ULTRAVIOLET SPECTROSCOPY OF HD 44179

MICHAEL L. SITKO
Department of Physics, University of Cincinnati, Cincinnati, OH 45221 and Space Science Institute, Boulder, CO 80301; sitkoml@ucmail.uc.edu, sitko@spacescience.org

LAWRENCE S. BERNSTEIN
Spectral Sciences, Inc., Burlington, MA 01080; larry@spectral.com

AND

ROBERT J. GLINSKI
Department of Chemistry, Tennessee Technical University, Cookeville, TN 38505; rglinski@tntech.edu

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ABSTRACT

We have reanalyzed the ultraviolet spectrum of HD 44179, the central star(s) of the Red Rectangle Nebula, providing improved estimates of the column density, rotational, and vibrational temperatures of the 4th Positive $A-X$ system of CO in absorption. The flux shortward of 2200 Å is a complex blend of CO features with no discernible stellar photosphere, making the identification of other molecular species difficult, and the direct derivation of the dust extinction curve impossible. We confirm that the spin-forbidden CO($a-X$) Cameron bands are likely produced by either collisional excitation or a chemical reaction, not photoexcitation, but with a higher internal vibrational excitation than previously determined. We also detect the spin-forbidden CO $a-X$, $d-X$, and $e-X$ absorption features. The hot CO($A-X$) bands exhibit a blueshift of $\sim 300$ km s$^{-1}$, likely occurring close to the white dwarf star(s) suspected as the original source of the ultraviolet flux in the system, and forming the base of the outflow of material in the Red Rectangle. The OH "comet-band" system near 3000 Å is also analyzed, and estimates of its rovibrational temperatures determined. The source of the molecules studied in this system is still unknown, but may be a combination of gaseous material associated with the star(s), or processed material from the surrounding dust torus.

Subject headings: circumstellar matter — molecular processes — stars: AGB and post-AGB — stars: individual (HD 44179) — ultraviolet: stars

1. INTRODUCTION

HD 44179 is a binary star system that is deeply embedded within a dusty torus at the heart of the Red Rectangle Nebula. In addition to the ubiquitous organic infrared emission bands (generally thought, but not yet proven, to be polycyclic aromatic hydrocarbons [PAHs]), the nebula possesses a broad emission band centered near 6400 Å and a series of narrower bands of molecular origin throughout much of the visible part of the spectrum (Cohen et al. 1975; Schmidt et al. 1980). Despite a quarter century of investigation, the origin of both the broad and narrow features remain unidentified.

The star system itself is a single-line spectroscopic binary with a highly eccentric orbit ($e \sim 0.38$) with an orbital period of $\sim 300$ days, and the masses of both components are subsolar (van Winckel et al. 1995; Waelkens et al. 1996; Waters et al. 1998; Men’shchikov et al. 2002). The visible star seems to be a post-asymptotic giant branch (post-AGB) spectral class F star with an ultraviolet (UV) excess. The nature of the secondary star is unknown. Waelkens et al. (1996) suggested that it was a low-mass main-sequence star, while Men’shchikov et al. (2002) favored a high-luminosity white dwarf. This latter model is based largely on the presence of an ultraviolet (UV) excess above that of the post-AGB primary, which was attributed to a white dwarf secondary. In fact, we will show that the UV flux cannot be entirely photospheric.

The UV spectrum of HD 44179 contains key information about the excitation of the Red Rectangle Nebula and the molecular chemistry required to produce the species responsible for its complex spectrum at longer wavelengths. Since their original discovery by Cohen et al. (1975) and Schmidt et al. (1980) both the broad red emission “hump,” now referred to as the “extended red emission” (ERE), and the narrower emission bands at visible wavelengths have defied any positive identification. In a study of NGC 7023, another nebula with significant ERE emission, Witt et al. (2006) concluded that the carrier of the ERE is made from the ionization of precursor species with photons having energies greater than 10.5 eV ($\lambda < 1180$ Å). Thus, information about molecular species bathed by ultraviolet photons in the vicinity of HD 44179 is central to understanding the chemical development of the carrier of the ERE and other emission features.

