DIRECT SPECTROSCOPIC EVIDENCE FOR IONIZED POLYCYCLIC AROMATIC HYDROCARBONS IN THE INTERSTELLAR MEDIUM

G. C. Sloan,¹,² T. L. Hayward,³ L. J. Allamandola,² J. D. Bregman,² B. DeVito,³ and D. M. Hudgins²

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ABSTRACT

Long-slit 8–13 μm spectroscopy of the nebula around NGC 1333 SVS 3 reveals spatial variations in the strength and shape of emission features that are probably produced by polycyclic aromatic hydrocarbons (PAHs). Close to SVS 3, the 11.2 μm feature develops an excess at ~10.8–11.0 μm, and a feature appears at ~10 μm. These features disappear with increasing distance from the central source, and they show striking similarities to recent laboratory data of PAH cations, providing the first identification of emission features arising specifically from ionized PAHs in the interstellar medium.

Subject headings: dust, extinction—infrared; ISM: lines and bands—ISM: molecules

1. INTRODUCTION

Gillett, Forrest, & Merrill (1973), in a spectroscopic study of planetary nebulae, discovered a series of unidentified infrared (UIR) bands, now known to include features at 3.3, 6.2, 7.7, 8.6, and 12.7 μm. Since then, the UIR bands have been observed in a rich variety of astronomical sources (see Allamandola 1996 for a recent review). Polycyclic aromatic hydrocarbons (PAHs) were first introduced as a possible carrier of the UIR bands by Leger & Puget (1984) and Allamandola, Tielens, & Barker (1985). While other carriers have been proposed for the UIR bands, none have proved as successful as the PAH model in explaining the observed spectral details. Nonetheless, many differences exist between earlier laboratory and astronomical spectra, including the relative strength of the features and the wavelengths at which they appear, and these differences have prevented the PAH model from gaining wider adherence.

A major discrepancy has been between the relative intensities of the features in the 10–13 μm region (C—Hout-of-plane bending modes) and the features in the 6–9 μm region (C—C modes and the C—Hplane bend at 8.6 μm). In laboratory spectra of neutral PAHs, the C—H modes are much weaker than the features at longer wavelengths, but in astronomical spectra, the C—H modes are stronger. Allamandola et al. (1985) originally suggested that most or all PAHs in the interstellar medium would be ionized because of their low-ionization potential (~6 eV), and laboratory investigations of PAH cations show better agreement with the observed UIR spectra in terms of both the relative feature strengths and positions (Hudgins, Sandford, & Allamandola 1994; Hudgins & Allamandola 1995a, 1995b, 1997, 1998; Szczepanski & Vala 1993a, 1993b; Szczepanski, Chapo, & Vala 1993; Szczepanski et al. 1995a; Szczepanski, Wehlburg, & Vala 1995b).

Here we make a detailed comparison between recently available laboratory data and spatial and spectral variations in the infrared spectrum from the SVS 3 region in the reflection nebula NGC 1333. SVS 3 is an early B star (Strom, Vrba, & Strom 1976; Harvey, Wilking, & Joy 1984), producing a much milder UV spectrum than found in other PAH emission regions (e.g., the Orion Bar, NGC 7027). Joblin et al. (1996) obtained discrete mid-infrared spectra in three locations in the reflection nebula using a 5″ beam and found that the strength of the 8.6 and 11.2 μm PAH features varied inversely to each other, with the 8.6 μm feature emitting more strongly near SVS 3 and the 11.2 μm feature emitting more strongly to the south, away from SVS 3. Since the 8.6 μm feature shows enhanced strength in the spectra of most PAH cations (along with the other bands between 6 and 10 μm), Joblin et al. concluded that SVS 3 was ionizing a larger fraction of the PAHs closer in than farther away. This dependence of the ionization fraction as a function of distance from the ionizing source could easily result from geometric dilution.

This investigation concentrates on emission features in the 10–13 μm spectral region that arise from out-of-plane bends of C—H bonds on the periphery of PAH molecules (see, e.g., Bellamy 1958, p. 65, and Allamandola, Tielens, & Barker 1989). For neutral PAHs, the strongest feature at 11.2 μm originates in PAH rings that contain only one C—H bond (the solo mode). The duo mode (two adjacent C—H bonds) produces a feature in the vicinity of 11.9 μm, but this feature is much weaker in astronomical sources. Since, in the laboratory spectra of PAHs, the wavelength of these bands can shift a substantial fraction of a micron, depending on the size and structure of the molecule, the positions of the solo and duo modes have long been used by chemists as a diagnostic of PAH structure. The trio mode produces a feature at ~13 μm, but the wavelength range of this mode overlaps with the quartet and quintet modes at longer wavelengths, making unambiguous identification difficult.

