Vehicle/Atmosphere Interaction Glows: Far Ultraviolet, Visible, and Infrared

G. Swenson
University of Illinois, Urbana, Illinois

Prepared for Marshall Space Flight Center
under Contract NAS8-40579
and sponsored by
the Space Environments and Effects Program
managed at the Marshall Space Flight Center

October 1999
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National Aeronautics and Space Administration

Marshall Space Flight Center • MSFC, Alabama 35812

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<td>AE</td>
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<tr>
<td>AO</td>
<td>atomic oxygen</td>
</tr>
<tr>
<td>CMG</td>
<td>control moment gyro</td>
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<tr>
<td>DE</td>
<td>Dynamic Explorer</td>
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<td>EISG</td>
<td>Experimental Investigation of Spacecraft Glow</td>
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<td>FUV</td>
<td>far ultraviolet</td>
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1. INTRODUCTION

Spacecraft glow studies have been performed with the Experimental Investigation of Spacecraft Glow (EISG), which flew on Space Transportation System (STS)–62 as part of the Office of Aeronautics and Space Technology (OAST) payload. The experiment flew as a Hitchhiker payload. Glow studies have been done since they were discovered on STS–3, as crewmen viewed the phenomena from the aft flight deck. Film pictures documented the early observed phenomena, as exemplified in figure 1.

Figure 1. An image of Shuttle glow taken on STS–8. The orange glow is observed to extend away from the surface—an early emission mystery.
The EISG payload consisted of visible, ultraviolet (UV) and far ultraviolet (FUV), and infrared (IR) instrumentation. A schematic of the payload is shown in figure 2, and a picture of the instrument integration nearing its completion is shown in figure 3. The instrumentation was built and assembled by a team managed from the Lockheed Palo Alto Research Laboratory (LPARL) but engineered with both LPARL and the Lockheed Engineering and Systems Company (LESC), Houston, TX. The NASA management of the experiment was provided by the materials group at NASA Johnson Space Center and the payload integration and operations support was performed by the Hitchhiker office at NASA Goddard Space Flight Center. It was truly a well chosen path for the payload system as engineering and support groups from both NASA and Lockheed worked as a team to accomplish the successful development and mission, achieving most of the objectives set initially.

Figure 2. Schematic describing the EISG, which participated in the STS–62 mission, shown integrated to the Hitchhiker bridge structure. The optical detectors viewed into a mirror which in turn directed the viewing fields over two material samples. The left sample was a white Chemglaze™ (A276) and the right, a black carbon paint (Z306).
On orbit, the integrated payload can be seen from the aft flight deck camera on the orbiter (fig. 4) and from the wrist camera on the remote manipulator system (RMS) (fig. 5). The wrist camera complemented the observations as the view along the plate clearly showed ram glow on the sample plate (fig. 6). The classical experiment to test atom exchange with molecular nitrogen (N) was performed with a gas release which released the plum on and in front of the sample plate. The experiment proved that molecular N did not atom exchange at the orbital velocity. In fact, the gas cloud sufficiently attenuated the ram oxygen (O) beam (and energy) which resulted in extinguishing the glow (fig. 7).

Figure 4. The payload bay from the aft flight deck near sunrise on STS–62. A glint of sunlight is seen on the orbiter bay. Glows can also be seen on the port side the vertical stabilizer and the starboard orbital maneuvering system (OMS) pod. The EISG payload is visible on the mission peculiar experiment support structure (MPRESS) in the lower left with the logo on the insulation blanket.
Figure 5. A view of the EISG experiment in the payload bay during an orbital daytime. The orbiter RMS wrist camera is being used to make the observation along the normal to the sample plate.

Figure 6. An orbital nighttime image from the same camera with the same viewing geometry as figure 5. Note the glow on the sample plate.
Figure 7. The same view as shown in figures 5 and 6. The glow visible on the sample plate in the top image is extinguished during the N$_2$ gas release in the bottom panel. This is due to the release gas becoming collisionally thick to the ramming O, and consequently, reducing the fluence and O velocity (and energy) necessary for the heterogeneous chemistry to take place which normally leads to the glow reaction.

This data analysis task has been supported by the Space Environmental Effects (SEE) program and managed from the materials group at the NASA Marshall Space Flight Center (MSFC). This study evolved models of the visible, FUV, and IR glows for engineering replication of glow processes. The models and suggested mitigation are contained in this report.
2. VISIBLE REGION

Visible glow has been reported on ram surfaces of the Atmospheric Explorer-E (AE), Dynamic Explorer-B (DE). The spectral observations from the AE and Space Shuttle suggest a "red" glow, although the course spectral shape is not the same for the two spacecraft. The AE and DE data support an O intensity dependence with altitude. This O dependence is qualitatively supported in the STS observations as well. The major difference between the two would appear to be the surface e-fold of the emission, determined to be 20 cm by Yee and Abreu and verified by Swenson et al. for the STS observations. The ram angle effect of the AE glow appears to closely follow a cosine relationship with the ram angle. The ram angle of the stabilizer seems to produce slightly more glow at high ram angles than the cosine effect, yet an unexplained phenomenon.