The first clues to the nature of molecular interaction with the UV flux occurred with the demonstration that the UV spectrum of HD 44179 is dominated by molecular emission and absorption by CO (Sitko 1983). All of the identified bands were due to electronic transitions from the $A^1\Pi - X^1\Sigma^+$ 4th Positive (4P) system of this molecule. Absorption features at wavelength shortward of 1600 Å from the vibrational states $(\nu', \nu'') = (0, 0)$ to at least $(8, 0)$ were visible (where $\nu'$ and $\nu''$ refer to the vibrational levels of the upper and lower electronic states, respectively). The entire emission band system seen at wavelengths longer than 1600 Å was then due to downward fluorescent emission from the upper $(\nu)$ electronic state to the ground $(X)$ electronic state. Based on the strengths of the $(0, 0)$ and $(1, 0)$ absorption bands, a column density $N_{CO} \gtrsim 10^{18}$ cm$^{-2}$ was derived. A similar amount of CO$_2$ was also suggested at the time, since CO$_2$ was known to be highly absorbing shortward of 1600 Å. Because the energy bandwidth and total estimated dereddened flux absorbed in this region were comparable to those in the ERE, it was speculated that the former might even be responsible for powering the latter.
Observations of HD 44179 with greater sensitivity and spectral resolution using the Hubble Space Telescope have provided more detailed information about the UV flux in this object. Glinski et al. (1996) identified the spin-forbidden $^3\Pi-^1\Sigma^+$ Cameron emission bands in between 1900 and 2200 Å, the first such detection in an astronomical object outside the solar system. The excitation mechanism for the Cameron bands remains unknown. Glinski et al. (1996) suggested collisions by charged particles with energies of 7–15 eV. Yan et al. (2000) favored dissociative recombination of electrons with HCO$^+$, leaving CO in an excited $^3\Pi$ state.

Glinski et al. (1997) found that fitting the 4P fluorescent (0, 1), (0, 2), and (0, 3) emission bands required the presence of at least 2 components (and a best fit with 4), one being cold with a rotation temperature of only $\sim$50 K, and another much hotter one at $\sim$3000 K. The spectropolarimetry studied by Reese & Sitko (1996) indicated that these emission bands also dilute the polarization, similar to what Schmidt et al. (1980) found for the ERE at visible wavelengths. Reese & Sitko also drew attention to the possible presence of OH emission bands near 3080 Å.

Glinski et al. (1996) and Reese & Sitko (1996) had identified other features between 2500 and 2800 Å as possibly arising from Fe II, but offered no model to describe the emission. The presence of eleven Fe II emission lines arising from three separate multiplets at visible wavelengths (Hobbs et al. 2004) supports the reality of these identifications.

Because of its importance to understanding the chemical evolution of the nebula and to better understand the detailed physics of the ultraviolet spectrum, we recently undertook an effort to reanalyze the UV spectrum of HD 44179. We discuss our results in the following sections.

2. OBSERVATIONS

The high-resolution observations of the (0–1), (0, 2), and (0, 3) $A-X$ emission bands of CO were obtained on 1993 March 20 (UT) using the G160M grating and the 2.0′′×2.0′′ Large Science Aperture of the Goddard High Resolution Spectrograph (GHRs) of the Hubble Space Telescope (HST). Details of the observations are described in Glinski et al. (1997).

All low-resolution spectra were obtained using the G130H, G190H, and G270H gratings of the Faint Object Spectrograph (FOS) of the HST, with the 1.0′′ circular aperture centered on the star. The G190H and G270H observations were obtained on 1992 November 11 (UT) in the spectropolarimetric mode, and total intensities reconstructed as described by Reese & Sitko (1996) and Glinski et al. (1996). Under ideal conditions, with calibration lamp exposures obtained immediately before or after the science exposure, and without grating movement, the wavelength calibration of the FOS was generally good to a fraction of a diode. At the longer G270H wavelengths, with a spectral resolution of 2 Å diode$^{-1}$, the calibration would normally be good to $\sim$1 Å. However, the insertion of the polarization optics introduces additional uncertainty. Usually, the spectra in the two orthogonal planes of polarization had slightly different wavelength calibrations and were simply shifted before merging (R. A. Allen 2007, private communication), degrading the absolute wavelength calibration further. We consider the absolute calibration to be no better than 2 Å, but the relative shift within a limited spectral region with the same grating to be more precise. These limitations will determine whether or not any conclusions can be drawn regarding the relative velocities of different emitting species.

The G130H observations were a series of seven identical observations on 1992 September 20 (UT) without the insertion of the polarimeter optics, and averaged with equal weighting.

