Role of excited N₂ in the production of nitric oxide

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Excited N₂ plays a role in a number of atmospheric processes, including auroral and dayglow emissions, chemical reactions, recombination of free electrons, and the production of nitric oxide. Electron impact excitation of N₂ is followed by radiative decay through a series of excited states, contributing to auroral and dayglow emissions. These processes are intertwined with various chemical reactions and collisional quenching involving the excited and ground state vibrational levels. Statistical equilibrium and time step atmospheric models are used to predict N₂ excited state densities and emissions (as a test against previous models and measurements) and to investigate the role of excited nitrogen in the production of nitric oxide. These calculations predict that inclusion of the reaction N₂[A³Σ_u⁺] + O, to generate NO, produces an increase by a factor of up to three in the calculated NO density at some altitudes.


1. Introduction

[2] Radiative decay of the excited states of N₂ has long been identified as the source of many auroral emissions, such as the Vegard-Kaplan (VK) bands from transitions between vibrational levels in A³Σ_u⁺ → X¹Σ_g⁻. It has been proposed [Swider, 1976; Gordiyets, 1977] that N₂[A³Σ_u⁺] and N₂[X¹Σ_g⁻ (ν' > 11)] react with O atoms to produce nitric oxide. These are potentially important chemical processes because although a minor constituent, NO is a vital component of the Earth’s atmosphere due to the critical role it plays in determining the thermal balance in the lower thermosphere [Killeen et al., 1997] and in the odd-nitrogen chemistry in the ionosphere [Strickland et al., 1999]. Its density is highly variable [Adler-Golden et al., 1999] and the production mechanisms leading to enhanced NO densities in aurora are complex and not fully understood [Solomon and Barth, 1999]. Similarly, NO production by sunlight is a subject of ongoing research [Barth and Bailey, 2004], as are the reaction rates involved [Duff et al., 2005].

[3] Motivated by the large variability in the measured and inferred auroral NO densities reported in the early 1970s, Hyman et al. [1976] examined the NO chemistry to determine upper limits on the NO concentration and its production rate. They concluded that the NO concentration ([NO]) cannot exceed 0.1 × [O₂] and called attention to the sensitivity of [NO] to the N(²D) and N(⁴S) atoms produced in the odd-nitrogen chemical cycle.

[4] Swider [1976] originally proposed the reaction N₂[A³Σ_u⁺] + O → NO + N(²D) as a source of NO, and it was included in a model of NO production by O’Neil et al. [1979]. In seeking to explain observations of higher concentrations of NO, Gordiyets [1977] showed that the reaction N₂[X¹Σ_g⁻ (ν' > 11)] + O → NO + N can also be an important source of NO and Vlasov et al. [1980] proposed it as the main source of NO in strong aurora. Its rate is quoted as 10⁻¹¹ cm³ s⁻¹ by Aladjev and Kirillov [1995], who ascribe it to Rusanov and Fridman in a publication in Russian. This value is larger than used in some recent work [Pintassilgo et al., 2005].

[5] Sharma et al. [2001] have shown that the enhancement of NO infrared emissions observed in the 6.8–10.8 μm band in the 110–130 km region during an auroral event can be explained by the prompt chemiluminescent emission from the nascent NO formed by N(²D) + O₂ → NO + O, which is initiated by local auroral energy deposition producing N atoms. In their model the chemical reactions responsible for producing NO in an aura of characteristic energy 8 keV, were determined to have the relative ranking:

N(²D) + O₂ → NO + O 172
N₂[A³Σ_u⁺] + O → NO + N(²D) 7.4
N(⁴P) + O₂ → NO + O 5
N(⁴S) + O₂ → NO + O 1

Thus Sharma et al. find that the production and loss of N(²D) is the critical element in the auroral production of NO, but note that the N₂[A³Σ_u⁺] reaction is included.