It was later found from the STS database that the intensity for similar altitudes showed significant differences in brightness. Analysis has revealed that the rammed surface temperatures were significantly different for these measurements. Subsequent analysis uncovered an explanation which is consistent with an earlier theory that supports the fact that nitric oxide (NO) is held in bond on the surface for ramming O atoms to form nitrogen oxide (NO\(_2\)) in three-body recombination, where the NO\(_2\) vibrationally relaxes to produce the glow. It was also suggested that the 0.14-eV bond energy of the NO would be consistent with the lesser intensity seen on the AE spacecraft since the AE spacecraft surface was much warmer than the STS tile surfaces. Yee et al. suggested that a portion of the AE glow was also likely a result of NO\(_2\) emission.

An analytical glow intensity model has been developed to fit the experimental results by Swenson and Mende which was later refined by Swenson and Lairden. This model assumes a cosine ram angle dependence, an O scale height dependence, and a surface temperature dependence. The validity of the model in the spectral range (4,000–8,000 Å) is supported by the good fit of the model to the observed slit spectral intensity flown on board the STS–41–D and STS–62.

The model takes the form:

\[
I_{x,\phi,T,\lambda} = 0.084 \cdot \frac{O_z}{O_{250}} \cdot \cos(\phi_r) \cdot e^{930/T_s} \cdot \cos(0.0005 \cdot |6,800 - \lambda|)^3
\]

where

- \(I_{x,\phi,T,\lambda}\) = intensity, Rayleigh/Angstrom
- \(O_z\) = atomic oxygen (AO) density at altitude, z
- \(O_{250}\) = AO density at 250 km
- \(\phi_r\) = the angle that a surface makes with ram, degree (0–90)
- \(T_s\) = surface temperature, K (190–450)
- \(\lambda\) = wavelength, Angstrom (4,000–8,000).

An example of the model output may be found in figure 8. The assumptions were a surface temperature of 300 K, ram angle of 45°, and altitude of 400 km.
2.1 Use of the Model

The model is coded in interactive digital language (IDL) with the following input and output parameters:

Input parameter:
- \( t_{\text{surf}} \): surface temperature, 190–450 K
- \( \text{ram\_angle} \): ram angle of the Shuttle position, 0°–90°
- Altitude: altitude of the Space Shuttle

Output parameter:
- Wavelength: wavelength of the investigation
- Intensity: spectral intensity of the surface glow.

The altitude dependence is calculated as the ratio of AO intensity at altitude \( z \) and at 250 km. AO intensity between 200–600 km with 20-km increments was derived using MSIS86 model\(^{12}\) using a set of default input parameters. Users can run the MSIS86 model using the parameters other than the default values to get the dependence for their special case.

To run the program, type “.run model_vs” at the IDL prompt and type the input parameters in response to the prompts. The resulted model will be plotted on the screen. The output parameters will be written into an ASCII file named “output.txt”. The output figure can also be plotted to a postscript file and printed to a postscript printer using the switch flag. A model result for the indicated conditions is shown in figure 8.

Enhanced visible glows 557.7- and 630-nm observed due to NO products from hydrazine doping of surfaces.\(^{13}\) Enhanced O 557.7- and 630-nm emissions were also observed.\(^{14}\) Visible spectral enhancements due to thrusters have not been modeled here.

![Shuttle Vertical Stabilizer Glow](image)

Figure 8. A model output of the visible glow for the following conditions: surface temperature = 220 K, ram angle = 10°, and altitude = 300 km. The glow is due to surface recombination which leads to NO\(_2\) emission.
3. ULTRAVIOLET REGION

Historically, UV and FUV emissions have been observed on a number of spacecraft. UV glows were reported in the 2,800 and 3,371 Å channels of the AE spacecraft. The UV glows were considerably weaker than glows in visible wavelengths on that spacecraft. It is assumed that the UV glows from NO might likely be on ram surfaces. Kofsky and Swenson and Meyerott have proposed recombination of N on the surface as being responsible. Swenson and Meyerott have found that a large source of N is likely in the plow cloud of low-altitude spacecraft due to atom exchange with the ramming atmosphere. This mechanism predicts a source flux and altitude distribution that supports the observations. Meyerott and Swenson have described a set of possible surface interactions likely with fast N recombination, including the possibility of 958 Å and 2,800–3,400 Å (Gaydon-Herman bands) from excited N recombination. Meyerott and Swenson presented a chemical explanation for fast recombination of N which would result in an N₂ upper state that would cascade to produce a source of N₂ Lyman-Birge-Hopfield (LBH) bands as observed by Torr and Torr. Hunton and Machuzak have verified the production of N and NO products in O-bombarded thruster plumes.

