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### Thermospheric nitric oxide from the ATLAS 1 and Spacelab 1 missions

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Abstract. Spectral and spatial images obtained with the Imaging Spectrometric Observatory on the ATLAS 1 and Spacelab 1 missions are used to study the ultraviolet emissions of nitric oxide in the thermosphere. By synthetically fitting the measured NO gamma bands, intensities are derived as a function of altitude and latitude. We find that the NO concentrations inferred from the ATLAS 1 measurements are higher than predicted by our thermospheric airglow model and tend to lie to the high side of a number of earlier measurements. By comparison with synthetic spectral fits, the shape of the NO  $\gamma$  bands is used to derive temperature as a function of altitude. Using the simultaneous spectral and spatial imaging capability of the instrument, we present the first simultaneously acquired altitude images of NO  $\gamma$  band temperature and intensity in the thermosphere. The lower thermospheric temperature images show structure as a function of altitude. The spatial imaging technique appears to be a viable means of obtaining temperatures in the middle and lower thermosphere, provided that good information is also obtained at the higher altitudes, as the contribution of the overlying, hotter NO is nonnegligible. By fitting both self-absorbed and nonabsorbed bands of the NO gamma system, we show that the self absorption effects are observable up to 200 km, although small above 150 km. The spectral resolution of the instrument (1.6 Å) allows separation of the  $N^+({}^5S)$  doublet, and we show the contribution of this feature to the combination of the NO  $\gamma$  (1,0) band and the N<sup>+</sup>(<sup>5</sup>S) doublet as a function of altitude (less than 10%) below 200 km). Spectral images including the NO  $\delta$  bands support previous findings that the fluorescence efficiency is much higher than that determined from laboratory measurements. The Spacelab 1 data indicate the presence of a significant population of hot NO in the vehicle environment of that early shuttle mission.

### 1. Introduction

An instrument flown on the ATLAS 1 shuttle mission [Torr, 1993] from March 24 to April 2, 1992, obtained spectral images of the nitric oxide gamma bands over latitudes ranging from the equator to approximately 65°N. The altitudes sampled ranged from 80 to 300 km. The Imaging Spectrometric Observatory (ISO) database provides additional information on this important upper atmospheric constituent. Specifically, the complete NO gamma band system is measured at relatively high spectral resolution, allowing the NO concentrations to be inferred from bands that are not subject to self-absorption and permitting spectral separation from spectrally blended features. The range of latitudes covered exceeds that previously available for the spectral features studied here. In addition to the determination of NO concentrations, the spectra are used to examine the feasibility of using these bands to derive thermospheric temperature profiles. In addition, the twodimensional imaging capability of the instrument allows altitude to be imaged in one dimension, while wavelength is imaged in the other (with all detector pixels being simultaneously exposed), so that in the lower thermosphere/upper

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Paper number 95JA01040. 0148-0227/95/95JA-01040\$05.00 mesosphere a temperature/concentration altitude image is obtained in each exposure in addition to the spectral image. Both the NO concentrations and temperatures are compared with models. The ATLAS 1 data are compared with similar data obtained during an earlier flight of the instrument on the Spacelab 1 mission in late 1983, which obtained the first simultaneously acquired altitude images of temperature and concentration in the lower thermosphere/upper mesosphere.

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The existence of significant NO concentrations in the upper atmosphere was first postulated by Nicolet [1945] in order to explain the measured D region ionization. He pointed out that NO is readily ionized by solar Lyman  $\alpha$  radiation. The first measurements of NO emissions were made almost 20 years later from a rocket [Barth, 1964] in which the NO  $\gamma$  (1,0) band at 2150 Å was observed with a spectrometer. Further measurements of this feature from rockets and satellites established the photochemistry of NO [Thomas, 1978; Cravens et al., 1979; McCoy, 1983].

NO also plays an important role in the energy budget of the lower thermosphere. Torr et al. [1980b] showed that between 120 and 200 km, the exothermic chemical reaction between NO and  $N(^4S)$  constitutes a major heat source. Kockarts [1980] demonstrated that the 5.3-µm radiation from vibrationally excited NO is an effective means of cooling the lower thermosphere. This was quantified in relation to other heating

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Table 1.	Relevant Parame	eters for Far-	and Near-UV
Imaging S	pectrometric Obs	ervatory Spec	ctrometers

Spectrometer	Wavelength Range, Å	Simultaneous Spectral Image Range, Å	Spectral Resolution, Å
3	2138-4282	236	2.3
4	1150-2400	177	1.6

and cooling rates by *Richards et al.* [1982] who showed that at altitudes below 150 km, the  $5.3-\mu$ m radiation is the dominant cooling process for the lower thermosphere.

Other studies have characterized the morphological behavior of NO. Work based on the 2150 Å (1,0) NO gamma band measured from the Atmosphere Explorer satellites focused on the diurnal, altitude, and latitudinal morphology [*Cravens and Stewart*, 1978; *Stewart and Cravens*, 1978]. These studies found the variability to be largest at high latitudes.

Data from the Solar Mesosphere Explorer satellite [Barth et al., 1988] were used to map the low-latitude variations in NO concentrations at 110 km, indicating a solar maximum to minimum ratio of 7.5. Another study using the Nimbus 7 solar backscattered ultraviolet (SBUV) data [McPeters, 1989], found that the equatorial NO column concentrations above 48 km

increased by a factor of 2 from 1980 to 1986, with smaller changes at higher latitudes.

At lower altitudes (mesosphere and stratosphere), NO is involved in the catalytic removal of ozone. There is a possible link to the NO produced at higher altitudes at high latitudes as the latter might be transported to lower altitudes during polar winter conditions [Solomon et al., 1982; Solomon and Garcia, 1984].

Much of what we know about NO in the thermosphere is based on measurements of the very bright (1,0) y band at 2150 Å. In the early 1980's there was some concern that these measurements might be compromised by the blended  $N^+({}^{5}S)$ doublet at 2139/2149 Å [see Victor and Dalgarno, 1982; Torr and Torr, 1985al. The N<sup>+</sup> feature was found to have a substantial effect on the higher-altitude observations and the contribution to the lower thermospheric measurements was relatively small. Cravens [1977] pointed out that the NO concentrations inferred from measurements of the 2150 Å band might be compromised in the lower thermosphere because of the fact that the emission might not be optically thin as had been assumed in earlier studies. The (1,0) transition producing the 2150 Å emission ends in the v' = 0 vibrational level, where the bulk of the NO molecules reside. A study of Spacelab 1 spectral/spatial images at 100 km by Erwin [1985] found that when the NO  $\gamma$  spectra were synthetically fitted,



Figure 1. Illustration of the spectral/altitude imaging capability of Imaging Spectrometric Observatory.



**Figure 2.** Ground track of tangent ray point during the various sequences used in this study: (dotted-asterisk line) altitude imaging mode with center of field of view at approximately 140 km; (dashed-crossed line) vehicle rolls to sweep field of view down through the atmosphere; (dashed-opened circle line) altitude imaging mode with center of field of view directed at 99 km.

with good fits to the (1,0) and (2,0) bands, that the (1,1) and (2,2) bands were systematically underfitted by the model. Had Erwin fitted the (1,1) and (2,2) bands he would, in fact, have measured the degree of self-absorption of the (1,0) and (2,0) bands. *Cleary* [1986] derived altitude profiles of NO concentration from the  $\gamma(1,0)$  and (2,0) bands. These profiles showed a roll over below 120 km, which the authors presumed to be due to self-absorption. More recently, *Eparvier and Barth* [1992] reported the results of a high-latitude rocket experiment from which they were able to clearly demonstrate the self-absorption of the 1-0 band below 130 km by comparison with the 0-1 band, as the latter does not end in the 0 vibrational level.