[6] In the current work we utilize a recently developed model that combines the radiative cascade processes in excited nitrogen with a simple photochemical description of atmospheric species [Campbell et al., 2006]. This model has been shown to predict intensities of the Lyman-Birge-Hopfield (transitions between vibrational levels in a¹Π_g →
X\(^{1}\Sigma_g^+)\) and VK bands [Campbell et al., 2005] in reasonable agreement with rocket and satellite measurements. Comparisons with measurements and other calculations are made here, to verify our model for its estimates of N\(_2[\Lambda^3\Sigma_u^+]\) densities. The predictions of NO density from our model are compared with those from other models and with measurements to show that our model is sufficiently accurate to be used to investigate the role of N\(_2[\Lambda^3\Sigma_u^+]\) and N\(_2[X^1\Sigma_g^+ (\nu' > 11)]\) in the production of NO. We find that in both equatorial daytime and high-latitude auroral conditions, N\(_2[\Lambda^3\Sigma_u^+]\) makes a significant contribution to the NO density, while the contribution from N\(_2[X^1\Sigma_g^+ (\nu' > 11)]\) is relatively small. As the inclusion of the N\(_2[\Lambda^3\Sigma_u^+]\) + O reaction will reduce the calculated 557.7 nm emission in aurora, we confirm that our predicted 557.7 nm emission is consistent with auroral measurements. The effect of atmospheric winds on the calculated altitude profile of NO density is also investigated.

2. Details of the Model

[7] The computational model has been extended to altitudes below 120 km using:

\[
F_h = F_{120} e^{-0.1(120-h)}
\]

where \(F_h\) is the calculated auroral flux at height \(h\) and \(F_{120}\) is the flux at 120 km [Campbell et al., 2006]. Equation (1) was chosen to match our predictions of O(\(^1\Sigma\)) excitation to those of Sharp et al. [1979]. Below 100 km the auroral flux is reduced linearly to zero at 89 km, similar to the lower height limit in Figure 4.28 of Vallance Jones [1974].

[8] Molecular and eddy diffusion of N\(^2\Sigma\) and NO have been incorporated: Coefficients of molecular diffusion between pairs of constituents [Pavlov, 1981] are combined [Pavlov, 1988] to give molecular diffusion coefficients. For the equatorial case, measured eddy diffusion coefficients [Sasi and Vijayan, 2001] for September are used for the altitude range 70–90 km. Above 100 km the often-used value of \(1 \times 10^6\) cm\(^2\) s\(^{-1}\) [e.g., Barth, 1992] is applied, with linear interpolation between this value and that of Sasi and Vijayan at 90 km used for the range 90–100 km. For the auroral case, eddy diffusion coefficients for 20 September are read from a graph of calculated results at 61°N [Garcia and Solomon, 1985]. The eddy and molecular diffusion coefficients are used to calculate diffusion rates using equation (E-10) of Bailey et al. [2002].

[9] Updated rates for the reactions N(\(^3\)D) + O\(_2\) \(\rightarrow\) NO + O [Duff et al., 2005], N(\(^2\)S) + O\(_2\) \(\rightarrow\) NO + O and N(\(^4\)S) + NO \(\rightarrow\) N\(_2\) + O [Pintassilgo et al., 2005] are used. Otherwise the photochemical model is as described previously [Campbell et al., 2006].

[10] A statistical equilibrium calculation is employed to determine the populations of the excited states of N\(_2\), which are then “imported” into a time step simulation [Campbell et al., 2006] that is used to calculate the NO density. In the latter the incremental change in density \(\Delta[z]\) for each species \(z\) is calculated at successive time steps of \(\Delta t\) as \(\Delta[z] = R[z][y] \Delta t\), where \(R\) is the rate for the reaction \(x + y \rightarrow z\). Calculations are made at either 0.5 s intervals in the auroral case or 1.0 s in the equatorial case. These intervals are too long for some fast reactions, for which an equilibrium calculation is performed at each time step.

3. Results and Discussion

[11] The current model is verified by comparison of its predictions of N\(_2[\Lambda^3\Sigma_u^+]\) densities, N\(_2[X^1\Sigma_g^+]\) production, \(A^3\Sigma_u^+ \rightarrow X^1\Sigma_g^+\) emissions and NO production rates with other models and measurements. It is used to calculate NO densities in conditions for which measurements are available. The calculation is made with and without the NO production due to N\(_2[\Lambda^3\Sigma_u^+]\) and N\(_2[X^1\Sigma_g^+]\) to assess its importance, including the implications for calculations of 557.7 nm emission. The effect of vertical winds in these calculations is assessed by a sample calculation.

