Analysis for Research and Application (MERRA) reanalysis of NASA’s Global Modelling and Assimilation Office (Rienecker et al., 2011). Above this level (i.e., in the region where the ion chemistry is particularly active), SD-WACCM transitions linearly to a free running model. For brevity, we omit the “SD-“ prefix, and refer simply to WACCM and WACCM-D, although we use the specified dynamics mode throughout this paper. Our simulations were made at a horizontal resolution of 1.9°×2.5° (latitude × longitude), cover the period July 2009 to June 2010, and were initialised from existing WACCM initial states. We particularly examine the period spanning April to May 2010 using daily-mean data, although the annual cycle from July 2009 to June 2010 will also be shown using monthly-mean data.

The parameterization of auroral electron precipitation in WACCM has been supplemented by the inclusion of additional ionisation due to MEEs. The details on the derivation of the MEE fluxes and the calculation of the associated ionization rates are given in the Appendix. Having twin simulations with and without MEEs allows us to separate the contributions to the HNO\(_3\) enhancement from the D-region ion chemistry from the MEE forcing. We focus on high latitudes where EEP is concentrated, and thus undertake polar averaging over all latitudes poleward of 50° to include the full meridional extension of the auroral ovals. In total, we carried out four simulations, i.e., with and without ion chemistry (WACCM-D or WACCM), and with and without the MEE forcing.
We examine the distribution of HNO$_3$ and of the following four key ion cluster groups: the water clusters H$^+$($H_2$O)$_n$ (with $n=1,5$), NO$^+$ ($H_2$O)$_n$ (with $n=1,3$), NO$_3^-$ ($H_2$O)$_n$ (with $n=1,2$), and the nitric acid ion cluster NO$_3^-$ ($HNO_3$)$_n$ (with $n=1,2$). Our model output is for an ion cluster group, i.e. for all $n$ and not for individual cluster orders. The individual ion clusters are treated as prognostic species only in WACCM-D, and are not included in the standard WACCM. Although individual ion clusters are short-lived, their abundances adjust to the transport and chemical production or loss of their longer-lived source species. In the thermosphere, the abundances of ion clusters become very low, and it is expected that the ion chemistry is driven by the 5 primary ions already included in WACCM.

The geomagnetic storm period of April 2010 was driven by a combination of both co-rotating interaction regions associated with high-speed solar wind streams and of coronal mass ejections associated with dense transient solar wind flows. This period is also described in more detail in Smith-Johnsen et al. (2017). We use the Disturbance Storm Time (Dst) index to characterise the storm onset and evolution. The minimum of -81 nT (nanoTesla) on the 5th of April classifies it as a moderate geomagnetic storm. The recovery period is interrupted by a second moderate geomagnetic storm (with Dst of -67 nT) on the 12th of April. Further storms of similar magnitudes, in term of Dst deflections, occurred in early May and June of that year. There were no large solar proton events during the simulated year.

3. Results

3.1. Annual evolution

3.1.1 Monthly mean polar-averaged HNO$_3$ distributions

The monthly-mean, polar-averaged (zonally and over latitudes poleward of 50 degree) evolution of HNO$_3$ throughout the modelled year (July 2009 to June 2010) from 40 to 110 km altitude is shown in Figure 1 for standard WACCM and for WACCM-D, as well as their ratio (WACCM-D divided by WACCM), for both hemispheres. In both WACCM and WACCM-D, there is a seasonal march in the HNO$_3$ abundance, which culminates in the winter months (e.g., June-July in the SH, in this case July 2009 and June 2010, and December-January in the NH, in this case December 2009 and January 2010). Inspection of the ratio shown in the lowest panels reveals that the abundance of HNO$_3$ is increased by over two orders of magnitude in WACCM-D in the altitude range 55-75km, pointing to the important role of ion chemistry in producing mesospheric HNO$_3$. The ratio is most strongly amplified in April-May 2010 in both hemispheres, raising abundances from $10^{-4}$ to a few $10^{-2}$ ppb and suggesting that the EEP events may play a role then. Above 90 km, the background abundance of HNO$_3$ is very low in both models, but the increase in the ratio in April-May 2010 remains substantial throughout the lower thermosphere (over a factor 10) in both hemispheres, although weaker than for the altitudes below. We will return to this influence of the precipitation events in section 3.2. We further note that, in the upper stratosphere, there is no large increase even though the ion chemistry could have been, in principle, active in the cold winter conditions.

Figure 2 (left panel) shows the monthly-mean seasonal cycle of HNO$_3$ averaged in the polar caps (poleward of 50 degrees latitude) in both hemispheres, and averaged over an altitude range of 70-
75 km. It reveals a peak-to-peak variation in the seasonal cycle that is twice as large in the SH as in the NH, and a maximum abundance in winter (0.07 ppb in the SH and 0.035 ppb in the NH).