#### 2. Observational Sequences

The Imaging Spectrometric Observatory (ISO), flown on the ATLAS 1 mission and the earlier Spacelab 1 mission, comprised five imaging spectrometers. Each spectrometer covered a portion of the wavelength range from the extreme ultraviolet to the near-infrared [*Torr et al.*, 1982, 1983]. The instrument was designed to measure emissions from the thermosphere and mesosphere under sunlit or dark conditions. Two of these spectrometers measured wavelength ranges that contained the various UV bands of NO. The parameters associated with these two spectrometers are summarized in Table 1.

Where the term altitude/spectral image is used, this means a

Table	2. (	Geophysica	l Parameters	for S	equences	Studied
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Parameter	March 29, 1992	March 25, 1992	March 26, 1992
Shuttle attitude	fixed tangent ray height (~140 km)	fixed tangent ray height (~99 km)	altitude scan via roll maneuver (300–105 km)
Universal time	0930-0950	0845-(0902)-0915	07:18-07:43
Local time	0922-1612	0545-(0852)-1438	05:22-10:53
Solar zenith Angle	66.6-63.9	94-(58)-62	99.6-55.5
Latitude	64.2°N-25.2°N	1°N(50°Ń)46°N	10.5°S-57.0°N
Longitude	1.9°W-95.6°E	37°W(10°É)92°E	27.9°W-60.8°E
$F_{10,7}$ -cm flux	193	<b>`186</b> ´	179
$F_{10,7}$ 90-day average	182	189	187
Ap	14	18	13
Kp (average)	3-	3-	2+

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 Table 3.
 Summary of Photochemistry Used in the Airglow Model