3.1. N\(_2[\Lambda^3\Sigma_u^+]\) in Daytime and Aurora

[12] Figure 1 shows the predicted number density of N\(_2[\Lambda^3\Sigma_u^+]\) produced by an IBC II' aurora and by sunlight for a solar zenith angle of 0°. It shows that the densities due to the aurora are greater than those due to sunlight, ranging from a factor of 15 at 130 km down to about 2 at 200 km and above. The predicted densities for aurora at 130 km without quenching are shown, to allow comparison with the predictions (scaled to the current N\(_2[\Lambda^3\Sigma_u^+, \nu' = 0]\) density) of Cartwright [1978] and Morrill and Benesch [1996]. The shape of the distribution differs from that of Cartwright due to the use of updated transition probabilities [Gilmore et al., 1992]. It is similar to that of Morrill and Benesch, who used the transition probabilities of Gilmore et al. However, the densities are about half of those calculated by Cartwright, with only about 10% of this discrepancy being due to the changes in the auroral spectrum and updated cross sections. The densities of Morrill and Benesch are also larger than the current values, but this is not an independent determination because they used excitation rates from Cartwright [1978].

[13] Relative distributions deduced in fitting synthetic spectra to observations of dayglow at 200 km [Broadfoot et al., 1997], auroral column emission [Degen, 1982], and aurora at 110 km [Eastes and Sharp, 1987] are also included in Figure 1 by scaling to the appropriate N\(_2[\Lambda^3\Sigma_u^+, \nu' = 0]\) value. There is good agreement in the shapes of the distributions for aurora, except that the current model does not predict the enhancement for \(\nu' = 6\) and \(\nu' = 7\) found by Degen. However, this enhancement is not seen in the measurements of Eastes and Sharp [1987] and also Degen states that for the vibrational populations, “The values adopted ... are still uncertain and somewhat arbitrary.” Hence that enhancement may not be real. For dayglow the measured values [Broadfoot et al., 1997] of the relative populations for levels 0, 1, and 2 agree with the present model. Broadfoot et al. also give theoretical predictions for levels 3–6 which are lower than the current predictions for levels 4–6. These discrepancies however are not accounted for by the processes (C'\Pi_u predissociation and resonance scattering) that are not in the current model but are included by Broadfoot et al. (The theoretical distributions calculated by Broadfoot et al. with these processes included are closer to our predictions than to their inferred measurements.)

[14] Given the discrepancies in both the absolute and relative values of the calculated \(\Lambda^3\Sigma_u^+\) populations with previous cases, the current model is tested indirectly by...
comparison of its predictions of other quantities with previous models and measurements. In Figure 2 calculated values for the production rate of “vibrational quanta” (the sum of the number of transitions to vibrational level $\nu''$ of the ground state, multiplied by $\nu''$), by direct electron impact and $A^3\Sigma_u^+ \rightarrow X^1\Sigma_g^+$ transitions, are compared with the results of Richards and Torr [1986]. The present calculation gives production rates of vibrational quanta by direct electron impact that are up to a factor of two different to those of Richards and Torr, being higher at lower altitudes and lower at higher altitudes. This is not unreasonable as the current model uses an updated set of electron impact cross sections [Campbell et al., 2004] for the vibrational levels. However, while the $A^3\Sigma_u^+$ contribution matches reasonably at higher altitudes, it is significantly lower at the lower altitudes. This is due to different assumptions about collisional quenching. Whereas we assume this returns excited $N_2$ to the ground state, Richards and Torr presumably use the assumption of Richards et al. [1986] that both radiation and atomic oxygen quenching leave $N_2$ in the same distribution of vibrational levels in the ground electronic state. When the current model is run without quenching, equivalent to the assumption of Richards and Torr that all deactivations of $A^3\Sigma_u^+$ end up in the same distribution of vibrational quanta, our production rates are in much closer agreement with theirs but larger by up to 50%. This difference is not unreasonable given that we have calculated the $A^3\Sigma_u^+ \rightarrow X^1\Sigma_g^+$ transitions in detail, whereas Richards and Torr presumably use the approximation [Richards et al., 1986] that all $A^3\Sigma_u^+ \rightarrow X^1\Sigma_g^+$ tran-
Figure 3. Adjusted measurements [Sharp, 1971] of the column emission rate for the downleg portion of a rocket flight into an IBC I+ aurora, for the sums of all A³Σ_u⁺ (ν’/ = 0) → X¹Σ_g⁺ (black filled circles) and of all A³Σ_u⁺ (ν’/ = 1) → X¹Σ_g⁺ emissions (black crosses), and the first negative (blue open circles). Predictions of the current model for the first negative (blue dash-dotted line), for ν’/ = 0 (black short dashed line), and for ν’/ = 1 with (black solid line) and without (black long dashed line) quenching and chemical reactions. Error bars at 120 km indicate the uncertainty on two example points that show the sum for only the subset of the transitions that were measured. All values for ν’/ = 0 are multiplied by 0.1 to avoid overlap on the plot.