However, the archived G130H spectra were severely mangled during processing and archiving, and appear to suffer from a poor background correction. In particular, the flux levels shortward of 1550 Å are inconsistent with previous spectra obtained using the International Ultraviolet Explorer (IUE, Sitko 1983), as well as LiF2A spectra at $\lambda \sim 1080$–1180 Å in the data archive of the Far Ultraviolet Spectroscopic Explorer (FUSE), which place an upper limit on the flux at 1180 Å of $F_\lambda \sim 5 \times 10^{-15}$ erg cm$^{-2}$ s$^{-1}$ Å$^{-1}$. In Figure 1 we compare the “preview” and final archived G130H spectra with that obtained using the IUE. Also shown are GHRS observations from the archive obtained a few years later with the G140L grating. When smoothed to the same approximate spectral resolution as the FOS data, the overall flux levels and general spectral shapes are very similar, but the core line depths deeper. In order to use a uniform data set throughout the entire 1300–3000 Å region for much of the modeling, and to keep the differences in time of observation small, we use the FOS spectra for the entire 1300–3300 Å region, except for the set of three CO bands covered by the GHRS spectra from 1993.

3. MODEL DESCRIPTION

3.1. Radiative Transfer Model

The approach used for computing the UV transmission and emission from a gas in non–local thermodynamic equilibrium (NLTE) is an extension of a previously developed radiative-transfer (RT) model (Sundberg et al. 1993) for NLTE IR emission from high-altitude reentry wake flows. It employs a single-line Voigt equivalent width approximation with Curtis-Godav resolution averaging to treat multisegment inhomogeneous paths. The path transmittance and radiance is computed separately for each line and a statistical line overlap correction is applied. The computational spectral resolution is limited by the largest single line equivalent width for the spectrum under consideration. For this study, we typically performed calculations at $\sim$1 cm$^{-1}$ resolution corresponding to $R = \lambda/\Delta\lambda > 3 \times 10^4$. This is higher resolution than the observations, and the model predictions were degraded to lower resolution, accounting for both velocity dispersion and sensor spectral resolution. The accuracy of this approximate RT approach is generally within ±10% of the more accurate, but several orders of magnitude slower, monochromatic line-by-line
method. Given the simplifying assumptions we used in the spectral analysis of the highly spatially complex RR, this level of accuracy for the RT model is appropriate.

The primary extensions of the IR model for UV applications are the generalizations of the NLTE source function and line strength. UV transitions involve upper and lower electronic states each with its own vibration-rotation energy level manifold. The total energies of each upper and lower transition level are factored into their separate electronic, vibrational, and rotational contributions. Populations are required for each of these six components. It is convenient to characterize the population for each degree of freedom in terms of an effective temperature. This often works well; however, an arbitrary vibrational level population distribution can be specified if required. In addition to the six effective temperatures for the internal degrees of freedom, a translational temperature is specified if required. In addition to the six effective temperatures for the internal degrees of freedom, a translational temperature is also specified, which governs the Doppler line width. In this study, the effect of velocity dispersion was introduced in terms of an effective translational temperature.

3.2. Line Parameters

Energy levels for the CO $^1\Pi$ state were determined from a deperturbed expression (Kurucz 1976). Although many of the $^1\Pi$ state energy levels are perturbed by nearby triplet states, these perturbations are not important at the spectral resolution of this work. Dunham coefficients were used for the energy levels of the $X^1\Sigma$ ground state (Guelachvili et al. 1983). The line strengths were based on Franck-Condon factors, $r$-centroids, and HönL-London factors taken from Kurucz’s compilation. The relative electronic transition moments were obtained from the DeLeon (1988) quadratic representation of the $r$-centroid. The absolute transition moments were tied to measurements of the radiative lifetimes (Field 1983). Transitions for $\nu' \leq 23$, $\nu'' \leq 37$, and $j \leq 130$ were included in the line strength database.

The CO Cameron line parameters used in the spectral fitting are based on those previously used in the analysis of laboratory flame (Burke et al. 1996) and space shuttle plume (Dimpfl et al. 2005) observations of CO Cameron emission. The line parameters correspond to vibrational transitions between the $0 \leq \nu' \leq 7$ and $0 \leq \nu'' \leq 9$ levels and for rotational levels up to $j',j'' \leq 100$. The Fe $\Pi$ line parameters were taken from the NIST atomic line compilation. A simplified emission model was adopted for the Fe $\Pi$ lines, principally as a means of demonstrating the close correspondence of data features to the known Fe $\Pi$ lines. The Fe $\Pi$ emission was modeled as optically thin and characterized by an effective emission temperature. While it is understood that neither of these assumptions holds true for the Red Rectangle, they suffice to unambiguously confirm the presence of strong Fe $\Pi$ emission.