Reaction Number	Reaction	Rate Coefficient ( $cm^3 s^{-1}$ ) or Rate ( $s^{-1}$ )	Reference
(1) (2)	$O^+ + e \to O + h\nu$ $O_2 + e \to O + O$	$\sim 4 \times 10^{-12} (T/300)^{0.7}$ 1.6 × 10 <sup>-7</sup> (300/T <sub>e</sub> ) <sup>0.55</sup> for T <sub>e</sub> $\geq$ 1200 K	Torr [1985] Torr and Torr [1981]
(3) (4)	$O^+ + O_2 \rightarrow O_2^+ + O$ $O^+ + N_2 \rightarrow NO^+ + N$	$2 \times 10^{-10} (300/T_e)^{-1.7} \text{ for } T_e < 1200 \text{ K}$ $2.1 \times 10^{-11} \{T_n + 2\text{Ti}/300\}^{-0.763}$ $1.533 \times 10^{-12} - 5.92 \times 10^{-13} (T_{\text{eff}}/300) + 8.60 \times 10^{-14} (T_{\text{eff}}/300)^2 \text{ for } 300 \le T_{\text{eff}} \le 1700 \text{ K}$	Mehr and Bional [1969] Chen et al. [1978] <sup>a</sup> St. Maurice and Torr [1978]; Albritton [1978]; Chen et al. [1978]
(5) (6)	$NO^{+} + e \rightarrow N + O$ $N_{2}^{+} + O \rightarrow NO^{+} + N$	$2.73 \times 10^{-12} - 1.155 \times 10^{-12} (T_{eff}/300) + 1.483 \times 10^{-13} (T_{eff}/300)^2 \text{ for } 1700 < T_{eff} < 6000 \text{ K}  4.3 \times 10^{-7} (T_e/300)^{-1}  1.4 \times 10^{-10} (Ti/300)^{-0.44} \text{ for } Ti < 1500$	Torr and Torr [1979] McFarland et al. [1974]; Torr
(7) (8) (9) (10) (11) (12) (13) (14) (15)	$N_{2}^{+} + e \rightarrow N + N$ $N_{2}^{+} + 0 \rightarrow O^{+} + N_{2}$ $N_{2}^{+} + O_{2} \rightarrow O_{2}^{+} + N_{2}$ $N^{+} + O_{2} \rightarrow O_{2}^{+} + N$ $N^{+} + O_{2} \rightarrow NO^{+} + O$ $N^{+} + O_{2} \rightarrow NO^{+} + O(^{1}D)$ $O_{2}^{+} + N \rightarrow NO^{+} + O$ $O_{2}^{+} + NO \rightarrow NO^{+} + O_{2}$ $O^{+}(^{2}D) + N_{2} \rightarrow N_{2}^{+} + O$	K $2.7 \times 10^{-7}$ $0.07 k_6 (T_i/300)^{0.21}$ $9.1 \times 10^{-11} \exp(-0.002T_{eff})$ $4 \times 10^{-10}$ $2 \times 10^{-10}$ $\beta = 0.7$ $1.2 \times 10^{-10}$ $4.4 \times 10^{-10}$ $8 \times 10^{-10}$	[1979] Abdou et al. [1984] McFarland et al. [1974] Lindinger et al. [1974] Huntress and Anicich [1976] Huntress and Anicich [1976] Langford et al. [1985] Fehsenfeld [1977] Lindinger et al. [1974] Rowe et al. [1980]; Johnsen and Biondi [1980]
(16) (17) (18) (19) (20) (21) (22) (23) (24) (25) (25) (26) (27) (28)	$O^{+}(^{2}D) + O \rightarrow O^{+}(^{4}S) + O$ $O^{+}(^{2}D) + O_{2} \rightarrow O_{2}^{+} + O$ $O^{+}(^{2}D) + e \rightarrow O^{+}(^{4}S) + e$ $O^{+}(^{2}P) \rightarrow O^{+}(^{2}D) + h\nu$ $O^{+}(^{2}P) \rightarrow O^{+}(^{4}S) + h\nu$ $O^{+}(^{2}P) + e \rightarrow O^{+}(^{4}S) + e$ $O^{+}(^{2}P) + e \rightarrow O^{+}(^{4}S) + e$ $O^{+}(^{2}P) + Q \rightarrow \text{products}$ $O^{+}(^{2}P) + O \rightarrow \text{products}$ $He + h\nu \rightarrow He^{+} + e$ $He^{+} + N_{2} \rightarrow N^{+} + N + He$ $He + + N_{2} \rightarrow N^{+}_{2} + He$ $O^{+} + H \rightarrow H^{+} + O$	$\begin{array}{l} 5\times10^{-12}\\ 7\times10^{-10}\\ 6.6\times10^{-8}(300/T_e)^{0.5}\\ 0.173\ \mathrm{s}^{-1}\\ 0.047\ \mathrm{s}^{-1}\\ 1.5\times10^{-7}(300/T_e)^{0.5}\\ 4.7\times10^{-8}(300/T_e)^{0.5}\\ 3.4\times10^{-10}\\ 4.0\times10^{-10}\\ 4.0\times10^{-8}-1.2\times10^{-7}\mathrm{c}\\ 1\times10^{-9}\\ 6.5\times10^{-10}\\ 2.2\times10^{-11}\ (T_i)^{0.5} \end{array}$	Abdou et al. [1984] Johnsen and Biondi [1980] Henry et al. [1969] Seaton and Osterbrock [1957] Seaton and Osterbrock [1957] Henry et al. [1969] Chang et al. [1993] Chang et al. [1993] Torr and Torr [1985] Adams and Smith [1976] Adams and Smith [1976] derived from Banks and Kockarts
(29)	$H^+ + O \rightarrow O^+ + H$	$2.5 \times 10^{-11} (T_n)^{0.5}$	[1973] derived from Banks and Kockarts
<ul> <li>(30)</li> <li>(31)</li> <li>(32)</li> <li>(33)</li> <li>(34)</li> <li>(35)</li> <li>(36)</li> <li>(37)</li> <li>(38)</li> <li>(39)</li> <li>(40)</li> <li>(41)</li> <li>(42)</li> <li>(43)</li> <li>(44)</li> <li>(45)</li> <li>(46)</li> </ul>	$ \begin{array}{l} {\rm O}^{+} + {\rm N}(^{2}D) \rightarrow {\rm N}^{+} + {\rm O} \\ {\rm NO}^{+} + e \rightarrow {\rm N}(^{2}D) + {\rm O} \\ {\rm N}_{2}^{+} + e \rightarrow {\rm N}(^{2}D) + {\rm N} \\ {\rm N}_{2}^{+} + 0 \rightarrow {\rm N}(^{2}D) + {\rm NO}^{+} \\ {\rm N}^{+} + {\rm O}_{2} \rightarrow {\rm N}(^{2}D) + {\rm O}_{2}^{+} \\ {\rm N}(^{2}D) + {\rm O} \rightarrow {\rm N}(^{4}S) + {\rm O} \\ {\rm N}(^{2}D) + {\rm O}_{2} \rightarrow {\rm NO} + {\rm O} \\ {\rm N}(^{2}D) + {\rm O}_{2} \rightarrow {\rm NO} + {\rm O} \\ {\rm N}(^{2}D) + {\rm O}_{2} \rightarrow {\rm NO}^{+} + {\rm O} \\ {\rm N}(^{2}D) + {\rm O}_{2} \rightarrow {\rm NO} + {\rm O} \\ {\rm N}(^{4}S) + {\rm O}_{2} \rightarrow {\rm NO} + {\rm O} \\ {\rm N}(^{4}S) + {\rm O}_{2} \rightarrow {\rm NO} + {\rm O} \\ {\rm O}_{2} + h\nu \rightarrow {\rm O}(^{1}D) + {\rm O} \\ {\rm O}_{2} + h\nu \rightarrow {\rm O}(^{1}D) + {\rm O} \\ {\rm O}_{2} + e \rightarrow {\rm O}(^{1}D) + {\rm O} \\ {\rm O}(^{1}D) + {\rm N}_{2} \rightarrow {\rm O}(^{3}P) + {\rm N}_{2} \\ {\rm O}(^{1}D) \rightarrow {\rm O}(^{3}P) + h\nu \\ {\rm N}(^{2}D) + {\rm O}_{2} \rightarrow {\rm O}(^{1}D) + {\rm NO} \end{array} $	5 × 10 <sup>-11</sup> $\beta k_5$ , where $\beta = 0.76$ $\beta k_7$ , where $\beta = 1.9$ $\beta k_6$ , where $\beta = 1.0$ $\beta k_{10}$ , where $\beta = 1.0$ $\sim 7 \times 10^{-13}$ $6 \times 10^{-12}$ $5 \times 10^{-10} (T_e/300)^{0.5}$ $1 \times 10^{-10}$ $4.4 \times 10^{-12} \exp(-3220/T)$ $3.4 \times 10^{-11}$ $\beta = 1; J_{x}(O2)_{SR} = (1.5-2.8) \times 10^{-6}$ $\beta k_2$ , where $\beta = 1.2$ $2.0 \times 10^{-11} \exp(107.8/T_n)$ $2.9 \times 10^{-11} \exp(67.5/T_n)$ 0.00934 $\sim 5 \times 10^{-12}$	[1973] Torr et al. [1979] Kley et al. [1977] Queffelec et al. [1985] Frederick and Rusch [1977] assumed Richards et al. [1981] Lin and Kauffman [1971] Frederick and Rusch [1977] Dalgarno [1970] Becker et al. [1976] Lee et al. [1978] Torr et al. [1980] Abreu et al. [1976] Streit et al. [1976] Fischer and Saha [1983] Rusch et al. [1978]; Torr et al. [1981]
(47)	$O_2^+ + e \rightarrow O(^1S) + O$	$\beta k_2$ , where $\beta = 0.08$	Bates and Zipf [1981]; Abreu et al. [1986]
(48) (49) (50) (51) (52) (53) (54)	$O_{2}^{+} + N \rightarrow O(^{1}S) + NO^{+}$ $O(^{1}S) + O(^{3}P) \rightarrow O + O$ $O(^{1}S) + O \rightarrow O + O$ $O(^{1}S) \rightarrow O(^{1}D) + h\nu$ $\rightarrow O(^{3}P) + h\nu$ $N(^{2}D) + NO \rightarrow N_{2} + O$ $O(^{1}S) + O_{2} \rightarrow O(^{3}P) + O_{2}$ $N_{2}(\mathcal{A}^{3}\Sigma_{+}^{+}) + O \rightarrow \text{products}$ $N_{2}(\mathcal{A}^{3}\Sigma_{+}^{+}) + O \rightarrow \text{products}$	$\sim 2 \times 10^{-11}$ $2 \times 10^{-14}$ $8 \times 10^{-12}$ 1.07 0.045 $7 \times 10^{-11}$ 4.9 × 10^{-12} exp (-1730/RT) $2 \times 10^{-11}$	Frederick et al. [1976] Slanger and Black [1981] Abreu et al. [1986] Kernahan and Pang [1975] Lin and Kauffman [1971] Zipf [1985] Piper [1982]
(55) (56) (57)	$N_2(A^{3}\Sigma_{u}^{+}) + O \rightarrow O(^{1}S) + N_2$ $N_2^{+} + O \rightarrow O^{+} + N_2$ $N_2 + h\nu \rightarrow N^{+} + N + e$ $N_2 + h\nu \rightarrow N_2^{+} + e$	$\beta k_{54}$ , where $\beta = 0.37$ ~2 × 10 <sup>-12 b</sup> (1.78-5.14) × 10 <sup>-18</sup> (3.06-8.82) × 10 <sup>-7</sup>	Piper [1982] Abdou et al. [1984]; Torr [1985] Torr and Torr [1985] <sup>c</sup>

#### Table 3. (continued)

Table 4.

Reaction Number	Reaction	Rate Coefficient ( $cm^3 s^{-1}$ ) or Rate ( $s^{-1}$ )	Reference
(58)	$O + h\nu \rightarrow O^{+}({}^{4}S) + e$ $O + h\nu \rightarrow O^{+}({}^{2}D) + e$ $O + h\nu \rightarrow O^{+}({}^{2}P) + e$ $O + h\nu \rightarrow O^{+}({}^{4}S) + e$ $O + h\nu \rightarrow O^{+}({}^{2}P^{e}) + e$	$\begin{array}{c} (0.98-2.81) \times 10^{-7} \\ (0.79-2.34) \times 10^{-7} \\ (0.45-1.38) \times 10^{-7} \\ (1.04-3.43) \times 10^{-8} \\ (0.46-1.42) \times 10^{-8} \end{array}$	Torr and Torr [1985]°
(59) (60)	$O({}^{1}D) + O \rightarrow O({}^{3}P) + O$ $O + e_{\text{therm}} \rightarrow O({}^{1}D)$	$\hat{8} \times 10^{-12'}$ $k_{60} = 1.1 \times 10^{-10} \sqrt{T_e} e^{-2.27 \times 10^4/T_e}$ $[0.406 + 3.57 \times 10^{-5} T_e - (0.333 + 1.83 \times 10^{-5}T_e)e^{-1.37 \times 10^4/T_e} - (0.456 + 1.74 \times 10^{-5}T_e)e^{-2.97 \times 10^4/T_e}]$	Abreu et al. [1986] Rees et al. [1967]

<sup>a</sup>Since the results of Chen et al. [1978] stop at 700 K, we normalize the converted drift tube data parameterized by St. Maurice et al. [1978] at this temperature.