The EISG was flown on STS-62 in March, 1994. Many spectra were recorded by an FUV spectrometer that confirmed the existence of the UV and FUV surface glow. The major bright glow appeared to be of sporadic nature in the 2,200–3,000 Å spectral region. In particular, in the middle grating position the FUV channel displayed sporadic anomalous emission in the decent portion of the EISG 6 and 7 elliptical orbits when the Space Shuttle took the position where the sample plate was directed into ram. This sporadic emission featured a concentration of emission of energy in the short wavelength range between 2,200 and 3,000 Å. In the laboratory experiment, a fast O beam was directed into the N₂ and a sample plate to simulate the ram fluence. The primary emission of the lab glow is NO. The STS-62 and laboratory measured spectra, and the NO band heads are shown in figure 9. Laboratory results of similar experiments with IR measurements have been reported by Upschulte et al. The similarity between lab and ram glow indicates that the spectral bands emitting are the same but the vehicle glow relative to the laboratory observation was a rotationally hot emission, likely due to the extremely high recombination temperature.

These constituents that produce the UV surface glow can be scavenged from the atmosphere. In addition, monomethyl hydrazine thruster effluents interacting with the atmosphere are observed to produce large amounts of N and NO, likely through O collisions with exhaust products of vibrationally enhanced molecular N. These products produce both gas phase and surface (heterogeneous) reactions, resulting in UV and FUV glows.
Figure 9. A plot of FUV data and model information. The heavy line is the flight spectrum of FUV glow subsequent to thruster effluent doping of the EISG surface. The thin line is a spectrum taken with the same instrument of a laboratory experiment. The dots describe NO $\gamma$ and $\beta$ bandheads which are shown to correspond to the intensity peaks in the spectra. The flight spectra is "filled in" due to the effective high temperature of the on-orbit chemistry.
3.1 Computer Simulation of the Ultraviolet Surface Glow

To further investigate mechanisms of vehicle glow, a spectral model has been developed. The spectral model is based on the assumption that the vehicle glow is mainly from the emission of the NO $\beta$, $\delta$, and $\gamma$ bands. For the vibrational band, the wave number of the spectrum is given by Herzberg,$^{26}$

$$
\nu = \nu + \omega_e'(v' + 1/2) - \omega_e x_e'(v' + 1/2)^2
+ \omega_e y_e'(v' + 1/2)^3 + \ldots - \omega_e''(v' + 1/2)
- \omega_e x_e''(v'' + 1/2)^2 + \omega_e y_e''(v'' + 1/2)^3 + \ldots \right)
$$  \hspace{1cm} (2)

The coefficients were given by Herzberg$^{26}$ in the form of a table. Tanaka$^{27}$ found that in the NO afterglow the emission intensity is about the same for $\beta$, $\delta$, and $\gamma$ bands systems. Therefore, in our model spectrum the weights for $\beta$, $\delta$, and $\gamma$ bands are taken to be equal. The magnitude of the composite spectrum is normalized to the measured spectrum. According to Gaydon,$^{28}$ the emission intensity from the higher vibrational state of the band is weaker. However, in our simulation we still include the higher vibrational transitions because the theoretical calculation shows emission from some higher vibrational transitions cannot be ignored. In fact, we included in our model up to quantum number 23 and to the upper energy level 15 for $\gamma$, 19 for $\beta$, and 5 for $\delta$ bands.

3.2 Running the Programs

A FORTRAN90 program "synth_new.f" calculates the bandhead positions for $\gamma$, $\beta$, and $\delta$ bands and uses these calculated bandhead positions to simulate the laboratory and the spacecraft surface NO glow spectrum. The compilation command on the IBM J40 is "xlf90 synth_new.f". (This command will vary according to different vendors). To run the program after compilation, type "a.out". The calculated bandhead positions are written into the file "gamma_band_position.txt", "beta_band_position.txt", and "delta_band_position.txt" for $\gamma$, $\beta$, and $\delta$ bands, respectfully. The simulated laboratory and spacecraft NO glow spectral intensities and the corresponding wavelengths are written into the files named "lab_simulation.txt" and "space_simulation.txt" in response to user choice. For comparison, the experimental spectral data are also written to this file.