3.1.2 Chemical pathways for HNO₃ production

Up to now, we have not yet discussed which pathways of ion or neutral chemistry described in Verronen and Lehmann (2013) are the most important for the production of HNO₃ in WACCM or WACCM-D. In the standard WACCM, there is some weak HNO₃ production (Fig. 1) arising from the neutral reaction

\[
\text{NO}_2 + \text{OH} + \text{M} \rightarrow \text{M} + \text{HNO}_3
\]

WACCM-D however includes the key ion-ion recombination reactions between water ion clusters and the nitric acid ion clusters, that produce HNO₃:

(reaction pathway R1: ion-ion recombination reactions forming HNO₃)

\[
\text{H}^+ (\text{H}_2\text{O})_n + \text{NO}_3^-(\text{HNO}_3)_m \rightarrow (m+1) \text{HNO}_3 + n \text{H}_2\text{O}
\]

\[
\text{NO}_4^+(\text{H}_2\text{O})_n + \text{NO}_3^-(\text{HNO}_3)_m \rightarrow \text{NO} + \text{NO}_3 + m \text{HNO}_3 + n \text{H}_2\text{O}
\]

Another set of ionic reactions involved in the production of HNO₃ in the stratosphere is the conversion of N₂O₅ into HNO₃ through a hydrolysis reaction involving hydrated water clusters (Kawa et al., 1995; Kvissel et al., 2012 and references therein):

(reaction pathway R2: hydrolysis of N₂O₅)

\[
\text{H}^+ (\text{H}_2\text{O})_n + \text{N}_2\text{O}_5 \rightarrow \text{H}^+ (\text{H}_2\text{O})_{n-1}(\text{HNO}_3) + \text{HNO}_3
\]

\[
\text{H}^+ (\text{H}_2\text{O})_{n-1} (\text{HNO}_3) + \text{H}_2\text{O} \rightarrow \text{H}^+ (\text{H}_2\text{O})_n + \text{HNO}_3
\]

This reaction pathway (R2) sequesters NOx from the N₂O₅ reservoir into the longer-lived HNO₃ reservoir, and mostly plays a role in the stratosphere, since the abundance of N₂O₅ falls rapidly with height in the upper stratosphere and mesosphere.

Figure 3 compares the winter-mean vertical profiles of several (neutral) nitrogen species between WACCM and WACCM-D, in each hemisphere. We first remark that the winter increase in WACCM-D is found not only in HNO₃ but also in NO₃ and to a much smaller extent in N₂O₅. HNO₃ is enhanced above 40 km altitude. The NO₃ production is not very efficient in the mesosphere and its background level is very low. As seen on Fig. 3, NO₃ is only very weakly enhanced below 65 km and strongly enhanced between 70 and 85 km. Ion-ion recombination reactions involving the several types of water ion clusters can explain - through ion chemistry - the increase in NO₃ by up to two orders of magnitude (from 10⁻⁶ to 10⁻⁴ ppb) in WACCM-D compared to WACCM. Such reactions include

\[
\text{H}^+ (\text{H}_2\text{O})_n + \text{NO}_3^-(\text{H}_2\text{O})_m \rightarrow \text{H} + \text{NO}_3 + (n+m) \text{H}_2\text{O}
\]

\[
\text{NO}_4^+(\text{H}_2\text{O})_n + \text{NO}_3^-(\text{H}_2\text{O})_m \rightarrow \text{NO} + \text{NO}_3 + (n+m) \text{H}_2\text{O}
\]

\[
\text{NO}^+ + \text{NO}_3^-(\text{H}_2\text{O})_n \rightarrow \text{NO} + \text{NO}_3 + n \text{H}_2\text{O}
\]

Ion chemistry also enhances the NOx production compared to the parameterisation used in standard WACCM (Andersson et al. 2016), which is another factor relevant to the increased NO₃
through neutral chemistry. \( \text{N}_2\text{O}_5 \) also increases only very weakly below 65 km; the increase is larger above 70 km, though still smaller than that for \( \text{NO}_3 \). The increase in this case is consistent with the neutral reaction

\[
\text{NO}_2 + \text{NO}_3 + \text{M} \rightarrow \text{M} + \text{N}_2\text{O}_5
\]

and the \( \text{N}_2\text{O}_5 \) increase is limited to the layer where \( \text{NO}_3 \) is produced by the ion-ion recombination in WACCM-D. In short, the hydrolysis of \( \text{N}_2\text{O}_5 \), pathway R2, is unimportant for the seasonal production of \( \text{HNO}_3 \) due to the small abundance of \( \text{N}_2\text{O}_5 \) in the mesosphere, itself tied to \( \text{NO}_3 \) (Fig. 3).