...sitions terminate in X¹Σ_g⁺ (ν” = 7). The main point is that despite the current A³Σ_u⁺ populations being less than previously calculated [Cartwright, 1978], the unquenched populations are larger than calculated in another model, so there is not a consistent trend that they are underestimated.

[15] In the absence of direct measurements for the densities of the A³Σ_u⁺ state, the current model can be indirectly validated by comparison with measurements of A³Σ_u⁺ (ν’) → X¹Σ_g⁺ (ν’/) emissions. In Figure 3 such measurements are shown for the downleg of a rocket flight into an IBC I+ aurora [Sharp, 1971]. These measurements have a calculated component, in that emissions for only some transitions were measured and the total emission for a particular state. This suggests that our predicted densities of the A³Σ_u⁺ levels are accurate enough for an investigation of the production of NO via the A³Σ_u⁺ state.

[16] To scale the auroral flux independently of the A³Σ_u⁺ densities, it was set to give a good match of our predictions of the 427.8 nm first negative (N₂[A³Σ_u⁺(0) → X¹Σ_g⁺(1)]) to those measured by Sharp above 115 km. Our predictions of Sharp’s adjusted measurements are shown by the curves in Figure 3. There is good agreement of our predictions of the rates of A³Σ_u⁺(0) → X¹Σ_g⁺(ν’/) with the adjusted measurements of Sharp, with an underestimation for ν’/ = 1. The discrepancies are of the same order as the errors in the measured values, while the errors in the adjusted measurements might be larger due to errors in the transition probabilities used by Sharp in calculating them. Hence the agreement of the current predictions with the measurements is good in view of the experimental variation.

[17] As quenching and chemical reactions cause depletion of excited nitrogen without producing emission, the model was run with these omitted. This results in predicted emissions that are much higher than the measurements at low altitudes. Hence reasonable agreement with the measurements is seen only when the chemical reactions and quenching are included.

[18] While the current model is in disagreement with some previous results, principally in that it gives lower values for the A³Σ_u⁺ densities than earlier auroral calculations [Cartwright, 1978; Morrill and Benesch, 1996], it is in reasonable agreement with other calculations [Richards and Torr, 1986] for the production of vibrational quanta from A³Σ_u⁺ in daytime and in good agreement with measurements [Sharp, 1971] of auroral emissions produced by radiative decay of the A³Σ_u⁺ state. This suggests that our predicted densities of the A³Σ_u⁺ levels are accurate enough for an investigation of the production of NO via the A³Σ_u⁺ state.

[19] A problem is that in comparing with the available measurements only one parameter is generally available. A useful set of measurements would be simultaneous values of the N₂[A³Σ_u⁺] vibrational distribution, the 557.7 nm emission and the NO (2 → 1) infrared emission.