Since the FORTRAN program cannot provide the graphic function, an IDL program "f_rslt.pro" is used to plot the spectrum. The synthetic spectrum was composed as a summation of the triangle pulses centered at each position with the pulse width of half of the input spectral resolution. The spectral magnitude of the model is scaled to the measured spectra from the STS–62 mission. Intensity in Rayleigh/Angstrom is plotted versus wavelength. An example output is shown in figure 10, along with an overlay of the flight spectra which was previously shown in figure 9 as it was compared with the laboratory-generated spectra formed by an O beam flowing onto a plate with NO doping.
Figure 10. Example output of intensity in Rayleigh/Angstrom versus wavelength. Also, an overlay comparison of flight spectra.
4. INFRARED REGION

The data from the infrared telescope (IRT), flown on STS-51-F Spacelab 2 mission, were originally reported by Koch et al.\textsuperscript{29} as originating from near orbital emissions, primarily water (H\textsubscript{2}O). IR measurements were made in the 1.7-3, 4.5-9.5, 6.1-7.1, and 8.5-14.5 \textmu m bands. These observations were made close to perpendicular to the velocity vector and interpreted as being solely due to excited H\textsubscript{2}O vapor. The excitation is caused by solar and Earth shine radiation and by collisions with ambient AO. The outgassing molecules travel with the spacecraft at \approx 7.75 km/sec which, when combined with a thermal velocity of the O atoms at \approx 1,000 K, correspond to a relative translation energy (RTE) between 1.93 and 3.51 eV. The signal in the 1.7- to 3-\textmu m band was observed to change each time the spacecraft passed through the terminator. Since the radiation excitation cross section and solar flux are known, these data provide a measure of the H\textsubscript{2}O column density. This combined data were then used to provide information on the collisional excitation cross sections of the low-lying vibrational levels of H\textsubscript{2}O by AO at an RTE of \approx 3 eV. The IRT data for the 4.5- to 9.5-\textmu m sensor and the nighttime data for the 1.7- to 3-\textmu m sensor are consistent with theoretical collisional excitation of H\textsubscript{2}O by AO. Diurnal variations of 4.5- to 9.5-\textmu m intensities follow the model-predicted O density for a full orbit. Cross sections for the collisional excitation process were derived by Meyerott et al.\textsuperscript{30}

4.1 Theoretical Simulation

Theoretical spectra of H\textsubscript{2}O vapor for the lowest vibrational/rotational \nu\textsubscript{1} (100)\rightarrow(000), \nu\textsubscript{2} (010)\rightarrow(000), and \nu\textsubscript{3} (001)\rightarrow(000) bands have been produced with Briss' algorithm\textsuperscript{30} as part of the investigation of the collisional excitation of the outgassing H\textsubscript{2}O by Orbital O\textsuperscript{(3P)} in the Space Shuttle environment.

4.2 Briss' Algorithm

Briss' algorithm allows the calculation of the line position and the relative line intensity of an asymmetric molecule once the Watson-type hamiltonian\textsuperscript{32} is given. The hamiltonian includes the quartic and sextic centrifugal distortions and takes the following form:

\begin{equation}
H = A_k J_z^2 + A_J J_z^2 + \frac{1}{2} a (J_z^2 + J_z^2) - D_k J_z^4 - D_j J_z^2 J_z^2 - D_j J_z^4 - \frac{1}{2} [d_k J_z^2 + d_j J_z^2, J_z^2 + J_z^2]_+ + H_k J_z^6 + H_{kj} J_z^2 J_z^4 + H_{kj} J_z^4 J_z^2 + H_j J_z^6 + \frac{1}{2} [h_k J_z^4 + h_{kj} J_z^2 J_z^2 + h_j J_z^4, J_z^2 + J_z^2]_+ ,
\end{equation}

where \([A, B]_+\) represents the anticommutator, and \([A, B]_+ = AB + BA\). The coefficients of the hamiltonian are determined by the least-square fit to the high-precision experimental spectra. The hamiltonian is in the base of the symmetry adapted symmetric rotor function,

\begin{equation}
|J, K, \pm, m\rangle = \left\{ |J, K, m\rangle \pm |J, -K, m\rangle \right\} / \sqrt{2}, \quad |J, 0, +, m\rangle = |J, 0, m\rangle .
\end{equation}
The matrix elements in this base are given by Briss. By construction of the matrix of the hamiltonian and conducting the matrix diagonalization, the eigenvalues of the hamiltonian operator can be obtained, hence the energy levels of the molecule. The algorithm further uses the symmetry transformation to obtain the eigenvectors of the hamiltonian operator and uses them to calculate the relative line intensities.