3.1.3 Monthly mean polar-averaged distributions of ion clusters

The corresponding seasonal evolution of the four ion cluster species is shown in Fig. 4. The abundances vary with season and height. The \( \text{H}^+(\text{H}_2\text{O})_n \) and \( \text{NO}^+(\text{H}_2\text{O})_n \) water clusters have a maximum abundance in summer. This seasonal-dependence and its height variation follows the seasonal march of water vapour abundance as indicated by two white contours on Fig. 4, corresponding to 2 and 5 ppmv, with the summer-to-winter pole meridional circulation in the mesosphere bringing up air enriched in water vapour at high latitudes in summer, and bringing down dry air in winter. The \( \text{NO}_3^-(\text{H}_2\text{O})_n \) water cluster and the nitric acid cluster \( \text{NO}_3^-(\text{HNO}_3)_n \) tend to maximize in winter, indicating that the abundance of \( \text{NO}_3^- \) is, in this case, the limiting factor. A higher abundance of \( \text{NO}x \) has been previously reported in the winter high latitudes due to the descent from the MLT (e.g., Randall et al., 2007). This coincidence in time between higher abundance of precursory \( \text{NO}x \) and the secondary winter maximum in \( \text{H}^+(\text{H}_2\text{O})_n \), weaker than the summer one and occurring when abundance of water vapour is low, is suggestive of a role for conversion of \( \text{NO}^+(\text{H}_2\text{O})_n \) to \( \text{H}^+(\text{H}_2\text{O})_n \) (Verronen and Lehmann, 2013).

3.2 Changes around the April and May geomagnetic storms

3.2.1 Daily polar-averaged \( \text{HNO}_3 \) distributions

The period of April and May 2010 is characterised by large enhancements in mesospheric and lower thermospheric \( \text{HNO}_3 \) in WACCM-D, following the Dst negative excursions that mark the occurrences of the storms. The daily-averaged, polar-averaged abundances in WACCM-D (Fig. 5) are higher than in WACCM by over two orders of magnitude. Close inspection of the ratio indicates that the short-lived relative enhancements clearly appear in bursts. These bursts are tied to the Dst fluctuations but also, in the mesospheric layer at 50-70 km in particular, the daily-varying increases related to the geomagnetic storms are superimposed on a seasonal cycle variation. In the SH, the daily-varying enhancements also occur when the \( \text{HNO}_3 \) abundance is increasing as the winter approaches, while it is decreasing in the NH (see Fig. 1).

Returning to Fig. 2 (right panel), we note that the \( \text{HNO}_3 \) increase due to the April 2010 EEP event in WACCM-D is much smaller than the corresponding increase after the January 2005 SPE, discussed by Andersson et al. (2016). They found an increase in \( \text{HNO}_3 \) of 0.5-1.0 ppb near 70-75 km during the SPE (their Figure 9, top panel) in the NH. By contrast, Fig. 2 (right panel) shows that, in the SH, the increase during the EEP events has a maximum value of 0.075 ppb near 70-75 km in WACCM-D, at the peak of the events. It is hence a factor 5 to 10 smaller than for the SPE. In the NH, the \( \text{HNO}_3 \) increase is only about 0.005 ppb, i.e. 20 times smaller than in
the SH. It is important to note that SPEs are comparatively rare events, while MEE electron precipitation is considerably more common. The EEP event occurred at a time of year (April) when background abundances of HNO₃ are low in both hemispheres (Fig. 2, left panel), but the abrupt increase in the austral polar cap during the EEP event brings HNO₃ close to its seasonal maximum.

3.2.2 Daily polar-averaged distributions of ion clusters.

The corresponding day-to-day evolution of the abundance of the 4 ion cluster species (Fig. 6) is marked by an abrupt increase in most of the clusters following the Dst negative excursions indicating storm onsets. While Fig. 4 had shown that the H⁺(H₂O)ₙ abundance followed the seasonal H₂O cycle, it is clear from Fig. 6 that the rapid enhancements during the EEP events occur in the declining phase of water vapour abundance in the SH or in its slowly increasing phase in the NH (see the white contours). Enhancements of hydrated water cluster ions during the EEP event are due to higher ionisation and higher abundances of ions, including precursory NO⁺(H₂O)ₙ ion clusters or also oxygen ion clusters O₂⁺(H₂O)ₙ, followed by the conversion to H⁺(H₂O)ₙ (Verronen and Lehmann, 2013).

