1-1-1977

Rocketborne Measurement of Mesospheric H2O in the Auroral Zone

James W. Rogers
A. T. Stair Jr.

Follow this and additional works at: https://digitalcommons.usu.edu/sdl_pubs

Recommended Citation
https://digitalcommons.usu.edu/sdl_pubs/113
ROCKETBORNE MEASUREMENT OF MESOSPHERIC H₂O IN THE AURORAL ZONE

James W. Rogers and A.T. Stair, Jr.

Air Force Geophysics Laboratory, Bedford, Massachusetts 01731

Thomas C. Degges

Visidyne, Inc., Burlington, Massachusetts 01803

Clair L. Wyatt and Doran J. Baker

Electro-Dynamics Laboratories, Utah State University
Logan, Utah 84321

Abstract. Infrared emission spectra of the long-wavelength wing of the 6.3 μm water band between 6.7 and 7.6 μm have been measured in the mesosphere using a cryogenic rocketborne spectrometer. The resulting zenith radiances have been compared with a high-altitude radiance model resulting in a volume mixing ratio of 3.5 ± 2.2 ppm between 49 and 70 km.

A liquid-helium-cooled, long-wavelength infrared (LWIR) spectrometer was launched by the Air Force Geophysics Laboratory onboard a Black Brant rocket on 22 March 1973 from the University of Alaska’s Poker Flat Research Range at Chatanika, Alaska. The spectrometer, which employed a circular-variable filter (CVF), was developed by AFGL and Utah State University (USU) and provided the first measurements of the altitude profile of the infrared spectrum of the upper atmospheric emissions between 7 and 24 μm.

The dominant emission features measured were from the 15 μm carbon dioxide (ν2) band between 65 to 150 km and from the 9.6 μm ozone (ν2) band between 45 and 100 km initial results of which were reported by Stair et al., [1974]. Final analysis of the rocket flight has been completed by Rogers et al., [1976] revealing emission data on the long-wavelength wing of the 6.3 μm water band between 6.7 and 7.6 μm.

The spectrometer has been described in detail elsewhere [Wyatt, 1975] and so only a brief summary of its salient features will be included here. The spectral region from 6.7 to 24 μm is scanned at a rate of two scans per second with a full-angle field of view of 2 degrees. The entire optical subsection is cooled to liquid-helium temperature, including an arsenic-doped silicon detector which is thereby operated with an effective zero radiation background. A key element of the spectrometer is the Optical Coating Laboratory, Inc., (OCLI) rotating circular variable interference filter. The spectral resolution of the filter was measured at AFGL by Condron [1973] and the best estimate of the half-bandwidth for the flight-configured spectrometer was 2.63 percent for the short wavelength filter half and 2.92 percent for the long wavelength filter half. A dynamic range of 3x10⁴ was achieved by providing four data channels with different gains. The spectrometer is the least sensitive at the shorter wavelengths with a noise equivalent spectral radiance (NESR) of 2x10⁻¹⁰ watt cm⁻²sr⁻¹μm⁻¹ at 7μm.

The payload was launched during a post-breakup auroral glow which had persisted at the IBC II level for nearly 10 minutes. Vertical viewing data were obtained during ascent from 96 km to spogee at 185 km while descent data were obtained down to 45 km. The overhead photon radiance at 3914Å as measured by an onboard photometer was about 5 KR below 100 km.

Figure 1 shows a typical emission spectrum obtained at 54 km for the short wavelength half of the rotating filter in the spectrometer. The dominant feature at 9.6 μm is due to the ν2 band of ozone. The emission between 6.7 and 7.6 μm has been attributed to the long wavelength wing of the ν2 fundamental band of H₂O. The altitude dependence of these two spectral features from 75 to 50 km is shown in Figure 2 by a three-dimensional presentation of the zenith spectral radiance versus wavelength as a function of decreasing altitude. The 9.6 μm ozone emission has been intentionally drawn off scale in the figure to show the weaker H₂O emission at the shorter wavelengths.

The altitude dependence of the zenith radiance obtained by integrating the spectral data from 6.7 to 7.6 μm is shown by the open boxes in Figure 3. The dashed curves are the ± 3σ error bars associated with the spectrometer data. The larger spread in the error bars at the higher altitudes indicates that the spectrometer’s lower sensitivity level is being reached.