3.2. N₂[A³Σ_u⁺] and N₂[X¹Σ_g⁺(ν’/ > 11)] Production of NO

[20] As an initial test the current model is run for similar conditions to those for the model of Barth [1992]. In Figure 4 the production rates of NO by various species in the current model are compared with those of Barth. The current model provides a reasonable emulation of that of Barth and so is adequate to investigate the importance of excited N₂. Also included in Figure 4 are the rates of NO production due to the reactions of N₂[A³Σ_u⁺] and N₂[X¹Σ_g⁺ (ν’/ > 11)] with O, as predicted by our model. These reactions appear unimportant compared to the reactions of N atoms with O₂. However, including them actually increases the calculated NO density by a factor of about 2–3 at lower altitudes, as will be shown below.

[21] To investigate the role of N₂[A³Σ_u⁺] and N₂[X¹Σ_g⁺ (ν’/ > 11)] in the production of NO, the model is run with and without the reactions:

\[ N₂ [A³Σ_u⁺] + O \rightarrow NO + N(²D) \]

and

\[ N₂ [X¹Σ_g⁺(ν’/ > 11)] + O \rightarrow NO + N(⁴S), \]

to compare with measurements by the SNOE satellite [Barth et al., 2003] of the NO density.
In Figure 5 predicted values of NO density are compared with measurements for moderate and high solar activity at the equator at about 1100 LT, averaged over several intervals around the spring and autumn equinoxes. The time step calculations are run for 107 hours from 0600, using the atmospheric densities, temperatures and the $F_{10.7}$ index of 208.8 from the MSISE-90 model (http://modelweb.gsfc.nasa.gov/models/msis.html) for 20 September 2000, a day representative of those with high solar activity that is within one of the observation intervals. It can be seen that the predicted NO densities are tending to values in good agreement with the measurements for high solar activity at 110 km and 150 km but up to ~30% lower at altitudes in between. This is not as good as the agreement obtained by Barth and Bailey [2004], but the 30% discrepancy is almost within the sum of the uncertainties that they quote (18% for the measurements and 10% for solar input in the calculation, as shown by error bars on the plot) and within the range of discrepancies between their model and the measurements on other days. The agreement with the measurements is substantially better than given by the empirical model of Mitra [1968]. Thus while the model is by no means perfect, its ability to produce reasonable NO densities and a satisfactory shape for the profile with height enables us to use it to investigate the role of excited N2 in the production of NO.

The comparison of the time step predictions at 1100 LT on the fifth day of calculations with the measurements is replotted in Figure 6, along with the result of calculations which exclude one or both of the reactions of $N_2[A^3Σ_g^+] + O$ and $N_2[X^1Σ_g^+(v' > 11)] + O$. The predicted NO densities are significantly less when both reactions are excluded and are outside the range of uncertainty, particularly in the region of the peak density at about 107 km where they are less by a factor of 1.7. As the reductions are not significantly less when only the $N_2[A^3Σ_g^+] + O$ reaction is excluded, it is deduced that the $N_2[X^1Σ_g^+(v' > 11)] + O$ reaction is only a minor contributor to the NO density, except above 150 km where its contribution is similar to that of $N_2[A^3Σ_g^+] + O$.

Replacing the updated rate [Duff et al., 2005] for $N(^2D) + O_2$ with the earlier rate [Herron, 1999] (in the case where the excited N2 is excluded) reduces the NO density above 130 km by amounts similar to omitting the excited N2 reactions. Below 130 km the effect of the excited N2 reactions is greater, particularly at the height of the peak density and below, where it is 2–4 times as large.

The enhancement due to the inclusion of excited N2 seems out of proportion to the relative rates of production in Figure 4, where the contributions from $N_2[A^3Σ_g^+]$ and $N_2[X^1Σ_g^+(v' > 11)]$ are an order of magnitude less than the maximum for the nitrogen atom rates. This disproportionate effect may be explained by the fact that nitrogen atoms react with NO to destroy it as well as produce it. In contrast, for $N_2[A^3Σ_g^+]$ and NO the production mechanism is dominant because the reverse reactions [Herron, 1999]:

$$N_2[A^3Σ_g^+] + NO \rightarrow N_2 + NO[A^2Σ^+]$$

and

$$N_2[A^3Σ_g^+] + NO \rightarrow N_2 + N + O$$


Figure 4. Comparison of current calculations (dashed line of filled circles) with results of Barth [1992] (solid lines) for production rates of NO due to reactions involving species as labeled.