The OH energy levels, transition frequencies, and HönL-London factors were determined based on the Dieke and Crosswhite formulas (Dieke & Crosswhite 1962). Corrections to the line strengths due to vibration-rotation interactions were based on the analysis by Anketell & Learner (1967). Franck-Condon factors calculated by D. L. Albritton (1989, private communication) were used and verified against previous calculations by others (Goldman & Gillis 1981). Absolute normalization of the line strengths were based on a well-established experimental value for the $(0,0)$ oscillator strength (Rouse & Engleman 1973). All bands with $\nu' \leq 4$ and $\nu'' \leq 2$ were included, and the maximum value of the rotational quantum number for each $\nu'$ level was constrained to be below the OH($A$) state dissociation limit of 2 eV.

3.3. Spectral Fits

The general approach adopted for spectral fitting of the GHRS and FOS data is briefly reviewed, with more details presented below for the specific data features. In all cases, a spectral baseline contribution, as indicated in Figures 2 and 3, was first subtracted. The binary stars are obscured from direct view by a physically thick and optically opaque dust disk, and the baseline contribution

See the NIST Chemical Kinetics Database, http://kinetics.nist.gov/kinetics/.
corresponds to scattered starlight that has not passed through the strong molecular absorption regions associated with the conical outflows. The baseline-subtracted FOS flux in the 1300–800 Å region, which is dominated by CO(A−X) absorption, was modeled as the product of an effective source function and a computed transmittance, taken as the product of single, homogeneous segments of optically thick hot and cold transmittance components. The primary parameters determined for this spectral region are the source function, the column densities of CO, and the effective vibration and rotation temperatures for the OH(A−X) ground-state components. The baseline-subtracted FOS and GHRS data in the 1600–2200 Å region, which is dominated by CO(A−X) emission, were modeled as a linear combination of optically thin, single, homogeneous segments of hot and cold emission components. However, significant optical opacity is required to account for the attenuation in one of the cold emission components. This emission arises from the reemission to longer wavelengths of the absorbed photons below 1600 Å (fluorescence). The primary parameters determined for the CO emission region include the effective vibration and rotation temperatures for the CO(A) state of the hot and cold components, and a very substantial wavelength-dependent extinction attributed to dust associated with the RR. Because the extinction curve cannot be determined directly (due to the heavy blanketing of molecular features), its properties must be estimated in order to derive self-consistent results for the lines. In addition, the higher resolution of the GHRS data enables the determination of velocity shifts for the hot and cold components. The baseline-subtracted FOS data in the 2900–3300 Å region, which is dominated by OH(A−X) emission, were modeled as a linear combination of optically thin, single, homogeneous segments of hot and cold emission components. The primary parameters retrieved for the OH emission include the effective vibration and rotation temperatures for the OH(A) state of the hot and cold components. As found for CO, optical opacity is also required to account for the attenuation of one of the cold emission features. The model parameters used for the spectral fits are summarized in Table 1. The fits to the individual spectral components will be described below.

### Table 1

**Summary of Temperature and Velocity Fit Parameters**

| Component | \( T_v \) (K) | \( T_r \) (K) | Velocity (km s\(^{-1}\)) |
|-----------|----------------|----------------|-------------------------|
| CO(A−X) GHRS Emission 1580–1735 Å | \( 2000 \) | \( 5500 \) | +294 \(^a\) |
| Very hot emission (0, 1), (0, 2), (0, 3) | \( 2000 \) | \( 50 \) | 0 \(^b\) |
| Cold emission (0, 1), (0, 2), (0, 3) | \( 2600 \) | \( 50 \) | 0 \(^b\) |
| Cold absorption (0, 1) | \( 500–1000 \) | \( 50 \) | 0 \(^b\) |
| CO(A−X) FOS Emission 1600–2400 Å | \( 2000 \) | \( 500 \) | 0 \(^d\) |
| Hot absorption | \( 1000 \) | \( 50 \) | 0 \(^d\) |
| Cold absorption | \( 1000 \) | \( 50 \) | 0 \(^d\) |
| CO(1−X) FOS Emission | \( 1000 \) | \( 50 \) | 0 \(^d\) |
| Chemical | \( \leq 250 \) | \( \leq 250 \) | \( \leq 250 \) |
| OH(A−X) Emission 2550–3300 Å | \( \leq 250 \) | \( \leq 250 \) | \( \leq 250 \) |
| Very hot emission | \( \leq 250 \) | \( \leq 250 \) | \( \leq 250 \) |
| Cool emission | \( \leq 250 \) | \( \leq 250 \) | \( \leq 250 \) |

