No. 103 THE INFRARED SPECTRUM OF JUPITER, 0.95—1.60 MICRONS, WITH LABORATORY CALIBRATIONS

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ABSTRACT

New spectra of Jupiter in the region 0.95—1.60 microns are presented with calibrations from laboratory studies of the bands of CH₄ and NH₃. Estimates of the quantities of these gases above the effective reflecting layer in the Jupiter atmosphere are given for each of several individual bands. The presence of CS₂, H,S, and HCN is investigated with the aid of composite spectra of these gases in addition to CH₄ and NH₃. We find as upper limits 8 cm-atm for CS₂, 50 cm-atm for H,S, and 10 cm-atm for HCN.

1. Introduction

On January 5, 1967, we obtained infrared spectra of Jupiter in the range 0.95—1.60 μ, having higher resolution than those previously published by Kuiper (1964) and Moroz (1966). Because the higher resolution increases the opportunity to search for minor constituents in the Jupiter atmosphere, and to study the absorption characteristics of different bands, these new spectra are presented with laboratory spectra of different amounts of methane, ammonia, acetylene, hydrogen sulfide, and hydrogen cyanide, at a resolution commensurate with the Jupiter spectra. The spectra of Jupiter given here have a resolution λ/Δλ ≈ 700 at 1.5 μ, while the apparent resolution of Moroz’ tracings is 250 and that of Kuiper’s 450.

2. The Spectrum of Jupiter

Three tracings of the spectrum of Jupiter were obtained with the infrared spectrometer described by Kuiper, et al. (1962) and the 61-inch NASA reflector of the Lunar and Planetary Laboratory. A lead sulfide detector of width 1.0 mm was used with an entrance slit of 1.0 mm (subtended angle 10 seconds of arc). The slit was aligned roughly parallel to the equator and included the equatorial zone and parts of the north and south equatorial belt. The spectrum was scanned at 1.72 Å/sec with a time constant of 10 sec. The three tracings were averaged and a hand-drawn composite prepared. The composite and the two best individual spectra are presented in Figures 1 through 3.
Fig. 1  Infrared spectrum of Jupiter, Part 1, 0.93–1.20 μ. Curve a is a composite of tracings b and c and a third, not reproduced. Band identifications are taken from laboratory spectra. The spectrum up to 1.12 μ can be observed with higher resolution by photographic spectroscopy.

Fig. 2  Spectrum of Jupiter, Part 2, 1.12–1.40 μ; as Fig. 1.
3. Laboratory Comparisons

In 1964, Kuiper published a series of laboratory spectra of different amounts of CH$_4$ and NH$_3$ with a resolution comparable to his spectrum of Jupiter. We have similarly prepared a series of these spectra with slightly higher resolution to correspond more closely to the spectra of Jupiter in Figures 1–3. A single pass absorption tube of length 40 meters with quartz windows and a quartz-iodine lamp was used. To increase the amount of gas in the path, it was necessary to increase the pressure; the effects of band growth shown in Figures 4–9 are not pressure independent. Figures 4–6 show tracings of different amounts of CH$_4$ within the wavelength interval 0.93–1.72 μ. The best overall fit to the Jupiter spectrum corresponds to approximately 56 m-atm CH$_4$ at 1.4 atm pressure. With this quantity of methane in the absorption tube, we added increasing amounts of NH$_3$ so that the gases were mixed inside the tube.

The spectra so obtained are shown in Figures 7–9. The composite spectrum of NH$_3$ and CH$_4$ is more easily compared to the spectrum of Jupiter than are spectra of the individual gases alone. Using these laboratory data we have attempted to identify the prominent features in the Jupiter spectrum, the results of which are indicated above the composite tracing in Figures 1–3. Several uncertainties remain in the identification and such features are indicated with a question mark.