In Fig. 7, we also compare the changes in WACCM-D during the EEP event, i.e., contrasting the quiet and active periods (April 1-3 and April 6-8, respectively), for the aforementioned three neutral species and the four key ion clusters. Starting with the neutral species, the changes during the event are qualitatively similar to those seen during the seasonal cycle. The layer where HNO₃ increases is somewhat narrower for the EEP-related changes than for the changes due to the seasonal cycle, since the latter are also influenced by transport, due to the longer time scales involved. Since the event took place in April, the abundance of N₂O₅, which is easily photodissociated, is smaller in the NH, which is more sunlit in boreal spring conditions. Again, the N₂O₅ change is constrained by the increase in NO₃ produced by the ion cluster chemistry during the EEP event. Turning now to the ion clusters and their changes between the quiet and active period, we again notice how their respective abundances decrease sharply with increasing altitude, reflecting the decreasing availability of H₂O or HNO₃, as seen earlier (Fig. 4). Above 90 km, the NO⁺(H₂O)ₙ ion cluster becomes the most abundant, reflecting the primary importance of the NO⁺ ion in the thermosphere. It is apparent that the additional production of HNO₃ and NO₃ between the quiet and active period is tied to the enhanced abundances of the various ion clusters in WACCM-D, which recombine in ion-ion reactions. For example, NO₃ is decreasing sharply above 80 km along with the two ion clusters tied to the negative ion NO₅⁻. Also, the lower boundary of the HNO₃ increase near 55 km is tightly constrained by the altitude where the abundance of hydrated ion water clusters starts to increase due to the influence of energetic electrons (this will be even clearer in Fig. 12). In short, these figures are consistent with ion cluster recombinations (R1) playing the dominant role in the formation of HNO₃ during the EEP event.

3.2.3 Geographical maps of nitric acid and ion cluster abundances

We focus hereafter on the changes in the geographical distribution of HNO₃ at 70 km between the pre-storm quiet period (a 3-day period, April 1-3) and the active storm period (a 3-day period, April 6-8). These are shown as geographic maps in Figs 8 and 9 for the NH and the SH, respectively. We chose the early April storm since it is easier to contrast to the quiet pre-storm conditions, but the same changes are seen in the simulations for the May event (not shown). It is
apparent that there is already some enhancement in WACCM-D compared to WACCM withinmost of the whole polar cap even during the quiet period, related to the seasonal cycleenhancement in WACCM-D, albeit at much lower levels than during the time of the storm. Thechanges between the active and quiet periods are already present in WACCM (see Fig. 5), butthe amplification is stronger and much more widespread in WACCM-D, up to $10^2$ rather than to$10^3$ ppb. In the active period, HNO$_3$ increases considerably along the oval regions, i.e., in theregion symmetric in geomagnetic coordinates which experiences enhanced ionisation (seeAppendix). The equatorward expansion of the ovals at the initiation of the storm can be noticedin the enhancements on the equator-side of the pre-storm oval edges (Figs. 8-9). The increase isstronger in the SH, as had been shown earlier in Fig. 2.

The geographical distributions of the 4 ion cluster species at 70 km (Figs. 10 and 11) readilyillustrate the enhancements of the ion cluster distributions along the ovals during the EEP event.That is, the chemical effect of EEP is not limited to NOy species and the enhancements in ionclusters play an important role as outlined in section 3.1.2. The distribution of the ion clustercontaining the NO$_3^-$ ion is more complex than the water ion clusters. The ion clusters are short-lived but their distributions reflect the changing distributions of precursory neutral species, likeNO$_2$ in the case of the NO$_3^-$ ion clusters. It is also apparent from Figs. 8-11 that, while HNO$_3$ andion clusters remain close to their production around the auroral oval in the NH, they are morereadily redistributed within the polar cap in the SH. The inter-hemispheric differences, namelythat the changes are confined to a narrow latitude band characteristic of the auroral oval in theNH, but spread into the polar cap in the SH, can be explained by the larger oval offset (i.e.,difference in the zonal direction between the magnetic and geographic coordinate systems) in theSH. Hence, in the SH, there is a tendency for the zonal winds to rapidly redistribute species withisopleths along magnetic coordinates. Also, the NH is more sunlit at this time of the year and NOand HNO$_3$ are more readily photolysed. The transport by the mean meridional circulation isstronger in April-May in the SH (during austral fall) than in the NH (during boreal spring) (e.g.,Smith et al., 2015). The inter-hemispheric differences during the EEP event, with stronger polarcap-averaged abundances of HNO$_3$ or ion clusters in the SH, can also be seen in the profilesshown in Figs. 7.

3.3 Role of the MEE ionization

Since the aim of this paper was to examine an energetic electron precipitation event, it appears sensible to include the forcing by medium-to-high energy electrons (MEE). The previous figureshence showed results from WACCM or WACCM-D simulations with the MEEs included. In thissection, we nevertheless compare those simulations with simulations where the MEE forcing isremoved, leaving only auroral electron precipitation, in order to delineate more specifically theeffect of this MEE precipitation.