4.3 The Model of a Water Molecule

An H$_2$O molecule is a highly asymmetric top molecule with the moments of inertia (in $10^{-40}$ gm cm$^2$) at equilibrium position $I_a=1.0243$, $I_b=1.9207$, and $I_c=2.9470$. The geometrical parameters of the molecule were determined from an analysis of the rotational structure of IR bands and by microwave spectroscopy:

$$R(O-H)=0.9572\pm0.003 \text{ Å; } \angle \text{HOH} = 104^\circ31'\pm3'. \quad (5)$$

The geographical configuration and the axes of the moments of inertia are shown in figure 11. This geometric configuration falls into the C$_{2v}$ group. The symmetry species of the dipole moment components $\mu_a$, $\mu_b$, $\mu_c$, direction cosines $\lambda_{za}$, $\lambda_{zb}$, $\lambda_{zc}$, normal coordinates $Q_k$ (k=1, 2, and 3) and the rotational wave functions $|J_{K_aK_c}\rangle$ of the H$_2$O molecule in the C$_{2v}$ group are given in table 1.

![Geographic configuration and the axes of the moments of inertia.](image)

Table 1. Symmetry species that determine the selection rules of the transitions.

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<td>$\varepsilon\varepsilon$</td>
<td>A$_2$</td>
</tr>
<tr>
<td>$\lambda_{zb}$</td>
<td>A$_2$</td>
<td>$\varepsilon\varepsilon$</td>
<td>B$_1$</td>
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<td>$\lambda_{zc}$</td>
<td>B$_2$</td>
<td>$\varepsilon\varepsilon$</td>
<td>B$_2$</td>
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The symmetry characteristics solely determine the selection rules for the IR transitions. The allowed transitions are either the $\mu_\text{b}$ type (B bands, between vibrational states of the same symmetry species) or of the $\mu_\text{a}$ type (A bands, between vibrational states of the $A_1$ and $B_2$ species). Allowed rotational transitions in the B-type bands are $ee\leftrightarrow oo$ and $eo\leftrightarrow oe$. In A-type bands, the allowed rotational transitions are $ee\leftrightarrow eo$ and $oo\leftrightarrow oe$.

In the $\text{H}_2\text{O}$ molecule, exchanging of the nuclei of the two hydrogen (H) atoms leads to two different wave functions—the symmetric and antisymmetric. The nuclear spin of the H atom is $I=\frac{1}{2}$. The total nuclear spin $T$ of $\text{H}_2\text{O}$ molecules are different—one for antisymmetric and zero for symmetric. According to the general discussion of the angular momentum, a system with a total spin $T$ has $2T+1$ degenerate state in the situation when there is no magnetic filed. Therefore, the antisymmetric rotational levels occur three times as frequently as the symmetric. As a result, the intensity of the transition is weighed differently—one for symmetric and three for antisymmetric.

### 4.4 Determination of the Relative Population Density

The intensity of a spectral line is give by Briss: \(^{31}\)

\[
I = C(T) g_i e^{-E_i/kT} (1 - e^{-h\nu/kT}) | i'\rangle \mu_i | i \rangle^2
\]

in which $C(T)$ is a constant containing the temperature-dependent partition function and $\nu$ is the frequency of the transition from the upper to the lower state. Factor $g_i$ corresponds to the nuclear statistical weight. Boltzman distribution is assumed whenever the temperature $K$ is input.

The $\text{H}_2\text{O}$ vapor in the Spacelab environment cannot be treated as equilibrium because the excitation is due to the collision with O atoms that are moving toward the Spacelab with a velocity $\approx 7.8$ km/sec; that is, far greater than the speed of the random thermal motion of the $\text{H}_2\text{O}$ molecules. Redmon et al.\(^{33}\) calculated the cross section for vibrational excitation in $\text{O}(^3\text{P}) + \text{H}_2\text{O}(000)$ using the quasi-classical method. They also give the final distribution of the cross sections for the pure rotation excitation $(000) \to (000)$, and the cross sections for the $v_1 (100) \to (000)$, $v_2 (010) \to (000)$, and $v_3 (001) \to (000)$ bands. The theoretical results of the cross section for the $(100)$ band was in reasonable agreement with the experimental shock tube results. The $(010)$ excitation cross section is larger than the corresponding experimental results. Nevertheless, results from Redmon et al.\(^{33}\) give us the rough estimation of the probability of the $\text{H}_2\text{O}$ molecules to be excited to a particular energy level due to the collision with the orbital O atoms. Furthermore, since we do not have enough information to estimate the relative population at different energy levels, we simply use the final rotational distribution given by Redmon et al.\(^{33}\) to calculate the line intensity.

### 4.5 Input Parameters

An important issue in producing $\text{H}_2\text{O}$ emission spectra is to find out the collection coefficients to build up the hamiltonian for different energy levels for $\text{H}_2\text{O}$ molecules. There are 15 coefficients in the hamiltonian operator. The coefficients used were for the $(000)$ and $(010)$ band published by Camy-Peyret and Flaud\(^{34}\) and the coefficients for $(100)$ and $(001)$ bands published by Flaud and Camy-Peyret.\(^{35}\) In their works, the interaction between different states was considered so that the hamiltonian contains
terms higher than the sextic order. To fit Briss’ algorithm, the higher order was simply truncated and coefficients up to the sextic order were used. Since some of the coefficients in the higher order were comparable with that for the sextic order, it unavoidably introduced some inaccuracies.