<sup>b</sup>The model computes this rate coefficient as a function of  $N_2^+$  vibrational temperature (Tv). The value reduces to that given by (6) when Tv =Tn. Inclusion of this process is an option available. The ranges given for the ionization frequencies indicate the variation over a solar cycle.

Species Reference Type

Summary of Photoionization and Photoelectron Excitation Cross Sections

Photoionization	0	Samson and Pareek [1985]
	$O_2, N_2$	Kirby et al. [1979]
Total photoabsorption	$O_2, N_2$	Torr et al. [1979]
Total electron impact ionization cross sections	$O_2, N_2$	see Richards and Torr [1990]
Total electron impact excitation cross sections	0	Zipf and Erdman [1985]; Vaughr and Doering [1986]
	$N_2$	Cartwright [1977]
	O(D), O(S)	Henry et al. [1969]



Figure 3. Simplified potential diagram for NO. (Modified from Gilmore [1986].)

Table :	5.	Photochemical	Sources	and	Sinks	of	NC	)
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	Reaction	Rate Coefficient (cm <sup>3</sup> s <sup>-1</sup> ) or Rate (s <sup>-1</sup> )	Reference
		Sources	
1	$N(^{2}D) + O_{2} \rightarrow NO + O$	$6 \times 10^{-12}$	Lin and Kaufman [1971]
2	$N(^{4}S) + O_{2} \rightarrow NO + O$	$4.4 \times 10^{-12} \exp(-3220/T)$	Becker [1969] and Groth et al. [1969]
3	$N_2 + O^+(^2P) \rightarrow N^+ + NO$	$1.0 \times 10^{-10}$	Flesch and Ng [1991]
4	$N(^{2}P) + O_{2} \rightarrow$	$2.2 \times 10^{-12}$	Piper [1993]
5	$\operatorname{NO}(X^2\Pi) + h\nu \to \operatorname{NO}(A^2\Sigma)$	$7.7 \times 10^{-6}$ photons molecule s <sup>-1</sup> for the (1,0) band	Barth [1966]
		Sinks	
6	$N(^{4}S) + NO \rightarrow N_{2} + O$	$3.4 \times 10^{-11}$	Lee [1978] and Michael et al. [1978]
7	$N(^{2}D) + NO \rightarrow N_{2} + O$	$7 \times 10^{-11}$	Lin and Kaufman [1971]
8	$O_2^+$ + NO $\rightarrow$ NO <sup>+</sup> + $O_2$	$4.4 \times 10^{-10}$	Lindinger [1974] and Fehsenfeld et al. [1974]
9	$NO^{+} + h\nu \rightarrow NO^{+} + e^{-\nu}$	$7.3 \times 10^{-7}$	Rusch [1973]
10	$NO + h\nu \rightarrow O + N(^4S)$	$5.7 \times 10^{-6}$	Nicolet and Cieslik [1980]

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Figure 4. Photochemical (a) sources and (b) sinks of NO  $A^2\Pi$  for a field line at 61°N, 46°E on March 26, 1992, at 0730 UT.

simultaneously acquired two-dimensional image with spectral information in one dimension and spatial information in the other, without scanning a grating or mirror to build up this image in a time series.

The spectrometer array was mounted in the payload bay of the shuttle and made observations in a variety of vehicle attitudes. Observations made with the vehicle in three different attitudes are used in this study. In the first of these (Figure 1), the nose of the vehicle was pitched down so that the field of view was directed at a tangent ray height of approximately 140 km and held at close to this altitude while six spectral scans were made of the full wavelength range of the instrument. These spectra are used here to illustrate the dayglow spectrum in the lower thermosphere. In the second attitude selected for this study, the vehicle was rolled about the nose-tail axis so that the field of view of the instrument was scanned down through the atmosphere while the shuttle moved along its orbital path. These data are compared with other observations and a photochemical model of the thermospheric nitric oxide. The third vehicle attitude was similar to the first, with the nose of the vehicle pitched down so that the field of view was directed into the lower thermosphere at approximately 99 km and held there while the shuttle moved along its orbit. In this configuration (also that illustrated in Figure 1) the grating was cycled between six of the grating steps to emphasize the major spectral features at higher time resolution than if a complete spectral scan had been done. These data will be used to illustrate the imaging capability. The geographical location of the tangent ray point in these various observation sequences is shown in Figure 2.

The geophysical parameters for the sequences used in this study are shown in Table 2.

In the case of the March 25, 1992, column shown in Table 2, the numbers in parentheses are for the particular location of the data segment shown in this paper.

#### 3. Airglow Model

The data from the roll sequence are compared with the column concentration of NO along the same line of sight computed from our interhemispherically coupled airglow model. This model, which has been described elsewhere [Torr et al., 1990], solves the time dependent continuity, momentum, energy, and photoelectron equations as a function of altitude, latitude, and longitude. The equations are solved along the magnetic field line at each grid point, from 80 km in one hemisphere to 80 km in the other hemisphere, avoiding the need for upper boundary conditions. Since the Torr et al. [1990] study, some upgrades have been made to the model and some rate coefficients have been changed. To avoid listing just the changes and requiring the reader to refer to the earlier papers, we list the entire photochemical scheme in Table 3. It also allows us to correct typographical errors that appeared in (25), (38), and (43) of Table 1 of Torr et al. [1990].

For modeling the daytime thermosphere, the three most important input parameters to the model are the solar EUV and FUV flux spectrum, the photoionization cross sections, and the total electron impact cross sections. For computing photoelectron impact ionization, the solar flux below 310 Å is needed; for computing photoionization, the solar flux below 1025 Å is needed; and, in computing photodissociation of  $O_2$ , the solar flux in the Schumann-Runge continuum (1250–1750 Å) is needed. Since the Spacelab 1 mission flew at a time of



**Figure 5.** Segment of measured spectral sequence during which the field of view of the instrument was directed at a tangent ray height of approximately 140 km. Also shown are the synthetically modeled components of the spectrum (the individual NO  $\gamma$  and N<sub>2</sub> LBH band systems and principal atomic lines).

relatively low solar activity, the solar flux model used for that case is based on the F74113 Hinteregger reference spectrum from Torr et al. [1979], with the fluxes doubled below 250 Å [Richards and Torr, 1984]. Uncertainties in the solar EUV fluxes have recently been discussed by Richards et al. [1994], who developed a new solar EUV flux model (EUVAC). The EUVAC model also uses the F74113 solar flux reference spectrum but uses the Atmosphere Explorer E measured relative variations for the solar activity scaling. The basic difference is that the old model uses only the daily  $F_{10.7}$  index for scaling whereas the new model also uses the 81-day average  $F_{10.7}$ index. For the period of the Spacelab 1 mission, there is a negligible difference between the solar EUV fluxes from the two models. For higher levels of solar activity, such as prevailed at the time of the ATLAS 1 mission, the flux is scaled in each of 37 wavelength intervals according to Torr et al. [1985].