Using the laboratory observations of CH$_4$ and NH$_3$, we have estimated the approximate equivalent absorbing pathlength above the scattering/reflecting layer in the Jupiter atmosphere for several different absorption bands. The estimates are given in Table 1. Because the laboratory observations were made with a single fixed-pathlength absorption cell, the observed band-development with increasing amounts of gas is not pressure independent. Further, our observations
were all made at room temperature (22°C) so that the appearance of the bands and the adjacent continuum may differ somewhat from those in the spectrum of Jupiter. The effects of temperature and/or pressure are especially evident in the NH₃ band at 1.255 µ wherein the two minima are reversed in relative intensity in the laboratory spectra, and in the CH₄ band at 1.31 µ, which is much weaker in the Jupiter spectrum than in the laboratory runs. The estimates in Table 1 are correct in the first approximation, but to higher accuracy must be understood as relative only to our laboratory data. There may be errors by as much as a factor of two, especially for the strong bands which, in the case of Jupiter, may form high in the planet's atmosphere where the pressure is less by a factor of 3 or more than that in the laboratory.

The wide range of values for CH₄ and NH₃ in Table 1 attest to the often neglected fact that a reliable estimate of the quantity of either of these gases cannot be made from the study of a single band. As Kuiper (1952, 1964) and others have pointed out, one must recall that strong bands reach absorption at much higher levels in the atmosphere than do weak ones.

It can be seen from Table 1 and the figures illustrating the ammonia bands that NH₃ is an important absorbing molecule in the Jupiter atmosphere. Danielson (1966 p.955) reached the opposite conclusion because his Stratoscope II spectra showed an intensity maximum instead of a depression at 1.53±0.02 µ, very near the 1.51 µ band of NH₃. This is purely an effect of the low resolution of the Stratoscope II spectra. It can be seen in Figure 3 that the "window"
TABLE 1
Prominent bands of CH₄ and NH₃ identified in the spectrum of Jupiter with estimates of the equivalent amount of gas giving the same absorption in laboratory conditions.

<table>
<thead>
<tr>
<th>MOLECULE</th>
<th>WAVELENGTH (µm)</th>
<th>EQUIVALENT AMOUNT ABOVE JUPITER CLOUD LAYER</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH₃</td>
<td>1.08</td>
<td>5 m-atm</td>
</tr>
<tr>
<td>CH₄</td>
<td>1.10</td>
<td>100</td>
</tr>
<tr>
<td>NH₃</td>
<td>1.22</td>
<td>0.8</td>
</tr>
<tr>
<td>CH₄</td>
<td>1.23</td>
<td>100</td>
</tr>
<tr>
<td>NH₃</td>
<td>1.26</td>
<td>2</td>
</tr>
<tr>
<td>CH₄</td>
<td>1.29</td>
<td>0.8</td>
</tr>
<tr>
<td>CH₄</td>
<td>1.31–1.32</td>
<td>25</td>
</tr>
<tr>
<td>CH₄</td>
<td>1.33</td>
<td>13</td>
</tr>
<tr>
<td>CH₄</td>
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<td>10–20</td>
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<tr>
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<td>20–60</td>
</tr>
<tr>
<td>NH₃</td>
<td>1.51</td>
<td>0.2</td>
</tr>
<tr>
<td>NH₃</td>
<td>1.53</td>
<td>0.2</td>
</tr>
<tr>
<td>NH₃</td>
<td>1.57</td>
<td>4</td>
</tr>
</tbody>
</table>

in the Jupiter atmosphere from 1.40–1.63 µm is strongly asymmetrical with the intensity peak at 1.59±0.01 µm. Danielson’s spectra smear out this window to a nearly symmetrical envelope with the peak shifted to shorter wavelengths. The asymmetry shown in our spectra is also well displayed in the records of Kuiper (1964) and Moroz (1966). The importance of telluric water vapor in shaping this envelope is small compared to that of CH₄ and NH₃. The H₂O band closest to the region of interest is the relatively narrow absorption (at this resolution) at 1.47 µm. Comparison of a spectrum of the moon made when the telluric water vapor concentration was similar to that when the Jupiter records were made (for example, Kuiper, 1964, Figure 11) shows that the intensity at the deepest part of the band is about 70 percent of the intensity at 1.59 µm. In the Jupiter spectrum in Figure 3, the intensity at 1.47 µm is 25 percent of the peak at 1.59µm. This illustrates that water vapor absorption in this region does not account for the discrepancy between the spectra of Danielson and those of Kuiper, Moroz, and the present authors.