Figure 5. A comparison of the calculated NO density at 1100 LT on successive days of the simulation with the SNOE measurements for high (black solid line) and moderate (black dash-dotted line) solar activity and with an empirical model (dashed line of open diamonds) [Mitra, 1968]. Error bars at 122.5 km show the estimated measurement uncertainty and the altitude range for the high solar activity case. The error bar shows the estimated uncertainty due to solar input at 122.5 km to the calculated curve for 107 hours.
only become significant at NO densities much higher than calculated here. A second factor that makes the N2[A^3Σ_u^+] metastable state important in producing NO is that the excited N(2D) atoms produced by N2[A^3Σ_u^+] + O then take part in other reactions to produce NO, thus further enhancing the contribution by N2[A^3Σ_u^+]. This was confirmed by rerunning the model with the N(2D) product replaced by N(4S), which at altitudes below 120 km reduced the increase in the NO density by 77–95%.

Figure 7. A comparison of calculated NO densities with measurements by the SNOE satellite at 65°N for high (red line with solid circles) and maximum (blue vertical line) solar activity. The current calculations for 0.5 erg cm^{-2} s^{-1} (red line with solid circles) and 1.0 erg cm^{-2} s^{-1} (blue line with solid squares) are for high solar activity with auroral activities of 0.5 erg cm^{-2} s^{-1} and are repeated without the contributions from N2[A^3Σ_u^+] only (pluses) and without both N2[A^3Σ_u^+] and N2[X^1Σ_g^+] (red line with open circles). The current calculation, without contributions from N2[A^3Σ_u^+] and N2[X^1Σ_g^+] and with the other updated reaction rates, is applied (black line with open triangles) for the conditions of the calculation by Barth.
The results of the calculations above suggest the following important conclusions: with models that yield a reasonable estimate of the atmospheric NO density, the reaction of \( \text{N}_2[\text{A}^3\Sigma_u^+] \) with O atoms leads to a significant increase in the calculated NO density, up to a factor of 3 at lower altitudes for high geomagnetic conditions. The inclusion of the reaction of \( \text{N}_2[\text{X}^1\Sigma_g^-] (\nu' > 11) \) with O leads to a much smaller increase that only becomes apparent at higher altitudes.

### 3.3. Implication for 557.7 nm Emission

[29] The flux of 1 erg cm\(^{-2}\) s\(^{-1}\) with high solar activity of \( F_{10.7} = 200 \) and an auroral energy flux of 1 erg cm\(^{-2}\) s\(^{-1}\). The similarity to Barth’s model supports the validity of the current calculations.

[30] In addition to the production of NO, \( \text{N}_2[\text{A}^3\Sigma_u^+] \) can be deexcited by a reaction with O atoms, producing \( \text{O}(^1\Sigma_g^+) \), which can then radiatively decay to \( \text{O}(^1\Delta) \), emitting 557.7 nm radiation [O’Neil et al., 1979]. As the reaction of \( \text{N}_2[\text{A}^3\Sigma_u^+] \) to produce NO competes with this process, an important check on our model is that the 557.7 nm emissions are consistent with the assumed atmospheric conditions. In Figure 8a we compare our predictions with measurements by Sharp et al. [1979] of 557.7 nm emissions. An auroral flux of 1.04 erg cm\(^{-2}\) s\(^{-1}\) scaled the electron spectrum in the calculation to direct measurements at 143 km by Sharp et al. This leads to predictions of \( \text{O}(^1\Sigma_g^+) \) excitation and 557.7 nm emission at 143 km that are close to those of Sharp et al., as shown in Figure 8a. The calculated values of \( \text{O}(^1\Sigma_g^+) \) excitation and 557.7 nm emission agree well with the values of Sharp et al. at 130–150 km, while overestimating by a factor of up to 2 at higher altitudes and considerably underestimating the 557.7 nm emission at lower altitudes. Note that the agreement for \( \text{O}(^1\Sigma_g^+) \) excitation below 120 km is expected because the values of Sharp et al. for this quantity were used to determine the height dependence (equation (1)) of the energy flux of our auroral spectrum in the altitude range 100–120 km.