\(^a\) The only well-determined shift from data.
\(^b\) Shift below resolution of the data.
\(^c\) Not a fit, just estimated.
\(^d\) Constrained by emission fit.
\(^e\) Constrained by model fit to UV photodissociation of H\(_2\)O observed for the space shuttle.
\(^f\) Constrained by estimate for cold absorption.
\(^g\) Constrained by fit to UV photodissociation of H\(_2\)O observed for the space shuttle.
Figure 4 show the limiting extremes of no attenuation and complete attenuation of the (0, 1) cold feature. The data display a small residual (0, 1) cold feature, corresponding to ~90% attenuation of the optically thin limit. We estimate that a $T_v$ in the 500–1000 K range and a column density of cold CO of order $\sim10^{16}$ cm$^{-2}$ is consistent with the data.

The modeled hot component is blue shifted by 294 ± 20 km s$^{-1}$ (–1.6 ± 0.1 Å) at 1600 Å, while no shift was required for the cold components. In order to fit the data, we had to further reduce the effective spectral resolution of the model calculations by applying a triangular slit function of width 0.32 Å ($\sim12.5$ cm$^{-1}$) attributed to a velocity dispersion of 28 km s$^{-1}$. This is somewhat higher than the upper limit of 15 km s$^{-1}$ observed in molecular lines at visible wavelengths (Hobbs et al. 2004), suggesting we are sampling material closer to the central illuminating source.

As an example of the spectral fitting process, the individual spectral components associated with the (0, 3) fit are depicted in Figure 5. We used the same baselines as that estimated from the FOS data as shown in Figure 3. Because of the very high effective vibration temperature for the hot component, there are overlapping contributions from a number of ($v',v''$) bands, not just the (0, 3). As a result, the hot component displays a complex and somewhat irregular line structure. While the cold component gives rise to the most distinct feature, the integrated intensity is dominated by the hot component. The general shape of the hot component is principally determined by the CO(4) rotational temperature, while the underlying continuum (i.e., not the baseline) is largely determined by the CO(3) vibrational temperature. It is also seen that the relatively sharp band edges for the hot bands allows for a good determination of the observed blue shift. This implies that if emission of comparable intensity from a corresponding redshifted component were present, it would be clearly discernible in the data. A possible origin of the one-sided velocity shift seen only for the hot emission component is discussed later.

Inspection of Figure 4 shows that the integrated intensity of the (0, 3) spectral region is approximately a factor of 1.5 times higher than that for the (0, 1) region. The relative integrated intensities of the CO(4–X) emission in these spectral bands are primarily controlled by the (0, 1) and (0, 3) Franck-Condon factors, since both regions arise from the same upper electronic state vibrational level. This would predict an intensity ratio of (0, 3) to (0, 1) of 0.7, as opposed to the observed 1.5, a factor of 2 discrepancy! We attribute this strongly wavelength-dependent effect to several potential causes, including, strong dust extinction in the “hidden” region in the general vicinity of the binaries, and the spectral albedo of the scatters in the conical outflows. This can be expressed as a reddening function that is applied to the CO(4–X) spectral predictions,

$$f_{\text{red}} = \exp[0.006(\lambda - 1600)],$$

where, for convenience in the spectral fitting, we normalized the function to unity at 1600 Å (this technique is analogous to using the Balmer decrement in ionized gas clouds to derive the extinction characteristics of the dust there). Physically, this means that the dust opacity increases by a factor of 3.6 in going from 2200 to 1600 Å, corresponding to the full spectral range of the CO(4–X) emission. This translates into a factor of 36 change in intensity. This large effect will be borne out subsequently when the FOS CO(4–X) emission fit is presented, which encompasses the full range of the emission spectrum.

3.5. FOS CO(4–X) Fits 1300–2400 Å

The fits to the FOS in the CO(4–X) absorption, 1300–1600 Å, and emission, 1600–2400 Å, spectral regions are depicted in Figure 6. The FOS emission region was modeled with the same parameters and spectral components as determined from fitting the GHRS data. The GHRS derived reddening function was also
used for the FOS fit. The dramatic influence of the reddening on the entire emission spectrum is evident in Figure 7. Without reddening, the long-wavelength tail of the CO emission spectrum would not be easily discerned beyond \( \sim 2000 \) Å.