2. OBSERVATIONS AND ANALYSIS

In order to investigate the spectral variations in the PAH emission at higher spatial resolution, we obtained long-slit 8–13 μm spectra of NGC 1333 SVS 3 at the 5 m Hale telescope at Palomar on the nights of 1996 September 29–30 (UT) using SpectroCam-10 (Hayward et al. 1993). The data have a spectral resolution of 0.19 μm and a diffraction-limited angular resolution of ~0.5″. The slit was oriented north-south and covered a 2° × 16′ region of the sky, including SVS 3 and the nebulosity to the south. We used standard chop-and-nod sequences

¹ School of Physics, University College, Australian Defence Force Academy, Canberra ACT 2600, Australia.
² NASA Ames Research Center, MS 245-6, Moffett Field, CA 94035-1000.
³ Center for Radiophysics and Space Research, 612 Space Sciences Building, Cornell University, Ithaca, NY 14853.

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Fig. 1.—The average spectrum from the PAH emission region south of SVS 3, determined from 2" away from SVS 3 to the south end of the spectrometer slit. The error bars represent 1σ uncertainties.

with 40" and 60" amplitudes east-west in order to correct for background emission from the telescope and sky. The data were flux-calibrated using spectra of β Peg taken immediately before or after the NGC 1333 observations, together with archival SpectroCam-10 ratio spectra of β Peg versus α Lyr and the absolute α Lyr model from Cohen et al. (1992). The data from the two nights were combined into a single two-dimensional spectral image from which individual one-dimensional spectra were extracted for plotting.

Figure 1 illustrates the spectrum summed from 2" south of SVS 3 to the end of the slit, showing the 8.6 μm feature on the shoulder of the stronger 7.7 μm feature, the 11.2 μm feature, and the emission plateau extending to the weaker 12.7 μm feature. Figure 2 shows how the strengths of the PAH features vary with position along the slit. While the 11.2 and 12.7 μm features grow progressively stronger toward the PAH emission ridge 10" south of SVS 3, the 8.6 μm feature is stronger closer in. This behavior confirms the results of Joblin et al. (1996).

Figure 3 shows how the shape of the 11.2 μm feature (solo mode) changes as a function of distance from SVS 3. Close to the source, the feature has an excess on the short-wavelength wing, which appears to consist of multiple components. As the distance from SVS 3 increases, the shorter wavelength portion of the wing (centered at 10.8 μm) disappears first, followed by the longer wavelength portion (at 11.0 μm).

We have extracted the flux from these two components (Fig. 4), which we describe as the “blue outliers” to the 11.2 μm feature, by averaging the profile of the 11.2 μm feature 8" south of SVS 3 (and beyond), normalizing this mean profile to each row, and subtracting it. The 10.8 μm outlier is summed from 10.6 to 10.9 μm, and the 11.0 μm outlier is summed from 10.9 to 11.2 μm. While the 10.8 μm outlier goes to zero 8" from SVS 3, the 11.0 μm outlier still makes a contribution, which we crudely estimate to be $0.91 \times 10^{-16}$ W m$^{-2}$ arcsec$^{-2}$ at this position by fitting a Gaussian to the blue edge of the main band at 11.2 μm. Both outliers increase in strength by a factor of 2 close to SVS 3, despite the fact that the stronger features at 8.6, 11.2, and 12.7 μm are at a minimum in this region (Fig. 2).

Our observations of NGC 1333 suggest that the 12.7 μm feature also develops a blue wing close to the central source. Because of the poor signal-to-noise ratio in this spectral region, the deep atmospheric absorption features (due to water vapor at 12.38, 12.44, 12.52, and 12.56 μm and CO$_2$ at 12.63 μm), and the complications introduced by possible [Ne II] emission

Fig. 2.—Equivalent fluxes from the PAH features at 8.6 μm (top), 11.2 μm (middle), and 12.7 μm (bottom), determined by fitting and subtracting a linear baseline from beneath each feature. The 11.2 μm spatial profile (dotted line) has been normalized and plotted (top and bottom) for comparison.