Danielson (1966, p.955 and note added in proof) correctly concluded that the broad, deep absorption
at 1.37 \mu is caused by CH₄. This is evident in Figures 5 and 6 where the growth of the band superimposed on the small water vapor absorption in the laboratory (corresponding to 18 microns precipitable H₂O), is well shown. These spectra are offered in lieu of those of Owen and Cruikshank (unpublished) mentioned by Danielson (1966, p.959).

Mention must be made also of Danielson's discussion of the absorptions at 1.15 \mu. He indicates that if a significant fraction of the absorption there is due to the 2-0 band of H₂, the depth of the bands corresponds to approximately 45 km-atm H₂ for a temperature of 200–225°K. By comparing Figures 2 and 4, it can be seen that the three individual minima at about 1.135 \mu (broad), 1.155 \mu, and 1.165 \mu correspond in detail to features of CH₄.

From these room temperature comparisons, the 1.135 \mu band is most closely matched by 36.8 m-atm CH₄ at 0.93 atm pressure, while the latter two bands agree well with 17.4 m-atm CH₄ at 0.434 atm pressure. Contributions from the relatively weak telluric H₂O band at 1.13 \mu are insignificant. If then, as it seems, much of the absorption in the 1.15 \mu region is due to CH₄, Danielson's estimate of the amount of H₂ in the absorbing path is too great, making the abundance in this band more nearly in accord with that of about 14 km-atm estimated from the H₂ absorptions near 2.25 \mu (see Appendix).

4. Acetylene, Hydrogen Sulfide, and Hydrogen Cyanide

The presence of minor constituents on Jupiter is important to the chemical equilibrium of the atmosphere. In order to make an accurate test for the presence of H₂S, HCN, and C₂H₂, we observed these gases against the background of absorptions of CH₄ and NH₃. These three gases were considered possible contributors to the array of features in the Jupiter spectrum partly on the basis of comparisons with the laboratory spectra of Cruikshank (1967). Recognition that composite spectra are superior to single spectra for the detection of individual lines or line
Fig. 7  Spectra of CH₄ with increasing amounts of NH₃ added. Part 1, 0.93–1.20 μ. Absorbing path length is 40 m for all spectra. Curve a is a blank run with no gas in absorption tube, but with 3 m air path in spectrometer (about 18 microns precipitable H₂O). All spectra in Figures 4–11 have same spectrometer air path. Curves b-i have 55.6 m-atm CH₄ at 1.42 atm partial pressure. Curve b has no NH₃; remaining spectra have NH₃ added in the following amounts with the indicated total pressure: 0.105 m-atm NH₃, 1.424 atm; d, 0.211 m-atm NH₃, 1.426 atm; e, 0.422 m-atm NH₃, 1.432 atm; f, 0.899 m-atm NH₃, 1.443 atm; g, 1.847 m-atm NH₃, 1.47 atm; h, 3.69 m-atm NH₃ at 1.51 atm; i, 7.91 m-atm NH₃, 1.62 atm.

Fig. 8  Spectra of CH₄ with increasing amounts of NH₃ added, Part 2, 1.10–1.40 μ; as Fig. 7.
blends supported the use of the following laboratory procedure.