For fitting the absorption region, 1300–1600 Å, we also used a two-component, hot and cold, model. The velocity shifts were constrained by those found from the GHRS emission analysis. However, it is noted that the effect of the velocity shift on absorption is opposite to that for emission; that is, the absorption is redshifted when the emission is blueshifted. The spectral fit in the absorption region was divided into two spectral pieces because of the strong discontinuity of the apparent source and baseline contributions at \( \sim 1450 \) Å (see Fig. 1). This sudden drop in flux is consistent with spectral models of the AGB component of the

Fig. 6.—Comparison of model fits to the FOS data in the CO(\( A-X \)) absorption (top) and emission (bottom) regions. Baseline-subtracted FOS data were used for the absorption region comparisons. The same constant CO column density of \( 2.7 \times 10^{17} \text{ cm}^{-2} \) was used for the entire absorption spectral region, 1300–1560 Å. The retrieved source function for each of the absorption regions is also displayed. Only CO(\( A-X \)) absorption and emission features are included in the fits. The (\( v',0 \)) band locations of CO absorption transitions originating from other electronic states are indicated in the middle panel with arrows. The lowest \( v' \) band for each sequence is labeled.

Red Rectangle (Men’shchikov et al. 2002). The CO(\( X \)) ground electronic state temperatures for the hot component were found to be \( T_{v'} = T_{v''} = 2000 \pm 500 \) K. The retrieved CO column density was \( 3 \pm 1 \times 10^{17} \text{ cm}^{-2} \). The cold component parameter values are not well determined from these low-resolution data, as the cold component only accounts for a small sliver of the spectrum near the strong band edges. However, it is required in order to fill in missing intensity near the band edge due to the redshifting of the hot component. Based on the GHRS fit for the cold component, we used \( T_{v'} = 1000 \) K and \( T_{v''} = 50 \) K. A column density comparable to that for the hot component was required in order to account for the missing level of absorption (see middle plot of Fig. 6). The \( ^{13}\text{CO}/^{12}\text{CO} \) isotope ratio was estimated to be 0.05, but this is not a well-determined parameter; it was slightly preferred over 0.02 and 0.10. A velocity dispersion corresponding to a Doppler temperature of \( T_D = 2 \times 10^3 \text{ K} (\sim 18 \text{ km s}^{-1} \text{ dispersion}) \) was required to obtain a sufficient level of absorption for the highly saturated CO features. The actual velocity dispersion may well be higher, but at this value the individual CO lines are highly overlapped, and further line broadening has little impact on the RT calculations. For simplicity, a constant source function was assumed and adjusted for a best fit. The source function is certainly more complex, as it is comprised of both the true stellar source function as well as reemitted light from the initially absorbed photons (see Fig. 7 for an estimate of the reemission spectrum in the absorption spectral region). Most of the absorption in the 1440–1600 Å is accounted for by the CO(\( A-X \)) bands, and the remaining absorption correlates well to other known bands of CO, in this case the spin-forbidden \( a'-X, d-X, \) and \( e-X \) bands, which are not included in the model, but whose wavelengths are indicated in Figure 6. Their presence at significant strength is confirmed by laboratory measurements of CO absorption (Rostas et al. 2000), which are compared to the FOS data in Figure 8. The laboratory measurements were performed at a much higher pressure, 50 torr, lower temperature, 298 K, and much higher column density, \( 2.4 \times 10^{19} \text{ cm}^{-2} \), than exist in the Red Rectangle. As a consequence, they correspond to a higher degree of opacity and saturation.

The spectral fit in the 1300–1450 Å region, using the same parameters as for the 1440–1600 Å, except for a lower baseline, is seen in Figure 6 to account for most of the observed absorption down to \( \sim 1380 \) Å where it begins to significantly underestimate
the level of absorption. However, while incomplete in this spectral region, the Rostas et al. (2000) data strongly suggest that the most, if not all, of the absorption is likely attributable to CO.