**Figure 6.** Comparison of the spectral segment taken at 25°N with one at 65°N. The solar zenith angles in the two cases differ by less than 3°. Also shown are the synthetically modeled NO  $\varepsilon$  bands.

The  $O_2$  dissociation frequency is determined according to *Torr* et al. [1980a]. The photoionization cross sections and the photoelectron impact cross sections are presented by *Richards and Torr* [1988, 1990] and *Fennelly et al.* [1992]. In summary (Table 4), the photoionization cross sections are best known; the electron impact cross sections are well known; and the total excitation cross sections are uncertain by a factor of 2. The calculated flux is inversely proportional to the total impact cross section.

A comparison of measured and modeled photoelectron spectra using the two different EUV reference spectra (F74113 and SC#21REFW), with the lower excitation cross sections in the case of the former, and the higher cross sections [*Strickland*  and Anderson, 1983] in the case of the latter, is shown by Richards and Torr [1988].

The neutral atmosphere is provided to the field line interhemispheric plasma (FLIP) model from the MSIS-86 model [*Hedin*, 1987]. When the airglow model is run for a particular day, with the appropriate solar flux and magnetic indices, the output provides a four-dimensional array (altitude, latitude, longitude, local time) through which the line of sight of the instrument can be projected, and the slant path intensity computed for any point along the orbit. Studies comparing the ISO observations of emissions from other atmospheric constituents obtained on the ATLAS 1 mission with the FLIP model have been reported elsewhere [*Torr et al.*, 1994, 1993a, b).

### 4. Photochemical Sources and Sinks of Nitric Oxide in the Thermosphere

A potential term diagram for NO is shown in Figure 3 (which is simplified from *Gilmore* [1986]). The  $\gamma$  bands arise from transitions from the  $A^2\Sigma$  state to the  $X^2\Pi$  state. The A state can be populated by various mechanisms, including resonance fluorescence of sunlight, cascade from higher levels, and photochemical sources. The (1,0) band is particularly bright in the dayglow because of its high-fluorescence efficiency [*Callear and Smith*, 1963]. Photochemical reactions are responsible for the removal of NO in addition to photoionization and photodissociation. Because the first ionization potential of NO is relatively low (9.25 eV), it is the only important atmospheric constituent that is ionized by the strong Lyman  $\alpha$  emission in sunlight (a fact already referred to above as the major source of the *D* region ionization).

The first measurements of the NO  $\gamma$  bands [Barth, 1964] yielded NO concentrations 2 orders of magnitude larger than those predicted by simple resonance fluorescence of sunlight [Barth, 1961]. In an attempt to bring the theory into better agreement with the measurements, Nicolet [1965a, b] introduced ionic reactions. Norton and Barth [1970] subsequently added a source of NO resulting from the interaction of N(<sup>2</sup>D) with O<sub>2</sub> and were able to obtain relatively good agreement in the magnitude of the concentrations. Lower-altitude measurements were then made by Meira [1971], and these required the development of models incorporating diffusive transport [Rusch, 1973] which were able to account for the shape at the lower altitudes. More recently, Siskind et al. [1990] have sug-



**Figure 7.** Example of spectral/altitude image. These are the same data shown in Figure 5 but with the altitude dimension shown resolved.



**Figure 8.** Measured spectrum at selected tangent ray heights as field of view is rolled down through the atmosphere. Also shown is the synthetic spectral fit to each spectral segment.

gested that solar soft X rays produce  $N_2^+$  which then in turn produces  $N(^2D)$  and hence NO.

The present photochemical scheme for the production of NO in the thermosphere is given in Table 5.

In Figure 4 we show the major photochemical sources and sinks of the  $A^2\Sigma$  state of NO for conditions appropriate to the March 26, 1993 roll sequence listed in Table 2 above. These calculations are for 61°N, 46°E. Over the entire altitude range of interest here, the major sources are the reactions of N(<sup>2</sup>D) and N(<sup>4</sup>S) with O<sub>2</sub>, and the principal loss mechanism is the reaction of N(<sup>4</sup>S) with NO, with a small contribution from the charge exchange reaction with O<sub>2</sub><sup>+</sup> near 110 km.

At high latitudes the long photochemical lifetime of NO makes it an important candidate for transport to lower altitudes where it becomes involved in the catalytic removal of ozone (NO +  $O_3 \rightarrow NO_2 + O_2$  and  $NO_2 + O \rightarrow NO + O_2$ ).

### 5. Spectral Images at 140 km Tangent Ray Height

Figure 5 shows a spectral segment obtained on day 89, 1992 (March 29) at 0950 UT. The configuration for this (illustrated



in Figure 1) was discussed in section 2 (nose of the orbiter pitched down so that the line of sight pointed at a nearly fixed tangent ray height (TRH) of approximately 140 km). During this sequence the full spectral range of the instrument was scanned 6 times, obtaining representative spectra covering latitudes from the equator to almost 60°N (Figure 2). The data shown in Figure 5 are taken at 25°N. The lower panels in Figure 5 show the synthetic spectra for the various systems in this portion of the spectrum (NO  $\gamma$ , N<sub>2</sub> Lyman-Birge-Hopfield (LBH) and the major atomic features). The synthetic spectra have been generated using the line-by-line code of *Arnold et al.* [1969].

As can be seen from Figure 5, the dayglow spectrum in the lower thermosphere in this wavelength window is dominated by NO  $\gamma$  and N<sub>2</sub> LBH bands. Figure 6 shows a comparison of the low-latitude spectrum of Figure 5 with the higher-latitude spectrum of this sequence taken at 65°N. The two are quite similar as far as the LBH bands are concerned. The NO  $\gamma$ bands are significantly (>2×) brighter in the higher-latitude case. Also, the NO  $\varepsilon$  bands are clearly present in the higherlatitude spectrum, even at these altitudes, as can be seen at the locations of the (1,0) and (3,1)  $\varepsilon$  bands. In the sections below we shall discuss further the contributions by the NO  $\varepsilon$  bands and the NO  $\delta$  bands. The overlaying of the synthetic spectral fit to the observations in the FUV has been shown by *Torr et al.* [1994].

The data shown in Figure 5 are averaged across the 10 altitude segments of the image. In Figure 7 we show a portion of this spectrum in which the adjacent altitude elements are resolved. The changing characteristic of the spectrum in this simultaneously acquired image is evident. The adjacent rows in the spectrum each represent 1.7 km of altitude, so that the full image spans 129.8 to 145.4 km. The N<sup>+</sup>( ${}^{5}S$ ) atomic doublet at 2139.7/2143.6 Å, which overlaps the left branch of the NO  $\gamma$  (1,0) band, can be seen to decrease in intensity with decreasing altitude in this altitude range.

The temperature characteristics of these spectral features are discussed further in section 6.

## 6. Altitude Variation of NO Concentration and Temperature in the Thermosphere

The data sets used here are those acquired in the vehicle roll maneuvers in which the crew first positioned the orbiter in such





Figure 9. Comparison of column concentration of NO derived from the data shown in Figure 8 with that computed from the global airglow model.

a way that the instrument line of sight was tangential to the orbit altitude of 300 km. The direction of the line of sight was back along the orbit, into the wake. The shuttle was then rolled at a controlled rate to scan the narrow dimension of the field of view  $(0.007^{\circ})$  down through the atmosphere. In the course of this observing sequence, the wavelength was cycled between six selected grating steps in each of the spectrometers, rather than executing complete wavelength scans



Figure 10. NO concentrations derived from the column concentrations inferred from the measured slant path intensities. Shown for comparison are the concentrations from the FLIP model, as well as various measured vertical NO profiles [Rusch, 1973; Gerard, 1984; Cleary, 1986; Siskind et al., 1990; Eparvier et al., 1992].