Covering the period around the EEP event, Figure 12 shows the April-mean zonally averagedmeridional-height cross sections of HNO$_3$ from both WACCM and WACCM-D, with andwithout the inclusion of MEEs, as well as their absolute differences. Hence, the differencesbetween the plots along a row (i.e., (b)-(a) or (e)-(d)) indicate the change due to the introductionof the MEEs in WACCM or WACCM-D, while the differences along a column (i.e., (d)-(a) or(e)-(b)) indicate the changes due to the introduction of the D-region ion chemistry. From Fig.12c, it is clear that, in the standard WACCM, the MEEs produce little change in HNO$_3$ in the
upper stratosphere or MLT region. It is only with the inclusion of the ion chemistry in WACCM-D that the EEP-induced production of HNO₃ is amplified (Fig. 12f). Although there is already a mesospheric and thermospheric HNO₃ increase in WACCM-D without the MEEs, it is further increased by the inclusion of that precipitation. This is an indication that energetic electrons, which penetrate deeper into the mesosphere region, further trigger the ionic reactions which produce HNO₃. Figure 12i contrasts the simplest and the most elaborate model, i.e., the standard WACCM without MEE and WACCM-D with MEE. Figure 12 hence stresses the important role played by MEEs during the event: in the absence of MEEs, the HNO₃ enhancements in WACCM-D are much weaker. To further stress the important triggering role played by particle precipitation and by MEE in particular on the HNO₃ production during the event, Fig. 13 compares the two versions of WACCM-D, with and without MEE, at the peak of the EEP event (April 6-8). Importantly, the abundances of the four key ion clusters clearly increase in the mesosphere with the inclusion of the MEEs above 55 km, which is consistent with the strong increase of the HNO₃ production by MEE demonstrated in Fig. 12.

Finally, returning to Fig. 2 (left panel; red lines), we can see that the forcing by MEEs is strengthening the seasonal cycle in both hemispheres outside of the period of large negative Dst fluctuations in April and May 2010.

4. Summary

In summary, EPP induces the formation of ion clusters, and the ion-ion recombinations (reaction pathway R1) between these clusters lead to the production of HNO₃. Our study of an EEP event indicates that the effect is 5 to 10 times smaller than during a SPE event, but the event occurred at a time of year when the seasonally varying background of HNO₃ is low. Our study also confirms the importance of the ion-ion recombination pathway (R1) over the hydrolysis of N₂O₅ (R2), as shown by Verronen et al. (2001) during SPEs. NO₃ is also produced by similar ion-ion recombinations. There is a seasonal component of this HNO₃ production, independent of the EEP event, which is stronger in WACCM-D compared to WACCM due to the background presence of ion clusters, and which is stronger in WACCM-D with the inclusion of MEE. Superposed on this seasonal cycle, the EEP event considerably augments the production of HNO₃ raising its abundance by two order of magnitude from 10⁻³ to a few 10⁻² ppb. This effect is enhanced by the inclusion of the MEE, allowing more production of the necessary ion clusters down to 55 km in the latitude bands characteristic of the auroral ovals.

Three-dimensional middle atmospheric modelling with ion chemistry models such as WACCM-D is in its infancy, and only the aforementioned handful of studies have been published in the last year. Such model simulations with WACCM-D open new possibilities to study the connection between the neutral middle-atmosphere and the D-region ionosphere. It has been already shown by Andersson et al. (2016) that the ion chemistry in WACCM-D lead to a better agreement with satellite observations of HNO₃, HOx and NOx during SPEs. Further studies of the short and long-term variability in mesospheric HNO₃ using satellite observations (e.g., from MLS, Michelson Interferometer for Passive Atmospheric Sounding (MIPAS), Atmospheric Chemistry Experiment-Fourier Transform Spectrometer (ACE-FTS), or other instruments) would be warranted in order to support the conclusions of this model study. This would however require that satellite retrievals are improved and validated in the mesosphere. It should be also
noted that WACCM-D ion chemistry can provide a realistic electron density also in the lower ionosphere (Verronen et al., 2016), so that comparison against ionospheric observations such as ground-based radars can be used to study, e.g., the quality of the simulated EPP forcing. Further work with WACCM-D can be made to test our overall understanding of important ions and their reactions, both in the MLT and the stratosphere.

Appendix: Producing the MEE forcing

The energetic electron precipitation (EEP) forcing applied in the current study was developed as part of an effort to include EEP in climate models, such as WACCM. The approach used here is based on direct satellite measurements. By contrast, recent efforts undertaken to produce the EEP forcing as part of the solar forcing dataset recommended for CMIP6, as described in van de Kamp et al. (2016) or Matthes et al. (2017), rely on a proxy-based parametrization in order to extend the series back in time, prior to the satellite era.