The coefficients and some parameters are summarized in table 2.

Table 2. Parameters for the hamiltonian for different energy levels: $A_k = A - (B + C)/2$, $A_j = (B + C)/2$, $a = (B - C)/2$.

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<td>1594.74498</td>
<td>3755.92948</td>
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<tr>
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Other parameters:
- Reduction indicator: 1
- Representation indicator: 1
- Nuclear indicator (ee, eo, oo, oe): 1, 3, 1, 3
4.6 Program Operation

This FORTRAN program calculates the theoretical spectra of H₂O vapor for the rotational and the lowest vibrational/rotational (100)–(000), (010)–(000), and (001)–(000) bands using the Briss algorithm. The spectrum of the collisional excitation of the outgassing H₂O by orbital O(3P) is simulated by including the distribution of the cross sections for the pure rotation excitation (000)–(000), and the cross sections for the (100)–(000), (010)–(000), and (001)–(000) bands given by Redmon et al.

This program can also produce the H₂O vibrational/rotational spectrum for the thermal equilibrium, assuming that the Boltzmann distribution stands.

The program read input file to get the coefficients of the hamiltonian is “input0-0, input1-0, input2-0, and input3-0”. It also reads file “phs7.input” for the output control. The output files are OUTPUT0, OUTPUT1, OUTPUT2, OUTPUT3, and SYNT, which contain the spectral intensity for the pure rotational band, the three lowest vibrational/rotational bands, and the composite band, respectfully.

The procedure to compile and execute the program is as follows:

• Move Asyrot.f, interface.f, input0_0, input1-0, input2-0, input3_0, and phs7.input to the same directory
• Compile the source programs using command “f77 *.f”
• Execute the executable file a.out
• Plot the desired spectrum using included IDL program.

4.7 Water Vapor Emission Spectra

Theoretical H₂O vapor spectra are shown in figure 12 for pure rotational band (000)→(000), v₁ (100)→(000) (fig. 13), v₂ (010)→(000) (fig. 14), and v₃ (001)→(000) (fig. 15), respectively. These spectra are the simulations of the emission of H₂O vapor excited by collision with O atoms in the orbital Space Shuttle environments. Each plot has 501 data points with a data interval of 11.9 cm⁻¹. A composite of these theoretical spectra is shown in figure 16.
Figure 12. Spectra of $\text{H}_2\text{O}$ generated by collisional excitation of $\approx 2$ eV energy, believed to be excited by ram atmosphere colliding with $\text{H}_2\text{O}$ which has offgassed from orbiter surfaces but is in the Shuttle atmosphere.

Figure 13. Same as figure 12 except for the $\nu_1$ band of $\text{H}_2\text{O}$. 
Figure 14. Same as figure 12 except for the $v_2$ band of H$_2$O.

Figure 15. Same as figure 12 except for the $v_3$ band of H$_2$O.
Figure 16. Same as figure 12 except a composite spectra of H$_2$O for $v_1-v_3$ and rotational bands shown in figures 12–15.
5. GLOW MITIGATION

5.1 Visible Wavelength Surface (Heterogeneous) Glows

The heterogeneous (surface stimulated) glows often seen on the vertical stabilizer of the STS are caused by surface-bound NO encountering ram O, forming excited NO₂. The nature of the surfaces which enhance NO to adhere in large numbers are (a) the hygroscopic nature of the surface; i.e., irregular surfaces such as Shuttle tile and carbon paints offer a large surface area for NO to stick; and (b) temperature; i.e., colder surfaces keep more NO on the surface. NO is found naturally in the upper atmosphere which easily provides a saturation source for surfaces. Thruster effluent produces large amounts of NO. When effluent is directed upstream to reencounter surfaces, orbiter surfaces become doped by the orbiter flying through the effluent.

The surface brightness is directly related to the rate at which NO₂* forms; i.e., linearly with the O fluence, surface-doped NO density, and cross section.

5.1.1 Surface-Doped NO Density

The element least known about is the surface-doped NO density. Since the glows are related to O fluence with altitude, then the NO (which is variable with altitude also) must be at saturation on a given surface for a given temperature. It must be the surface sites of the material that make the physi-bond and NO unite, which is subsequently broken with a ram O recombination. This would suggest further then that porous surfaces are better than nonporous to hold a lot of NO. There is another contributor in orbit to make NO on surfaces from the recombination of N and O on catalytic surfaces (nickel is used in the laboratory for this purpose). It is the fluence of natural NO from the atmosphere that collides and physi-bonds to surfaces. This is the NO source in low-Earth orbit (LEO); however, a catalytic surface such as nickel (Ni) could add more NO through surface catalytics N and O recombinations. These surfaces would not glow any brighter as the NO saturation attribute is what seems to be important.