Figure 11. Illustration of the sensitivity of the relative magnitudes of the P and R branches of several NO  $\gamma$  bands to temperature.

over the entire range accessible to the instrument. During the course of one of these rolls, the ground track of the vehicle covered almost a quarter of an orbit. The latitude/longitude covered is illustrated in Figure 2. Parameters such as date, local time, solar zenith angle and magnetic activity are shown in Table 2.

Figure 8a–8j shows the spectral segment from 2090 to 2270 Å (a simultaneously acquired spectral image) at the various tangent ray heights at which the measurements were made. Also shown in the figure is the synthetic fit to the spectrum at each altitude. Included in the spectral fit are the NO  $\gamma$  bands and N<sup>+</sup>(<sup>5</sup>S) atomic features. The principal features are labeled. In addition to these, bands from the NO  $\varepsilon$  system can be seen at the lower altitudes.

The self-absorption of the (1,0) band at 105 km is evident in the fact that the synthetic spectrum is much larger than the

measurements for this band, while the (1,1) band is well fitted. The self-absorption can be seen up to approximately 200 km. Above 200 km, a good fit is obtained to the (1,0) band. By 145 km the N<sup>+</sup>(<sup>5</sup>S) doublet, which lies on the left hand branch of the  $\gamma(1,0)$  band, starts to become evident. The two lines of the doublet at 2139 and 2143 Å are clearly resolved. By 269 km, this is the dominant feature in this portion of the spectrum. We return to the N<sup>+</sup>(<sup>5</sup>S) feature below.

Two parameters can be derived from the spectral fits to the NO  $\gamma$  bands shown in Figure 8. The first is the column concentration of NO. The second is the effective NO temperature along the line of sight.

**Column concentration:** From the data measured in this sequence we can determine the total column concentration of NO along the line of sight at each point along the orbit and obtain an altitude/latitude profile. For example, using the (1,1) band, the



Figure 12. Comparison of temperatures derived from the synthetic spectral fit to the data shown in Figure 8 with the neutral temperature given by MSIS-86 at each tangent ray height (TRH), and the computed effective temperature that would be seen looking along this line of sight.

column concentration of nitric oxide is obtained by using the resonance fluorescence g factor for the (1,1) band as follows:

 $[NO]_{col} = I(Rayleighs) \times 1 \times 10^{6}/g_{1,1} molecules/cm^{2}$ (1)

where I is the measured intensity of the (1,1) band in Rayleighs and the factor of  $10^6$  converts from Rayleighs to photons per



**Figure 13.** NO contribution function as a function of altitude for the concentration profile derived from these dayglow measurements and shown in Figure 10.

second. We have used the value of  $2.1 \times 10^{-6}$  photons molecule<sup>-1</sup> s<sup>-1</sup> for the g factor of the (1,1) band, and  $7.7 \times 10^{-6}$  for the (1,0) band [*Barth*, 1965].

The results are shown in Figure 9 with the column concen-



**Figure 14.** Contribution to the signal measured at a TRH of 80 km from each 10 km layer between 80 km and 300 km. The full line curve is computed using the FLIP model [NO] and the dashed curve is that using the [NO] inferred from these data.



Figure 15. Temperature altitude image obtained with the center of the field of view directed at approximately 140 km, with the computed effective temperature shown for comparison.

trations predicted by our interhemispherically coupled airglow model for comparison. The model is found to agree well with the data above  $\sim 230$  km (which corresponds to lower latitudes), but below this the model predicts less column abundance of NO than is obtained from the dayglow measurements by a factor of 2-3. Figure 10 shows the NO concentration profile derived from the column concentration profile of Figure 9 (using the very simple spherical layer inversion technique), with various other measurements and our FLIP model shown for comparison. Bearing in mind that the profile measured from ATLAS 1 is one that covers quite a range of latitude, while the earlier measurements are strictly vertical profiles, the present results lie on the high side of the six profiles shown for comparison. However, [NO] is known to exhibit great variability with solar and geomagnetic activity and with latitude. The ISO inferred concentrations also fall off less steeply with height than does our model.

**Temperature:** The shape and relative size of the P and R branches of the  $\gamma$  bands are dependent on the NO temperature, and hence the neutral temperature. This is illustrated in Figure 11 in which we show a number of these bands modeled for a range of thermospheric/mesospheric temperatures from 150 to 850 K. By fitting the bands shown in Figure 8, we obtain the temperatures shown in Figure 12. This approach was used by *Erwin* [1985] and also *Bosserman* [1989] to determine NO temperatures using the data obtained with the ISO during the Spacelab 1 mission. Subsequently, the approach was used by *Barth and Eparvier* [1993] to obtain temperatures from a highlatitude, daytime rocket experiment. We have fitted the (1,0) band above 150 km and the (1,1) band at the lower altitudes. Above 150 km, the self-absorption of the (1,0) band is small and this band has the advantage of considerably brighter intensity. In the region of overlap, these two bands agree in temperature. Also shown in Figure 12 is the neutral temperature at the tangent ray height from the MSIS-86 model. However, as is the case with the column concentrations, the measurement is an integral of all the NO along the line of sight, and may not be well represented by the temperature at the tangent point itself, even though the [NO] is falling steeply with altitude. In other words, we need to estimate the extent to which this temperature fit is modified by the emission from the higher (hotter) altitudes along the line of sight. To do this we have computed the contribution function from each 10 km layer between the shuttle altitude of 300 km and the tangent ray height. The results are shown in Figure 13. When looking at a TRH of 100 km, for example, 30% of the NO in the line of sight is in the 10 km wide layer centered at 100 km. The remaining 70% is from higher altitudes. When looking at a TRH closer to the shuttle altitude, say 250 km, a larger percentage (50%) is from the region of interest. Figure 14 shows the contribution to the measured signal when looking at a TRH of 80 km, from each 10-km layer between the orbital altitude and 80 km. This function is shown for both the FLIP model [NO] and the [NO] inferred from these measurements. It can be seen from Figure 14 that the contribution of the hotter NO at higher altitudes falls off rapidly with increasing height. Using the model or measured [NO], and the temperatures at the tangent point from the MSIS-86, we can compute the effective (column) temperature at each point from:

$$T_{\rm eff} = \sum_{\rm TRH}^{300} Tn[\rm NO]ds \bigg/ \sum_{\rm TRH}^{300} [\rm NO]ds$$

where the shuttle is orbiting at 300 km, and the line of sight passes through the TRH and on out to 300 km on the far side. This effective temperature is also shown in Figure 12. Below  $\sim$ 130 km, the data indicate a cooler temperature than is predicted by the model.

From Figure 12 it can be seen that near 140 km the neutral temperature is changing by about 200 K over 20 km. Returning to the data used in section 5, in which the length of the spectrometer entrance slit was positioned perpendicular to the horizon, a simultaneous altitude image of 17 km is obtained. The 0.65° field of view is imaged onto 10 adjacent lines on the detector, with each line spanning 1.7 km at this tangent ray height. In the case of the spectra shown in Figure 5, we averaged the data across the adjacent altitude segments. In Figure 7 we showed the same spectral segment resolved into the altitude slices. By spectrally fitting each of the 10 components, we obtain a slant path temperature for each 1.7 km altitude segment, and these are shown in Figure 15 together with the effective temperature expected for this particular line of sight



**Figure 16.** Illustration of the self-absorption of the NO  $\gamma$  (1,0) band as a function of altitude: (a) comparison of NO  $\gamma$  (1,1) and (1,0) band slant path intensities, also showing the (1,1) band normalized to the (1,0) band at 220 km; and (b) ratio of the band intensities shown in (a), with a weighted smoothed curve drawn through the ratios.