The solid curve in Figure 3 is a theoretical calculation for the water radiance profile in this wavelength region using the model of Degges [1974]. In this calculation 20% of the total 6.3 μm water band emission lies between 6.7 and 7.6 μm. This estimate was obtained by calculating the spectral bandshape using the line positions and intensities of McClatchey et al., [1973] and integrating to obtain the percentage of resulting emission in the wavelength region of interest. The effects of temperature on this percentage are small since the result varied by only one percent over the range from 200 to 2500K. The primary excitation processes for H₂O at these
Fig. 1 Spectral radiance values between 6.7 and 12.4 \( \mu \text{m} \) during descent at 54 km.

Altitudes are the absorption of earthshine or radiation emitted lower in the stratosphere,

\[
h\nu + \text{H}_2\text{O} (000) \rightarrow \text{H}_2\text{O} (010)
\]

vibrational energy exchange with molecular oxygen through the near resonant collision mechanism

\[
\text{H}_2\text{O} (010) + \text{O}_2 (\nu = 0) + \text{H}_2\text{O} (000) + \text{O}_2 (\nu = 1)
\]

and interconversion of thermal and vibrational energy by collision with other molecules,

\[
\text{H}_2\text{O} (010) + \text{M} \rightarrow \text{H}_2\text{O} (000) + \text{M}
\]

De-excitation of vibrationally excited \( \text{H}_2\text{O} \) is by spontaneous radiation which occurs with an Einstein A coefficient of \( 20.3 \text{ s}^{-1} \) (McClatchey et al., [1973].) The rate coefficients used for collisional de-excitation are those given by R.L. Taylor [1975], \( 2.70 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \) for exchange with \( \text{O}_2 \) and \( 4.8 \times 10^{-11} \exp(-43/T^{1/3}) \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \) for quenching by \( \text{N}_2 \) and \( \text{CO}_2 \). Excitation rates are obtained from de-excitation rates by applying detailed balancing. Excitation by exchange of vibrational energy with \( \text{O}_2 \) is the strongest collisional process and is more effective than absorption of radiation at all altitudes below 75 km. Collisional de-excitation is faster than spontaneous emission at all altitudes below 85 km.

Accurate modeling of the \( \text{H}_2\text{O} \) emission requires detailed knowledge of the altitude profiles of the atmospheric temperature, density \([M]\) and water concentration \([\text{H}_2\text{O}]\). The calculation shown in Figure 3 used a standard arctic atmosphere with 3.5 ppm by volume of \( \text{H}_2\text{O} \) and a temperature profile obtained by an extrapolation of sounding rocket temperature measurements made during the night of the spectrometer flight. These temperatures were 216, 222 and 236K at 30, 40 and 50 km respectively (Atmospheric Sciences Laboratory, White Sands Missile Range, New Mexico, private communications). The mesospheric temperatures obtained by this extrapolation introduce a factor of two uncertainties in the theoretical radiance profile over a 30° temperature range.

The water concentration used was determined by at least square fit of the calculated to the measured radiance values. The resulting average \( \text{H}_2\text{O} \) concentration is \( 3.5 \pm 1 \text{ ppm} \) by volume between 39 and 70 km. The error bars are increased to \( 3.5 \pm 2.2 \) by the additional error of a factor of two introduced by the temperature uncertainty.

The mixing ratio of \( \text{H}_2\text{O} \) has been measured extensively in the stratosphere. Mastenbrook [1968], [1971] has performed a comprehensive stratospheric humidity sounding program carried out with an automatic frost-point sensor and has reported volume
Acknowledgments. The authors wish to thank the personnel from various organizations who made significant contributions to the success of this experiment including: Dr. H.C. Fitz, Jr. and Mr. Herb Mitchell (presently with R&D Associates) of the Defense Nuclear Agency (DNA); Ned Wheeler, Tom Condron and Ed McKenna of AFGL and Gary Frosham of Utah State University. This experiment was sponsored by DNA under Subtask L25AAKHK3C, Work Unit 20 as part of ICECAP (Infrared Chemistry Experiments—Coordinated Auroral Program).

References


Stair, A. T., Jr., J. C. Ulwick, D. J. Baker, C. L. Wyatt and K. D. Baker, Altitude Profiles of Infrared Radiance of O3 (8.6 μm) and CO2 (15 μm), Geophysical Research Letters, 1, 117-118, 1974


(Received July 12, 1977; accepted July 22, 1977.)