3.6. FOS CO(\(A\rightarrow X\)) and Fe II Fits 1800–2800 Å

The CO(\(a\)) and Fe II model fits to the FOS data for the CO Cameron emission region are shown in Figure 9. For the CO(\(a\)) fit, two emission mechanisms were considered, direct photoexcitation of CO by the strong UV flux in the vicinity of the binary system, and chemical excitation which likely occurs in a different spatial region. As discussed below, only the chemical mechanism was found to provide a reasonable fit and is the only one displayed in Figure 9. For the chemical mechanism, the retrieved rotational temperature for the CO(\(a\)) emission was \(T_\text{rot} = 1000 \pm 200\) K. A well-defined vibrational temperature could not be determined. Rather, an approximately equal population for all \(v', v''\) levels yielded the best fit. The fit shown in Figure 9 corresponds to equal populations for all levels. While some improvement to the fit could be obtained by allowing for individual adjustments to the populations, this is not warranted, given the as yet many uncharacterized absorption and emission features due to other species in this spectral region. For instance, we note that the strong model-predicted \((2, 0)\) emission is not evident in the data, presumably due to the overlapping strong C II absorption feature. For the photoexcitation mechanism, equal vibrational populations were also assumed, and a somewhat higher rotational temperature of \(T_\text{rot} = 3000\) K was used. The reddening effect found for the CO(\(A\)) emission was included for the photoexcitation mechanism but not for the chemical mechanism, for reasons discussed below. There is a good correlation of spectral features and relative intensities for the chemical mechanism. The photoexcitation mechanism may account for some of the small residual intensity above \(\sim 2550\) Å not accounted for by the CO(\(a\)) chemical and Fe II emission channels. However this is quite speculative and requires more careful modeling, particularly for the Fe II lines, to corroborate.

The spectral fit to the CO(\(a\)) emission for the chemical mechanism indicates a very high degree of vibrational excitation of the \(v'\) levels. The distribution determined here is considerably more excited than that derived previously by more approximate analysis (Glinski et al. 1996; Yan et al. 2000). The current, much hotter distribution seems more consistent with collisional excitation by an ion, as discussed by Yan et al. However, the chemical mechanism cannot be ruled out, as significant differences in the local environment in which the chemistry occurs (i.e., Red Rectangle vs. laboratory) may effect the observed product CO(\(a\)) vibration and rotation state distributions. The fit for the chemical mechanism did not require the reddening effect that was required for fitting the CO(\(A\)) emission. This suggests that the chemical excitation of CO(\(a\)) occurs in a spatial region that is further away from the binaries than that associated with the direct photoexcitation that produces CO(\(A\)) and possibly some CO(\(a\)).

The modeled Fe II emission lines demonstrate a close correspondence with emission or absorption features seen in the data. The modeled spectrum was based on an effective emission temperature of 12,000 K. However, no great significance should be attributed to this temperature; it was merely used to adjust the relative emission intensities of the various Fe II line groups to fall within the range of the data. It is noted that several of the stronger modeled Fe II emission lines, such as those near 2350 and 2400 Å, correlate with absorption features in the data. This indicates that these lines are optically opaque. Photons absorbed in these lines escape the Red Rectangle via optically thinner transitions at longer wavelengths that originate from the same upper states but terminate on different and higher lying lower states.

3.7. FOS OH(A→X) Fits 2750–3350 Å

The fit to the FOS OH(A→X) emission spectrum is displayed in Figure 10 and the fit components are shown in Figure 11. Because of the noisy appearance of the baseline-subtracted data, spectral fitting was performed on spectrally smoothed data corresponding to a spectral resolution of \(\sim 10\) Å. The spectrum was modeled as the sum of two components, a very hot component due to UV photodissociation of H₂O, and a cold component due to stellar-induced fluorescence of OH. The hot spectrum is based on a model fit to UV observations of OH(A) emission from space shuttle plumes (Bernstein et al. 2003). This spectrum is not well characterized in terms of effective OH(A) vibration and rotation temperatures and corresponds to a high degree of excitation in both the vibrational and rotational energy level populations.

The retrieved rotational temperature for the cold components was \(T_\text{rot} = 250 \pm 50\) K. The approximately equal contributions of the cool \((0, 0)\) and \((1, 1)\) components are consistent with
stellar pumping of vibrationally cold OH(X); this is also consistent with the lack of observed (2, 2) and (2, 1) cold features. Typically, the (1, 0) feature, which occurs around 2850 Å, is observed with about a third of the intensity of the (1, 1) feature. The fact that it is not seen in the RR spectrum indicates that the (1, 0) transition is optically thick. The cold $\nu' = 1$ state is excited by stellar pumping of the (1, 0) transitions; however, the reemitted photons preferentially “escape” via the optically thin (1, 1) transition.