 Table 6. Geophysical Parameters for Spacelab 1 Sequence

Parameter	December 7, 1983
Shuttle attitude	Fixed tangent ray height (~104 km)
Local time	1607
Solar zenith angle	84.8
Latitude	37°N
Longitude	175°E
$F_{10,7}$ -cm flux	98
Ap	11

from the model. As in the case of the roll data discussed above, the 140 km TRH data indicate that the NO temperature is in reasonable agreement with the model at these altitudes.

Self-absorption: We have already discussed the fact that the  $\gamma(1,0)$  band is self absorbed at the lower altitudes. In Figure 16a we show a comparison of the altitude profiles of the (1,1) and the (1,0) bands. In Figure 16a we also show the results of normalizing the slant path intensities of these two bands at 220 km. While the fall off in the (1,0) band below 150 km is evident, there is absorption (although small) indicated up to  $\sim 200$  km, as was also the case in the spectral fits shown in Figure 8. In Figure 16b we show the ratio of the two normalized intensity profiles.

N<sup>+</sup>(<sup>5</sup>S) **Doublet:** A prominent feature in the spectra shown in Figure 7 at the higher altitudes is the doublet due to N<sup>+</sup>(<sup>5</sup>S) at 2139.7/2143.6 Å. In Figure 17 we show the intensity of this feature as a fraction of the sum of the N<sup>+</sup>(<sup>5</sup>S) doublet and the NO  $\gamma$  (1,0) band. From Figure 17 it can be seen that the atomic doublet would contribute ~10% or less of the unresolved N<sup>+</sup>(<sup>5</sup>S) and NO  $\gamma$  (1,0) emission below 200 km.

# 7. Altitude Imaging of the Lower Thermosphere/Upper Mesosphere

Using the same altitude imaging configuration discussed above and illustrated in Figure 1, several segments were flown during the mission with the center of the field of view directed down into the lower thermosphere and upper extent of the mesosphere. In Figure 18 we show an example of a spectral/ altitude image segment from an observation sequence taken on March 25, 1992, at 56°N, 63°E. The ground track for the full segment is shown in Figure 2. This particular image spanned the altitude range from 93 to 107 km with each row on the detector corresponding to a 2 km altitude segment. We have derived temperatures by fitting the (1,1) band, and these fits are shown in Figure 19, as compared with the anticipated effective temperature computed from our modeled NO and the MSIS neutral temperature model. Our NO model is not rigorous below 100 km at present because it neglects odd N production from photoelectrons with energies greater than 100 eV. As a result, the computed effective temperature is a rough estimate of what we would expect to measure at the low altitudes and is shown in Figure 19 to provide a point of reference rather than a quantitative comparison. Given the uncertainties in the model, the agreement in temperature is good, and we believe that the structure apparent in the data represents real structure. Any effect on the temperatures due to the higher altitudes would be common to each of these adjacent altitude segments and unlikely to introduce such variability. This altitude structure is probably similar to that reported in various ground-based lidar observations. This particular sequence ob-



Figure 17. Percentage contribution to unresolved NO  $\gamma$  (1,0) band and N<sup>+</sup>(<sup>5</sup>S) doublet of the N<sup>+</sup>(<sup>5</sup>S) doublet.

tained a number of such temperature images over a range of latitudes. (As the transitions ending in the v' = 0 level are self-absorbed at these altitudes, the populations are not correctly adjusted for these.)

The first spectral/spatial images of lower thermospheric NO was obtained with the ISO during the Spacelab 1 mission on December 7, 1983. As was done in the case of the ATLAS 1 sequence shown here, the field of view was directed at lower thermospheric altitudes, with the entrance slit perpendicular to the horizon, by controlling the orientation of the shuttle. This orientation was maintained while the shuttle covered a full orbit, during which time a series of six spectral scans were made with the instrument, covering a range of latitudes. The ground track for that set of observations is shown in Figure 20. An example of one of the Spacelab 1 spectral images is shown in Figure 21. These measurements were made with the tail of the orbiter pitched down toward the Earth so that the field of view was centered at approximately 104 km. Relevant parameters for this data set are summarized in Table 6.

The temperature profile derived from the measurements is shown in Figure 22 compared with the effective temperature computed using MSIS neutral temperatures and modeled [NO] for the ATLAS 1 roll sequence shown for comparison. The Spacelab 1 data (again, fitting the (1,1) band at these lower heights) show an effective temperature that is much hotter than can reasonably be expected from the model. The exospheric temperature at the time of the Spacelab 1 mission would have been closer to 1000 K, lower than the 1200 K of the ATLAS 1 mission, so the modeled effective temperature for that case would be even lower than that shown for the ATLAS 1 case. The Spacelab 1 spectral data have shown repeated evidence for significant vehicle induced emissions. ATLAS 1 and the later missions appear to be relatively clean in terms of contamination. Major changes in the prelaunch processing of the vehicle and the payload have been introduced since 1983. It is therefore possible that the hot temperatures indicated in Figure 22 are due to a significant vehicle induced population of hot NO, as NO is a likely major component of the induced environment.

We do not have accurate pointing information for the earlier Spacelab 1 data, and for that case the altitude uncertainties could be as large as 5 km or more at lower thermospheric altitudes. However, for the ATLAS 1 mission the pointing knowledge was greatly improved. The ISO tangent ray heights are computed from post mission orbiter position and attitude data. The dominant source of uncertainty is the determination of the instrument line of sight direction from the orbiter IMU (the primary vehicle star pointing system) attitude data. Pointing uncertainty for the ISO is believed to be less than 0.1° (3 sigma). The Pointing and Alignment Work Station (PAWS) investigation during ATLAS 1 determined a total pointing uncertainty of 0.2° for pallet mounted instruments, including thermal drift (0.08°) and instrument alignment variations (0.15°). ISO observations were conducted from attitudes in which the payload bay would have been thermally stable compared to the solar pointing conditions under which the PAWS determination was made, so thermal drift should be negligible for the ISO during these observations. Furthermore, the alignment of ISO with the pallet as measured on the ground before and after the mission, varied by only 0.02°. Using post mission position data, the uncertainty in orbiter position is a negligible source of error in tangent ray heights. The tangent ray height calculations are done using a fully ellipsoidal Earth, and there are no approximations in the calculation other than assuming that there are no higher-order terms in the geoid. The maximum error from this source would be 100 m. For the ISO the



**Figure 18.** Example of lower thermospheric simultaneous spectral/altitude image. The center of the field of view is directed at 99 km. Also shown are the synthetic fits to the spectral images. Note: the intensity scale must be multiplied by 20 to get the actual intensities.

200 250

1

(A02/A) YTIZNƏTVI



### (A02/A) YTISNƏTVI

17,406

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Figure 18. (continued)

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Figure 19. Temperature inferred from the NO  $\gamma$  bands shown in Figures 18. Also shown is the computed effective temperature.

uncertainty in the tangent ray height for these sequences should be less than 3 km (3 sigma).