The next step in this area of research, to establish validation, is to perform some experiments in the laboratory. Leone and Swenson began doing some of these studies at Lockheed in the early 1990's, but a dedicated effort is needed. What comes to mind is to dope surfaces with NO where the surface temperatures are controlled, then bombard those surfaces with an electron beam or, using laboratory methods, study the dopant density versus temperature and material. Until this is done, the only consideration for doping is the assumption that it is only the material temperature which affects the saturation limit of surface bound NO, and consequently the brightness results.

5.1.2 State Change

Does the cross section change for a state change at very low temperatures, where NO goes into an ice state (121 K)? Observations have been made for T>160 K (estimated), but suggest that on surfaces <121 K, the NO could layer up as ice, and as such, would have a much reduced cross section for glow.
However, this would likely introduce the additional problem in very LEO of NO ice on cryogenic mirrors.

Section 5.1.3 describes some ideas for mitigation. Sections 5.1.3.1 and 5.1.3.5 are the natural, most sensible considerations, based on the observations made to date. Sections 5.1.3.2 and 5.1.3.3 should be followed where possible. Before proceeding with section 5.1.3.4, the laboratory work described above must be addressed.

5.1.3 Mitigation

The following discussion addresses several ideas on how this glow may be eliminated, only from the standpoint of what will eliminate the glow and not from the practicality or necessity of eliminating or reducing it.

5.1.3.1 Glow Brightness. Orbiting at higher altitudes reduces glow linearly with the decrease in O. The O scale height (e folding distance) is $\approx 50$ km, thus 50-km higher altitude will reduce the glow brightness by $e$. In addition, the atmospheric source of NO is reduced. It is not clear from existing measurements that the altitude at which NO no longer saturates the surface continuously is reached, but it is likely between 300–350 km. Above that altitude, glow brightness will be reduced by the product of the decrease in O and atmospheric NO.

5.1.3.2 Gas Releases. The release of inert or nonreactive gases into the upstream (into the velocity vector) interact with the ramming atmospheric O before it reaches orbiter surfaces. This was demonstrated on STS–62, with an N$_2$ release. It was originally suspected that N$_2$ + O could result in a significant production of NO at orbit velocities. This was found not to be the case. N$_2$ is nonreactive with ram O and acts to reduce the O reaching the surface. Consequently, gas releases can reduce glow; although not a practical long-term solution, it certainly is very effective for short periods. A gas density sufficient to chase a ram atom to make a single collision (i.e., a column density of $\approx 10^{15}$ cm$^{-2}$) before ramming the surface is likely a sufficient density to be highly effective.

5.1.3.3 Thruster Effluent. Monomethyl hydrazine is a type of gas release that produces a large amount of NO. On the Shuttle, some of the thrusters are directed so that a large amount of effluent encounters the surfaces. That effluent is stopped by the surface to form a near-stationary cloud above it, which then can be directed by the momentum exchange of the ram atmosphere to carry the effluent into downstream (away from $\gamma$) surfaces. This is frequently seen on the orbiter, where the vernier jets dope the top of the port wing, for example, and the ram atmosphere carries the effluent to become a dopant of bay payloads and Shuttle surfaces. In addition, the NO densities are initially so large in the gas cloud that bright gas phase NO$_2$ emissions occur. Mitigation here is pretty simple. In future designs, the thrusters should be directed so they do not impinge material back onto surfaces in the thrust. The effluent in the burn will then escape quickly, and only the bright gas phase emission will be visible for a brief time. Optically, it is much better to use a nonthrust system altogether, such as the SKYLAB control moment gyro (CMG) system for attitude control where optical instruments are a major player in the payload. In that design, thrusters were used infrequently to dump momentum from the wheels, but not as the primary attitude control system.
5.1.3.4 Orbiter Surfaces. Nonhygroscopic surfaces are better since, at a given temperature, less real surface area per subtended area is available to hold surface sticking molecules, such as NO. Surfaces such as Shuttle tile and black carbon paints have this trait. This trait in the carbon paints has a very low-reflection coefficient and an isotropic scatterer. Consequently in optical baffles, for instance, the trait is needed for these attributes. This is a speculative issue, and the laboratory experiments described above would be recommended before imposing constraints or guidelines for orbiter surfaces exposed near glow sensitive detectors.