The single-pass 40-meter absorption tube of the Lunar and Planetary Laboratory was filled with 55.6 m-atm CH₄ plus 0.211 m-atm NH₃ at a total pressure of 1.4 atm. Because of the small amounts of C₂H₂ required, it was not possible to introduce the gas directly into the tube with CH₄ and NH₃. Thus, a 20-cm glass absorption tube with plane glass windows containing the C₂H₂ was varied independently for the different amounts shown in Figure 10. There are no identifiable lines of C₂H₂ in the Jupiter spectrum, but in the region of 1.51–1.55 μ where C₂H₂ absorbs, the Jupiter profile is concave. In the laboratory runs with CH₄ and NH₃ alone, this region is convex. The absorption by about 5 cm-atm C₂H₂ changes the profile to match more nearly that of Jupiter. It must be understood, however, that effects of temperature and/or pressure discussed above may account for the concave CH₄-NH₃ profile in Jupiter, and the identification of C₂H₂ must be regarded only as a possibility. Owen (1965) used the 10372 Å band of C₂H₂ to establish an upper limit of 3 m-atm of this gas in the Jupiter atmosphere. Using the stronger infrared absorptions, in particular the 101 band at 1.538 μ considered above, we can establish an upper limit of 8 cm-atm. While 5 cm-atm would affect the shape of the profile, as noted above, 8 cm-atm should result in recognizable absorption features at the resolution of the laboratory and Jupiter data. A positive identification is not possible from these unresolved spectra; other gases absorb in this region, for example, HCN which is discussed below. Also this region lies on the shoulder of a strong band of NH₃, and as demonstrated above, different parts of the fine structure in a given band correspond to different amounts of absorbing gas. Failure of the laboratory spectra to match the curve of growth produced in planet's atmosphere can easily effect small changes in the slopes and valleys of a band.
Fig. 10 Spectra of CH₄, NH₃, and C₂H₆ combined, 1.40–1.63 μ. All spectra have 55.6 m-atm CH₄ plus 0.211 m-atm NH₃ with path length of 40 m, total pressure 1.4 atm. Spectra b-e have absorptions of C₂H₆ superimposed. The C₂H₆ gas was contained in an absorption tube in tandem with the 40 m tube, and was used in the following amounts, always at 700 mm pressure: b, 2.3 cm-atm; c, 4.6 cm-atm; d, 9.3 cm-atm; e, 18.6 cm-atm.
Fig. 11 Spectra of CH₄, NH₃, and H₂S combined, 1.40–1.63 μ. All spectra were made with 40 m absorption tube and have 3 m laboratory air path in spectrometer superimposed (about 18 microns precipitable H₂O). All spectra have 55.6 m-atm CH₄ plus 0.211 m-atm NH₃. Spectrum a has only CH₄ plus NH₃, total pressure 1.40 atm; b, CH₄ plus NH₃ plus 0.211 m-atm H₂S, total pressure 1.403 atm; c, same as b, but with 0.632 m-atm H₂S, total pressure 1.415 atm. Small segment of H₂S spectrum inserted has only 0.65 m-atm H₂S at pressure 0.93 atm, shown at higher resolution.
Hydrogen cyanide (HCN) was observed in three different amounts for comparison with the spectrum of Jupiter. Figure 12 shows the 002 band at 1.533 μ; there is an indication of incipient rotational structure in the P branch, just beyond the spectral resolution limit. Using the HCN cells in tandem with the 40 m tube containing CH₄ and NH₃, the composite spectra in Figure 13 were made with the resolution adjusted to compare with that of the Jupiter spectra in Figures 1–3. The P and R branches of HCN are easily distinguished from the background of methane and ammonia absorptions even for 20 cm-atm of the gas. We estimate that 10 cm-atm in the double-transmission path in the Jupiter atmosphere would be visible in the Jupiter spectrum in Figure 3. This represents an improvement of a factor of 20 over the upper limit determined by Owen (1965).