### 8. NO ε Bands

As was shown in Figure 6, the NO  $\varepsilon$  bands are evident in the high latitude spectra, even above 140 km. The NO  $\varepsilon$  bands are relatively weak but observable in the lower thermospheric data as can be seen from Figure 23. Figure 23 shows the average of the nine lines that comprise the image of 93 to 107 km. During

the ATLAS 1 mission, spectral images near 100 km were not taken at a wavelength increment that includes the brighter  $\varepsilon$ bands at shorter wavelengths, but these data were acquired on the earlier Spacelab 1 mission (Figure 21). Using the Spacelab 1 data, *Bosserman* [1989] was able to empirically infer the Frank-Condon factors. Since the emissions from the (0,0), (0,1), (0,2), and (0,3) progression all have the same excitation mechanism, the relative intensities of these bands are determined by the Frank-Condon factors. Figure 24 shows Bosser-



Figure 20. Ground track of shuttle during the sequence of lower thermospheric observations taken during the Spacelab 1 mission. A sequence of eight complete spectral scans was made and the numbers shown on the track indicate the location at which the NO gamma (1,0) and neighboring bands were measured. The data shown in Figure 21 were taken from cycle 2, looking back along the track.



**Figure 21.** Spectral image measured during the Spacelab 1 mission on December 7, 1983. The data were obtained at a tangent ray height of 104 km, at 37°N and 175°E. The local time was 1007 and the solar zenith angle was 85°.

man's comparison of the Frank-Condon factors used in his synthetic spectral fit [*Nicholls*, 1964; *Spindler et al.*, 1970] versus those determined from the data.

### 9. NO δ Bands

The  $\delta$  bands have been reported in the twilight and nighttime emission spectrum [Cohen-Sabban and Vuillemin, 1973;



Figure 23. Example of spectral image shown in Figure 18, including the grating step to the short wavelength side. Several nonoverlapped  $\varepsilon$  bands are evident, as well as weaker NO  $\gamma$  bands.

Feldman and Takacs, 1974; Gerard, 1975]. Gerard [1975] reported a maximum total system intensity of 220 R, with a (0,1) band intensity of 50 R. Feldman and Takacs [1974] reported a comparable 83 R for the (0,1) band. The source was believed to be preassociation of oxygen and nitrogen atoms above 140 km:

$$N(^{4}S) + O(^{3}P) \rightarrow NO(a^{4}\Pi) \rightarrow NO(C^{2}\Pi, v' = 0)$$
$$\rightarrow NO(X^{2}\Pi) + hv_{s}$$



Figure 22. Temperature inferred from the NO  $\gamma$  bands shown in Figures 21. Also shown is the computed effective temperature.



Figure 24. Theoretical versus empirically derived Frank-Condon factors for the NO  $\varepsilon$  bands [Bosserman, 1989] measured during the Spacelab 1 mission.

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This source process is supported by the relative intensity of the  $\gamma$  (1,0) and  $\delta$  (0,0) bands in the *Feldman and Takacs* [1974] data, in which these two bands were found to be comparable in intensity. If the source were fluorescence, the  $\gamma$  (1,0) band would be considerably brighter. The chemiluminescent  $\delta$  bands are more easily seen at night because they are not swamped by the much brighter fluorescence bands of the  $\gamma$  and  $\varepsilon$  systems of NO.

Laboratory measurements by *Callear and Pilling* [1970a, b] were found to be consistent with a fluorescence efficiency of 0.03, and it is this value that has been used in determining the branching for NO photodissociation products. Using this low value for the fluorescence efficiency, one would not expect to see the NO  $\delta$  bands in the dayglow. However, *McCoy* [1983] and *Cleary* [1986] have reported the detection of the  $\delta$  (1,0) band in daytime rocket experiments, and inferred the fluorescence efficiency to be much higher, 0.2 and 0.25 respectively. In fact, *Cleary* [1986] measured a column emission rate for the  $\delta$  (0,1) band at TRH of 100 km of 1–2 kR and concluded that the



Figure 25. Portion of spectrum from ATLAS 1 roll sequence showing 2 TRHs: (a) 107 km and (b) 162 km: with NO  $\gamma$  and  $\varepsilon$  and N<sub>2</sub> LBH band synthetic fits.



fluorescence source of the  $\delta$  bands is considerably stronger than the chemiluminescent source.

In an assessment of the Spacelab 1 ISO data shown in Figure 21, *Bosserman* [1989] inferred a fluorescence efficiency of 0.32 based on the intensity of the feature at 1984 Å which he assumed to be the  $\delta$  (0,1) band.

The ATLAS 1 data provide an opportunity to look for confirmation of the presence and relative intensity of  $\delta$  bands. In laboratory measurements of the associative recombination process, only the v' = 0 level is populated [Groth et al., 1971], but if the results of Cleary [1986] are correct and fluorescence is the dominant daytime source of the  $\delta$  bands, then bands from higher v' levels may also be present.

In the lower thermospheric imaging sequences made from the ATLAS 1 mission, in order to increase the temporal resolution, not all grating steps were used. As a result, we do not have data containing the 1984 Å feature in this set. We do, however, have this portion of the spectrum in the roll maneuver sequence, and Figure 25a shows the lower-altitude (TRH = 107 km) image at this grating step. There is clearly a bright feature near 1984 Å. Figure 25b shows a higher-altitude image (TRH = 162 km) at this same grating step. Also shown in Figure 25a and 25b are the NO  $\gamma$  and  $N_2$  LBH synthetic spectral fits.

The slant path intensity of the  $\delta(0,1)$  band is ~1 kR at a TRH of 107 km. By 162 km (correcting for the underlying LBH (6,13) band) it has dropped to ~100 R. These results appear to support the conclusion of *McCoy* [1983] and *Cleary* [1986] that the fluorescence efficiency of the  $\delta$  bands is considerably higher than that determined by *Callear and Pilling* [1970a, b].

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#### 10. Conclusions

In this paper we have analyzed FUV data over the wavelength range of approximately 1900 to 2400 Å taken during the Spacelab 1 and ATLAS 1 shuttle missions with the Imaging Spectrometric Observatory. These data are used to derive altitude profiles of NO concentrations and temperature which are compared our global model and with other measurements. The NO concentrations during the ATLAS 1 mission lie to the high side of a sample of earlier measurements and are higher than predicted by our model. Because of its two-dimensional focal plane detectors, allowing imaging of wavelength and altitude, the flights of the instrument on the 1983 Spacelab 1 and 1992 ATLAS 1 missions have provided the first simultaneously acquired altitude/spectral images of NO concentrations in the thermosphere. From the shape of the branches of the NO  $\gamma$ bands, we have determined the neutral temperature. The results show that this is a viable technique for remote sensing of temperature down to the 80 km altitudes studied here, provided that data are also available on the NO at altitudes between the spacecraft and the tangent ray point, as the effect of the overlying NO is not insignificant for probing at these lower altitudes. The lower thermospheric temperature profile is quite structured in altitude. Temperature images obtained from the earlier Spacelab 1 mission indicate much higher than expected temperatures, probably due to a hot NO population in the induced environment of that mission.

The NO  $\delta$  band intensities are consistent with other findings of a higher fluoresence efficiency than had been earlier determined from laboratory measurements.

The spectral resolution of the instrument is sufficient not only to separate the NO  $\gamma(1, 0)$  band and the N<sup>+</sup>(<sup>5</sup>S) doublet but also to resolve the two lines in the doublet. Below 200 km, the N<sup>+</sup>(<sup>5</sup>S) feature contributes less than 10% to the unresolved combination of N<sup>+</sup>(<sup>5</sup>S) and NO  $\gamma$  (1,0). The selfabsorption of the NO  $\gamma$  (1,0) band is evident up to 150 km and appears to be present up to approximately 200 km.

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