[31] Figure 8b shows the result of running the calculation for an auroral flux of 2.54 erg cm\(^{-2}\) s\(^{-1}\), which is the value quoted by Sharp et al. [1979] for a satellite measurement that was coordinated with the rocket measurement. In this case the predictions are in agreement with the measurements below 120 km.

[32] In Figure 8c the 557.7 nm column emission rates calculated for the two auroral energy fluxes (2.54 and 1.04 erg cm\(^{-2}\) s\(^{-1}\)) are compared with the values measured by Sharp et al. [1979]. The current calculation using the energy flux of 2.54 erg cm\(^{-2}\) s\(^{-1}\) agrees well at the lowest altitude, while above 120 km there is good agreement using the energy flux of 1.04 erg cm\(^{-2}\) s\(^{-1}\).

[33] Thus we are able to predict the correct magnitude of the 557.7 nm emission measured by Sharp et al. [1979] at lower altitudes by using the primary electron flux (measured from a satellite) or at higher altitudes by scaling to the secondary electron spectrum measured by a rocket at 143 km. The apparent discrepancy we find in these two cases may be due to auroral variability, coupled with different locations for different observations [Rees et al., 1977]. This may also explain the discrepancy between the shape of the calculated and measured profiles in the height range 100–120 km, as this is the range over which the auroral energy flux seems to change. However, it could also be due, at least in part, to our addition of the new pathway \( \text{N}_2[\text{A}^3\Sigma_u^+] + \text{O} \rightarrow \text{NO} + \text{N}(^2\Delta) \), which would reduce the 557.7 nm emission.

[34] In Figure 8b the result of running the model without the NO production pathway is shown for the energy flux of 2.54 erg cm\(^{-2}\) s\(^{-1}\). It indicates that the 557.7 nm emission is...
predicted to be up to 30% higher, although there is little difference in the shape of the altitude profile. Thus the effect of including the NO production pathway is smaller than the discrepancies between the model and measurements, although the results when it is included are closer to the measurements for both the definitive values at 143 km and the column emission rate at 100 km.

3.4. Effects of Atmospheric Winds

Significant vertical winds due to auroral heating have been measured [Price et al., 1995], so it is necessary to consider whether the predicted density gradients of NO could exist in the presence of wind. Information on the vertical winds over the time interval and latitude band of the SNOE measurements is not available, so here we calculate the effect of a postulated wind profile based on isolated measurements. Wind is included in the model by calculating, in each time step, the change in density in each height interval due to the amount of NO that is added or removed at the top and bottom of the height interval by the wind. Measurements of vertical wind, deduced from auroral emissions, at 120 km over one night [Ishii, 2005] are used. It is assumed that the wind pattern is repeated on subsequent nights, and linear interpolation is used over the daylight interval and where the measurement error is large, as shown in Figure 9. The wind speed at 240 km and above is set to 3 times the speed at 120 km, based on the ratios for simultaneous measurements at different heights [Price et al., 1995], with linear interpolation of the speeds between 120 km and 240 km. Below 120 km, assuming conservation of mass, the speed is scaled as the reciprocal of the atmospheric density. These speeds are divided by 20, as we are considering an auroral flux of 0.5 erg cm$^{-2}$ s$^{-1}$, whereas the measurements are presumed to have been made during an intense aurora. A comparison of the results with these vertical winds with a “no wind” case from Figure 7 is shown in Figure 10. The wind affected case is labeled “downward” as this is the direction prior to the 1100 LT NO measurements. The “upward” case is calculated by running the model with the winds reversed. The postulated vertical winds produce changes of the same order of magnitude as the discrepancies between the calculations and measurements at some altitudes, and also tend to reduce the NO density at 115 km, irrespective of the wind direction. However, the large gradient in NO density remains despite the effect of vertical winds.