The energetic electron precipitation flux was determined from the experimental measurements made from 1998-2012 by the Space Environment Monitor 2 (SEM-2) instrument packages onboard the constellation of Polar Orbiting Environment Satellites (POES). The POES are a long-lived series of spacecraft in Sun-synchronous high-inclination orbits at roughly 800-850 km altitude. Included in the POES SEM-2 instrument is the Medium Energy Proton and Electron Detector (MEPED) and the Total Energy Detector (TED) that together monitor electron fluxes over the nominal energy range from 50 eV to 2.5 MeV (Evans and Greer, 2004; Rodger et al., 2010a). Here we focus on the electron fluxes observed by the 0° MEPED telescope, as this principally measures electrons with pitch angles inside the Bounce Loss Cone (Rodger et al., 2010b), which will be lost locally on short time scales (i.e., on the order of seconds). Observations from this telescope can provide an indication of the EEP input into the atmosphere at a location joined by the geomagnetic field line to the satellite in question. During the period studied here, up to 6 distinct SEM-2 carrying satellites were operational simultaneously (see Table 2 of Neal et al. (2013) for a list with summary information), each of which had orbital periods of ~100 min.

The MEPED electron observations are of integral electron counts for the nominal energy ranges >30 keV (e1), >100 keV (e2), and >300 keV (e3). Previous studies have identified significant contamination in the electron channels by protons with energies of hundreds of keV (Yando et al., 2011). Unfortunately, such protons can reach considerable levels during geomagnetic storms, which is also when EEP is likely to be most active. We correct for this proton contamination by applying a NOAA-developed algorithm (Lam et al., 2010), validated by Whittaker et al. (2014). In addition, solar proton events and the high-energy protons in the South Atlantic Magnetic Anomaly are known to swamp the electron detectors, leading to no meaningful electron measurements. Hence, all solar proton event periods, as well as observations from inside and around the South Atlantic Magnetic Anomaly, are excluded. We note that for the period studied in this paper, there were no data excluded due to solar proton events.

The original MEPED electron observations have 2 s resolution. In order to produce long-term EEP fluxes, the electron observations from each integral channel were combined from all operational POES instruments by zonally averaging the measurements in geomagnetic coordinates with 3 hour time resolution and 0.5 \( L \) resolution (where \( L \) is the McIlwain \( L \)-
parameter (McIlwain, 1961)). We restrict ourselves to the $L$ range from 2.25-9.75, which encompasses the outer radiation belt. It has previously been shown that power-laws are an accurate representation of the EEP flux spectrum, i.e., through a comparison of high energy resolution DEMETER electron flux observations with POES MEPED measurements (Whittaker et al., 2013). Hence, we fit a powerlaw spectrum to the three $0^\circ$ electron telescopes (e1, e2, e3) to obtain the energy spectral gradient ($k$) for the precipitating electrons.

Unfortunately, the small angular size of the MEPED/POES telescope results in low sensitivity at lower flux magnitudes (Yando et al., 2011), which causes the MEPED electron observations to have a comparatively high minimum detectable flux ($\sim 10^2$ el.cm$^{-2}$ s$^{-1}$ sr$^{-1}$ in each integral channel). Due to experimental noise, periods with lower EEP fluxes may appear to have near constant EEP at this minimum detectable level, although they have been shown to be inconsistent with other experimental datasets (e.g., Neal et al., 2015). So as not to bias the fluxes during the relatively common situation of geomagnetic quiet times (when EEP levels are likely near-zero), we set all EEP flux magnitudes $<250$ cm$^{-2}$ s$^{-1}$ sr$^{-1}$ to zero. The near noise-level fluxes can also produce artificially hard power law spectra; we therefore limit $k<-1$, and set any periods with $k\geq-1$ to $k=-1$. Finally, we follow Rodger et al. (2013) and remove all electron observations when the MEPED P7 omni-directional detector reports $>36$ MeV protons, i.e. set them to zero. This should suppress the impact of solar proton event contamination. However, it is important to note that this means the EEP flux representation provided here will not be fully accurate during solar proton events, where no meaningful experimental measurements of EEP are available from POES.

The combination of these processes should ensure that unrealistically high levels of EEP are excluded during quiet times or during solar proton events, an issue that has been identified in other EEP products derived from POES electron observations (e.g., Neal et al., 2015).