5.1.3.5 Temperature Relationships. Warmer temperatures of surfaces result in lesser NO molecules being bound to the surfaces, and thus not available for heterogeneous processes. This is likely a very practical solution for elimination of baffle glows, for instance, where changing the material is not practical. The temperature relationship for a given material follows closely with $I=Io\exp(E/kT)$, where $E=0.14$ eV, $k$ is Boltzman’s constant, and $T$ is temperature in Kelvin.

5.2 Infrared, Extended Glows

5.2.1 Background

IR glows observed on the Shuttle are primarily the result of the ram atmosphere interacting with the H$_2$O vapor outgassing from the Shuttle. Findings suggested that it is the ionospheric O$^+$ (ions) colliding with H$_2$O vapor molecules, which resulted in emissions $=2\mu$, and that atmospheric O bombarding H$_2$O vapor was responsible for a broad spectrum ranging from 1–40 $\mu$.

On STS–62, it was also discovered from the IR spectrometer measurements that NO bands were observed in thruster plumes and afterglows. NO is a burn product of the thruster, and thrusters which directly impinge effluent back to Shuttle surfaces cause NO doping and glows.

5.2.2 Mitigation of H$_2$O Emissions

Section 5.1.3.1 applies slightly in that the O density decreases with altitude, reducing the O fluence to the gas cloud of H$_2$O. The IR brightness will be directly related to the O fluence, which $e$-folds every 50 km. Hygroscopic surfaces such as Shuttle tile hold large amounts of H$_2$O vapor (described in sec. 5.1.3.2). The main mitigation is to keep the orbiter materials dry before launch. The large emissions observed from early Shuttle missions before the Scotchguard™–type treatments have seemingly been very effective. This has been effective in accumulating H$_2$O before getting into orbit. Also, once in orbit, exposing surfaces to a hot soak will drive the surface-accumulated species off the surfaces more quickly; i.e., as described in sec. 5.1.3.5).

5.2.3 Mitigation of NO Emissions

Thruster-induced NO emissions described in section 5.1.3.3 are believed to be responsible for most NO emissions. Gas phase N produced in burns can readily combine with ram O to form excited NO. Consequently, the burn product emissions cannot be avoided, although the rate of emission is related to the altitude (and the ram O density).
Surface conductors can also lead to a recombination glow in NO, similar to that formed by N and O flows over conductive surfaces such as Ni in laboratory-produced glows and afterglows. Nonconductive surfaces will significantly reduce (or eliminate) this process.

5.3 Ultraviolet and Far-Ultraviolet Glows

5.3.1 Background

Ultraviolet glows in the 200–350 nm region are a result of NO emissions and FUV glows in the 140–200 nm region from N\textsubscript{2} LBH bands. These emissions are primarily sporadic and dominated by thruster dopants. There appear to be UV glows in the 300-nm region, leaving many surface-related astronomy experiments misunderstood.

5.3.2 Mitigation of NO Emissions

5.3.2.1 Thruster Effluent. The dominant glows observed on the Shuttle by the EISG experiment were NO bands in the 200–300 nm spectral region. Nearly all the emission observed was NO in this region (secs. 5.1.3.3 and 5.2). The same mitigation ideas hold for the FUV NO band emissions as did the IR NO band emissions described in section 5.2.

5.3.3 Mitigation of N\textsubscript{2} LBH Bands

The observation of the N\textsubscript{2} LBH bands on STS–62 could not be repeated. Although it was observed in a laboratory experiment prior to flight, the instrument sensitivity may have been too low in the 130–150 nm regime. The LBH bands seen by the Infrared Space Observatory (ISO) experiment on STS–9 were due to surface recombination of N, where N was produced by thruster burns whose effluent came into the Shuttle bay by thruster effluent impinging on Shuttle surfaces and by ram atmosphere upstream from the orbiter momentum exchanging with the effluent. This exchange allows it to slow sufficiently until the Shuttle velocity overtakes the gas. The only mitigation is to eliminate the N sources; i.e., the thruster.

Glows occurring at wavelengths <140 nm have not been addressed here.
REFERENCES


## REPORT DOCUMENTATION PAGE

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<td>Spacecraft glow information has been gathered from a number of spacecraft including Atmospheric and Dynamic satellites, and Space Shuttles (numerous flights) with dedicated pallet flow observations on STS–39 (DOD) and STS–62 (NASA). In addition, a larger number of laboratory experiments with low energy oxygen beam studies have made important contributions to glow understanding. The following report provides information on three engineering models developed for spacecraft glow including the far ultraviolet to ultraviolet (1400–4000 Å), and infrared (0.9–40 μ) spectral regions. The models include effects resulting from atmospheric density/altitude, spacecraft temperature, spacecraft material, and ram angle. Glow brightness would be predicted as a function of distance from surfaces for all wavelengths.</td>
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NSN 7540-01-280-5500

Standard Form 298 (Rev. 2-89)
Prescribed by ANO Std. 238-19
298-102