It has been suggested that horizontal winds could reduce the NO density by transporting NO out of the auroral zone. Barth et al. [2003] postulated a meridional wind of 30 ms$^{-1}$ to account for an observed increase in NO at 106 km at midlatitudes following a large geomagnetic storm. This was questioned by Richards [2004], who suggested that it is unlikely that significant transport of NO from high latitudes occurs at this altitude and that other mechanisms could account for the production of the mid-

![Figure 9](image1.png)

Figure 9. Measurements of vertical neutral wind (positive upward) [Ishii, 2005] with low (black dashed line) and high (blue dashed line) error and the fit (red solid line) to these measurements used in this work. The time of the NO measurements (1100 LT) is indicated.

![Figure 10](image2.png)

Figure 10. A comparison of calculated NO densities with measurements by the SNOE satellite of average densities at 65$^\circ$N for high geomagnetic activity (red solid line). The calculations are for high solar activity with auroral activity of 0.5 erg cm$^{-2}$ s$^{-1}$, without vertical wind (red line with solid circles) and with winds that are upward (blue line with open circles) and downward (blue line with open triangles) before the measurement time.
latitude NO. However, even if this transport occurs, the measurements [Barth et al., 2003, Figure 11] show the postulated transport of NO out of the auroral zone does not occur for most of the time and that when it does the NO density in the auroral region is above average. Thus the one-dimensional calculations here are appropriate for the time-averaged measurements presented by Barth et al. It is expected that horizontal winds would smooth out horizontal variations in NO density created by patchy auroral activity and associated vertical winds. This is consistent with the absence in the measurements of the wind-induced reduction at around 115 km predicted when vertical winds are included in the calculations.

4. Conclusions

An atmospheric model which combines the radiative cascade through the excited states of molecular nitrogen with chemical reactions involving excited nitrogen is used to predict the absolute densities of the vibrational levels of N2[A3Σu+g] and N2[X1Σg−] and associated emissions. These predictions are compared with various previous simulations and measurements in order to assess the validity of the model, which is then used to investigate the role of vibrationally and electronically excited N2 in the production of nitric oxide in the atmosphere.

In general, predictions of the N2[A3Σu+g] densities and the associated production of vibrational quanta are consistent with previous simulations, although in some cases discrepancies of a factor of 3 were noted. However, predictions of emissions due to radiative decay of the N2[A3Σu+g] state compare well with measurements in an auroral environment, which supports the overall accuracy of the model employed in this study.

The model was used to predict the NO density in the atmosphere for cases where measurements have been made by satellite: high solar activity at the equator and high geomagnetic activity at high latitudes. At the equator there are discrepancies between the predictions and measurements of ~30% at some altitudes, but there is much better agreement (to ~10%) at the altitude of the peak density. In the auroral case there are discrepancies of up to a factor of two at some altitudes which may be due to the characteristic energy of the auroral model being high. Using two different auroral energy fluxes within a “standard” range, the maximum predicted NO density is shown to be similar to either the average for high geomagnetic activity or to the maximum value observed.

Emulation of a previous calculation showed good agreement for calculated NO densities and suggests that the model employed in this study is sufficient to investigate the role of excited N2 in the production of NO. This was achieved by running the new model but leaving out either of the production mechanisms involving N2[A3Σu+g] and N2[X1Σg−]. Omitting the N2[X1Σg−] pathway caused a small reduction above about 150 km and had a negligible effect at lower altitudes. Omitting the N2[A3Σu+g] production mechanism reduced the NO density in the region of the peak by a factor of ~1.7 in the equatorial case and by 2–3 in the auroral case. At altitudes below 130 km the increase in NO density due to excited N2 is greater than the increase produced by changing to an updated reaction rate for N(2D) + O2.

Inclusion of the NO production pathway will reduce the predicted 557.7-nm emission, so predictions of the 557.7 nm emission were made, with and without the NO production pathway, and compared with measurements. Inclusion of NO production from N2[A3Σu+g] improved the predictions of 557.7 nm emission, but the improvements were small in comparison to the discrepancies between the calculations and measurements. Inclusion of a vertical wind profile changed the NO density by the same order as the discrepancies between calculations and measurements but did not reduce the large NO density gradients calculated. Thus the effect on the calculated NO density of the N2[A3Σu+g] + O reaction is predicted to be large even in the presence of vertical winds and this conclusion is not invalidated by considering its effect on the 557.7 nm emission.

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References


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