The power-law fitted EEP parameters were then used to determine ionization rates assuming the EEP had a differential power-law flux spectrum covering the energy range from 50 keV - 2 MeV using 78 logarithmically spaced bins. The power-law assumption for the spectrum is the same as the one described by van de Kamp et al (Section 2.1, 2016). The ionization rate calculation is based on a continuously-slowing-down approximation and a normalized energy dissipation distribution function for electrons; this method is described in detail by Rees (1989). A prior WACCM simulation was used to provide daily, zonal mean neutral background data for the ionization rate calculation, specifically the concentrations of the main atmospheric constituents N$_2$ and O$_2$, and O. The ionization rates were calculated in 3-hour time resolution for each of the L shell bins (latitudes), after the differential electron fluxes were integrated over pitch angles 0-80$^\circ$ and azimuth angles 0-360$^\circ$ assuming an isotropic angular distribution. The ionization rates were calculated for the WACCM altitude (km) grid which changes slightly from day to day but corresponds to a fixed pressure level grid. The L shell-dependent ionization rates were then converted to magnetic latitude and, with the assumption of uniformity on magnetic longitude, projected onto the WACCM geographic (latitude-longitude) grid.

**Acknowledgments and Data**

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Terrestrial Coupling through High Energy Particle Precipitation in the Atmosphere: a Norwegian contribution”. YO is also supported by the Norwegian Research Council under contracts 223252/F50. DRM is supported in part by NASA LWS Grant NNX14AH54G. The National Center for Atmospheric Research (NCAR) is sponsored by the U.S. National Science Foundation (NSF). PTV was supported by the Academy of Finland through the project #276926 (SECTIC: Sun-Earth Connection Through Ion Chemistry). DRM, PTV, MAC, and CJR would like to thank the International Space Science Institute, Bern, Switzerland for supporting the project "Quantifying Hemispheric Differences in Particle Forcing Effects on Stratospheric Ozone" (PI: D. R. Marsh). WACCM is a component of the Community Earth System Model (CESM), which is supported by NSF and the Office of Science of the U.S. Department of Energy. WACCM is an open-source community model. All model simulations were performed at NCAR, are archived on the NCAR High Performance Storage System, accessible via the Earth System Grid (https://www.earthsystemgrid.org/), and are also accessible at the Norwegian Center for Research Data (http://www.nsd.uib.no/nsd/english/index.html).

References


Evans, D. S., and M. S. Greer (2004), Polar Orbiting environmental satellite space environment monitor - 2 instrument descriptions and archive data documentation, NOAA technical Memorandum version 1.4, Space Environment Laboratory, Colorado.


Captions

FIGURE 1: Monthly-mean polar cap average of HNO$_3$ (in ppb, log10 scale). Period is from July 2009 until June 2010, for both the southern hemisphere (a,c,e, over latitudes poleward of 50°S) and northern hemisphere (b,d,f, over latitudes poleward of 50°N), from 40 to 110 km. (a,b) standard WACCM, (c,d) WACCM-D, and (e,f) the ratio (WACCM-D / WACCM), also in log scale. All simulations are with MEEs included.

FIGURE 2: Time evolution of HNO$_3$ (in ppb). (left) Monthly-mean annual (July 2009-June 2010) evolution of HNO$_3$ in WACCM-D with MEE included, for the southern hemisphere (black dashed line, over latitudes poleward of 50°S) and the northern hemisphere (black solid line, over latitudes poleward of 50°N) averaged over the 70-75 km layer. Corresponding results form simulations without MEEs are also shown (red lines). (right) Same but showing daily evolution in April and May 2010 for simulations with MEEs included.

FIGURE 3: Winter-averaged vertical profiles of the three neutral species (in ppb, log10 scale). Shown are the three neutral species HNO$_3$, NO$_3$ and N$_2$O$_5$ in WACCM (dashed lines) and WACCM-D (solid lines) in JJA for the southern hemisphere (left column, latitudes poleward of 50°S), and in DJF for the northern hemisphere (right column, latitudes poleward of 50°N). All simulations are with MEEs included.

FIGURE 4: Monthly-mean polar cap average of key ion clusters (in ppb, log10 scale). (a,b) NO$_3^-$ (H$_2$O)$_n$, (c,d) H$^+$ (H$_2$O)$_n$, (e,f) NO$^+$ (H$_2$O)$_n$ and (g,h) NO$_3$ (HNO$_3$)$_n$ from July 2009 until June 2010, for both the southern hemisphere (a,c,e,g, over latitudes poleward of 50°S) and northern hemisphere (b,d,f,h, over latitudes poleward of 50°N) in WACCM-D. The two white lines are the contours of 2 and 5 ppm of H$_2$O (upper and lower contours, respectively). All simulations are with MEEs included.

FIGURE 5: Daily-mean polar cap average of HNO$_3$ (in ppb, log10 scale). The period is April-May 2010, for both the southern hemisphere (a,c,e, over latitudes poleward of 50°S) and northern hemisphere (b,d,f, over latitudes poleward of 50°N) from 40 to 110 km. (a,b) standard WACCM, (c,d) WACCM-D, and (e,f) the ratio (WACCM-D / WACCM), also in log scale. (g,h) the Dst-index for the same period to highlight the timing of geomagnetic activity. The two white lines are the contours of 2 and 5 ppm of H$_2$O. All simulations are with MEEs included.