Fig. 3.—Normalized spectral profiles of the 11.2 μm feature after removing a linear baseline, color-coded by position. The lengths of the vertical bars above the profiles illustrate the average 1σ uncertainty in the data for each spectral strip. These strips are 1" wide.
3. DISCUSSION

A component in the vicinity of 11.0 \( \mu \text{m} \) has appeared before in the spectra of several PAH sources, most strongly in TY CrA (at 11.05 \( \mu \text{m} \); Roche, Aitken, & Smith 1991), but also in Elias 1 (at 11.06 \( \mu \text{m} \); Hanner, Brooke, & Tokunaga 1994), and more weakly in several other sources, including Elias 14 (at 10.8 \( \mu \text{m} \); Hanner, Brooke, & Tokunaga 1995) and WL 16 (DeVito & Hayward 1998). The WL 16 data (also obtained with SpectroCam-10 at Palomar) show spatial behavior similar to our NGC 1333 data, with the blue wing on the 11.2 \( \mu \text{m} \) feature becoming more pronounced closer to the central source and fading farther away (Fig. 3 in DeVito & Hayward 1998).

To determine the nature of the blue outliers and the 10 \( \mu \text{m} \) feature, we compare our astronomical data with the database of spectra from the Astrochemistry Laboratory Group at NASA Ames Research Center (Hudgins et al. 1994; Hudgins & Allamandola 1995a, 1995b, 1997, 1998) and the theoretical models by Langhoff (1996).

For neutral PAHs, Langhoff (1996) finds that the position of the 11.2 \( \mu \text{m} \) feature depends on the geometry of the molecule. Moving from a small molecule like anthracene (three adjacent rings) to tetracene (four rings) to pentacene (five rings), the feature shifts from 11.3 to 10.9 \( \mu \text{m} \). This shift raises the possibility that the blue outliers might result from a change in the composition of the PAH mixture. However, the laboratory data of Hudgins et al. show a much smaller shift in wavelength as a function of molecular size. From anthracene to pentacene, the feature only shifts from 11.3 to 11.1 \( \mu \text{m} \).

The relative strengths of the 11.2 and 12.7 \( \mu \text{m} \) bands provide a crude means of probing the size of the PAHs. In larger PAHs, the solo mode (11.2 \( \mu \text{m} \)) would dominate, since any ring along a straight edge of the molecule would have only one C—H bond, while the trio mode (12.7 \( \mu \text{m} \)) would appear more frequently in smaller PAHs, where a larger fraction of the rings occupy corners and not straight edges. As Figure 2 shows, the spatial behaviors of both the 11.2 and 12.7 \( \mu \text{m} \) features agree with each other within the uncertainties, pointing to closely related sets of carries and not variations in the size of the PAHs. Consequently, we do not consider it likely that the blue outliers to the 11.2 \( \mu \text{m} \) feature result from changes in the size distribution or molecular geometry of the PAHs sampled by the slit.

Figures 6 and 7 compare the laboratory and theoretical spectra of neutral PAHs and PAH cations. In the cations, most of the C—H out-of-plane bending modes have shifted 0.4 \( \mu \text{m} \) to shorter wavelengths (Fig. 6), providing a straightforward interpretation of the blue outliers seen close to SVS 3. Figure 7 shows that the 10 \( \mu \text{m} \) feature seen near SVS 3 may also arise from PAH cations, since PAH cations consistently produce features in this wavelength region while neutral PAHs do not.

Joblin et al. (1996) argued that the fraction of PAH cations decreases further from SVS 3 because of decreasing fluxes of ionizing photons. In our data, the blue outliers to the 11.2 \( \mu \text{m} \) feature and the 10 \( \mu \text{m} \) feature show just this spatial dependence. The combination of this spatial behavior and the appearance of similar spectral features in laboratory spectra of ionized PAHs leads us to identify these features with PAH cations.

This identification substantially strengthens the case for...
PAHs as carriers of the UIR bands. The identification of the 3.29 μm band with the aromatic C—H stretch and the bands in the 11–13 μm region with out-of-plane C—H bends requires that the carrier consist of aromatic hydrocarbons, but the nature of these aromatic hydrocarbon molecules has remained in doubt (see, e.g., Sellgren 1994; Tokunaga 1997; Uchida, Sellgren, & Werner 1998). Energy requirements discussed in the proposal of PAHs as possible carriers of the UIR bands (Leger & Puget 1984; Allamandola et al. 1985) lead to estimates that these molecules must contain 40–80 carbon atoms. Therefore, they must be polycyclic. But these molecules might exist within a larger matrix of hydrocarbons and other molecules, commonly described as hydrogenated amorphous carbon (HAC; see the recent review by Duley 1993). In order for the molecules to be ionized, they must be free molecules; i.e., they must be separate from any HAC-like matrix. In the face of these combined arguments, individual gas-phase PAHs must be the dominant emitters of the narrow UIR bands.

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