A similar search was made for H₂S. This is a weakly absorbing gas, thus requiring large amounts. H₂S was therefore added directly to the tube containing 55.6 m-atm CH₄ and 0.211 m-atm NH₃. The resulting spectra are given in Figure 11 with a high resolution spectrum of H₂S alone (62 cm path at pressure 705 mm Hg) added to show the band shape and partly resolved component lines. As in the case of C₂H₂, no bands could be positively identified but the laboratory spectral profile in the region 1.58–1.61 μ was modified to match more closely that of the Jupiter spectra. From the spectra in Figure 11 we estimate that 0.3 m-atm H₂S could account for the shape of the Jupiter spectrum in this region, subject to the limitations of pressure and temperature noted above for acetylene. We suggest an upper limit of 0.5 m-atm for the presence of H₂S in the Jupiter spectrum (double-pass).

APPENDIX

In his discussion of the Jupiter spectrum in the region 2.0–2.5 μ, Danielson (1966 p. 956) noted that methane absorptions alone are insufficient to account for the broad absorption between about 2.0 and 2.5 μ. His statement was based on the appearance of the relatively weak absorption shown in the laboratory spectra of small amounts of CH₄ (Kuiper and Cruikshank, 1964). Danielson concluded that there might be absorption in this region in addition to CH₄ and NH₃, and considered the collision-induced dipole transitions in H₂.

More recent laboratory studies of large amounts of CH₄ have shown that the appearance of the 2.0–2.5 μ region is considerably different from what one might expect using the laboratory spectra of Kuiper.

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Fig. 12 Spectra of the 002 band of HCN, all at 1 atm pressure, (a) 20 cm path, (b) 38 cm path, (c) 1 m path.
and Cruikshank alone. In Figure 12 we give a family of curves showing the increase in effective band width of the CH$_4$ absorptions as the amount of gas and pressure are increased to values more appropriate for Jupiter’s atmosphere. The shortward progression of the very strong absorption terminates at about 2.09 $\mu$ for 120 m-atm CH$_4$. The massive absorption in this region is a blend of the 0110 (2.20 $\mu$), 0011 (2.32 $\mu$), and 1001 (2.37 $\mu$) bands. With this amount of CH$_4$, the 0012 (1.79 $\mu$) band also absorbs to about 2.04 $\mu$. As seen in Table 1, this quantity of methane is not inconsistent with some of the bands in the Jupiter spectrum. The spectra in Figure 14 do not include the effects of the strong NH$_3$ bands at 2.22 $\mu$ and 1.98 $\mu$, which would further alter the profile of the remaining window transmitted by the Jupiter atmosphere.

It is therefore inappropriate to draw conclusions on the abundance of hydrogen from absorptions in this spectral region where methane and ammonia

![Figure 13](image-url)  

*Fig. 13* Spectra of CH$_4$, NH$_3$, and HCN combined, 1.40–1.60 $\mu$. All spectra have 55.2 m-atm CH$_4$ plus 0.20 m-atm NH$_3$ mixing in 40 m tube, at total pressure 1.39 atm. HCN contained in glass tubes external to 40 m tube. (a) no HCN, (b) 20 cm HCN at 1 atm pressure, (c) 38 cm HCN at 1 atm pressure, (d) 1 m HCN at 1 atm pressure.
Fig. 14 Spectra of CH₄, 1.90–2.22 μ, medium resolution. All spectra were made with 40 m absorption tube and have 3 m laboratory air path in spectrometer superimposed (about 18 microns precipitable H₂O). (a) blank run, tube evacuated, (b) 1.68 m-atm CH₄ at 0.042 atm pressure, (c) 3.37 m-atm at 0.084 atm, (d) 13.7 m-atm at 0.34 atm, (e) 26.3 m-atm at 0.66 atm, (f) 52.6 m-atm at 1.32 atm, (g) 120 m-atm at 3 atm.
bands are so strong and complexly blended, unless the resolution is much higher than that of Stratoscope II spectra or our own observations with the scanning spectrometer.

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