FIGURE 6: Daily-mean polar cap average of key ion clusters (in ppb, log10 scale). (a,b) NO$_3^-$ (H$_2$O)$_n$, (c,d) H$^+$ (H$_2$O)$_n$, (e,f) NO$^+$ (H$_2$O)$_n$ and (g,h) NO$_3$ (HNO$_3$)$_n$ for April-May 2010, for both the southern hemisphere (a,c,e,g, over latitudes poleward of 50°S) and northern hemisphere (b,d,f,h, over latitudes poleward of 50°N) in WACCM-D. (i,j) the Dst-index for the same period to highlight the timing of geomagnetic activity. The two white lines are the contours of 2 and 5 ppm of H$_2$O. All simulations are with MEEs included.

FIGURE 7: Vertical profiles of abundance of ion clusters and neutral species (in ppb, log10 scale). Three-day averages in quiet conditions (April 1-3, dashed lines) and active conditions (April 6-8, solid lines) (southern hemisphere in left column, latitudes poleward of 50°S, and northern hemisphere in right column, latitudes poleward of 50°N) of the vertical profiles of
abundance of the four key ion clusters and the neutral species HNO₃, NO₃ and N₂O₅ in WACCM-D. All simulations are with MEEs included.

FIGURE 8: Geographic maps of HNO₃ (in ppb, log10 scale) over the northern hemisphere. Maps correspond to 3-day averages at 70 km, in quiet (left column for April 1-3) and active (middle column for April 6-8) conditions. Upper row shows the standard WACCM, second row shows WACCM-D. In this figure and all following figures with maps, latitude circles are at 45°, 60° and 75°, and zero degree longitude is at the bottom. All simulations are with MEEs included.

FIGURE 9: Geographic maps of HNO₃ (in ppb, log10 scale) over the southern hemisphere.
(As in Fig. 8)

FIGURE 10: Geographic maps of key ion clusters (in ppb) over the northern hemisphere. Shown are NO₃⁻(H₂O)ₙ, H⁺(H₂O)ₙ, NO⁺(H₂O)ₙ and NO₃⁺(HNO₃)ₙ from WACCM-D at 70 km, in quiet (left column for April 1-3) and active (right column for April 6-8) conditions. All simulations are with MEEs included.

FIGURE 11: Geographic maps of key ion clusters (in ppb) over the southern hemisphere.
(As in Fig. 10).

FIGURE 12: April-mean latitude/height zonal-mean cross-sections of HNO₃ (in ppb, log10 scale) for all simulations. (a,b,c) standard WACCM without and with MEE and their absolute difference (the later minus the former), respectively. (d,e,f) standard WACCM without and with MEE and their absolute difference (the later minus the former), respectively. (g,h) absolute difference between WACCM-D and WACCM without and with MEE. (i) absolute difference between WACCM-D with MEE and WACCM without MEE. Difference are expressed in ppb.

FIGURE 13: Vertical profiles of abundance of ion clusters (in ppb, log10 scale) in simulations with and without MEEs. Profiles of the four key ion clusters in WACCM-D correspond to 3-day averages during the active period (April 6-8) for southern hemisphere (left, latitudes poleward of 50°S) and northern hemisphere (right, latitudes poleward of 50°N), with MEEs included (solid lines) and without (dashed lines). Colour coding for the ion clusters is the same as in Fig. 7.
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FIGURE 6: Daily-mean polar cap average of key ion clusters (in ppb, \( \log_{10} \) scale). (a,b) \( \text{NO}_3^- \) \((\text{H}_2\text{O})_n\), (c,d) \( \text{H}^+(\text{H}_2\text{O})_n\), (e,f) \( \text{NO}^+(\text{H}_2\text{O})_n\) and (g,h) \( \text{NO}_3^-\text{(HNO}_3\text{)}_n\) for April-May 2010, for both the southern hemisphere (a,c,e,g, over latitudes poleward of 50°S) and northern hemisphere (b,d,f,h, over latitudes poleward of 50°N) in WACCM-D. (i,j) the Dst-index for the same period.
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Figure 1.
Figure 2.
Figure 3.
Figure 5.
Altitude [km]

SH quiet vs active

NH quiet vs active

log 10 [ppb]

April 1.-3.

April 6.-8.

HNO3
NO3
N2O5
H+(H2O)n
NO+(H2O)n
NO3-(H2O)n
NO3-(HNO3)n
April 1–3, 2010

April 6–8, 2010

WACCM

WACCM-D

log10 [ppb]

log10 [ppb]
Figure 10.
Figure 11.
Figure 12.