4. DISCUSSION OF THE MODELING

4.1. Hot CO(A) Emission Vibrational and Rotational Temperatures

The extremely high retrieved CO(A) emission temperatures, $T_v = 20,000$ K, are a direct consequence of the efficient stellar pumping to high $\nu'$ levels. From inspection of Figure 6, it is evident that all of the CO absorption bands are highly saturated. This means that the populations of the CO(A) $\nu'$ levels will be determined more by the available stellar flux as opposed to the inherent oscillator strengths. It is clear that levels beyond $\nu' = 9$, the highest observed level, are also significantly populated. In terms of an equivalent temperature the $\nu' = 9$ level corresponds to 17,000 K. Thus, the retrieved effective vibrational temperatures are in line with expectations based on the stellar pumping. On reflection, we should have adopted a two-temperature model for the vibrational distribution because of the strong discontinuity of the source flux at 1450 Å, although it is noted that the single-temperature model yielded a respectable fit to both the FOS and GHRS data.

The CO(A) state rotational temperature retrieved from the hot component of the emission data was $T_v = 5500$ K, in contrast to the much lower temperature retrieved for the CO(X) absorption data [the CO(A) excitation source] of $T_v = 2000$ K—a discrepancy of 3500 K! The resolution of this apparent discrepancy is also attributed to the strong saturation effects for the absorption. Analogously to the $\nu'$ population distribution, the population of the associated $j'$ levels will also be governed more by the available flux than their inherent line strengths. The effect of saturation is to enhance the population of high $j'$ levels relative to lower $j'$ levels. This enhancement is then directly observed in the emission spectrum, because these transitions are optically thin.

4.2. Previous GHRS Model

Glinski et al. (1997) previously analyzed the GHRS CO(A–X) emission spectrum. They employed a four-component model consisting of a first principles spectroscopic model for the hot and cold rotational emission components and empirical Gaussian models to account for cold absorption and broad emission features near the band origins. Using these components and allowing for separate velocity shifts and dispersions for each, semiquantitative agreement with all three GHRS spectral regions was obtained. The current first principles–based model results in an improved fit based on just two, hot and cold, emission components. Because of the semiempirical nature of the previous approach, it is not possible to compare it in detail to all aspects of the current modeling approach. However, it is noted that both models use the same rotational temperature for the cold emission features. The rotational temperatures for the hot emission component both indicate a high temperature but are significantly different—3000 K versus 5500 K—for the former and present model. In part, this difference is related to the assumption in the earlier model that only a single ($\nu'$, $\nu''$) band contributes to each of the GHRS spectra, whereas the current modeling indicates that multiple bands contribute to each. The previous approach found a significant cold absorption component for each spectrum, whereas the current approach found cold absorption only for the (0, 1) band. This difference results in very different estimates of the CO(X) ground-state vibrational temperatures, $\sim 5000$ K versus 500–1000 K for the former and present model. Very different velocity shifts were retrieved for the two models, for the hot emission component, +0.1 Å versus −1.6 Å for the former versus current model.

4.3. Geometry of the Emitting Region

Figure 12 depicts a radiative-transport geometry that is consistent with the different observed velocity shifts for the absorption by the hot CO below 1600 Å, and the hot and cold CO emissions above 1600 Å. The overall geometry is based on that deduced from optical scattering data by Waelkens et al. (1996) and the HST observations (Cohen et al. 2004), which indicate an opening angle of 40° for the conical flows. The binary system is obscured from view by a thick dust disk. In the optical spectral region, all the observed light has been attributed to dust scattering in the unobscured portions of conical outflows (Waelkens et al. 1996). The direct emission from the source species is not observed and presumably originates in the obscured region. We speculate that there is a very fast CO outflow, $\sim 300$ km s$^{-1}$, that originates from and is localized near the binary system. Assuming that the CO absorption in the portion of the conical outflows not obscured by the disk is optically thin, then the CO emission from the stellar-excited fast CO can only be observable via scattering in the outflow. Because of the pinched biconical geometry, only photons emitted in a narrow cone, $\sim \pm 20^\circ$, aligned with the CO flow are available for scattering into the observational line of sight. The scattering process will preserve the velocity shift due to the fast CO, resulting in a blueshifted hot emission component. Any hot redshifted absorption would be produced deep in the cavity more than 10° counterclockwise to the upward normal in the figure, and would likely go undetected. We also require the presence of a much colder and slower CO flow. This will lead to cold emission features with small to negligible blueshifts and corresponding
absorption features with small to negligible redshifts. This is consistent with the observation that the sharp absorption band edges of the CO appear to be unshifted, where as the emission band edges appear highly blueshifted.

In Figure 13 we compare the GHRS spectrum of the (0, 2) $A\rightarrow X$ band with the polarization reported by Reese & Sitko (1996). It is clear that the slow cold CO band and C i band are located at the polarization minimum, indicating that they are undergoing less scattering than the hot fast CO, whose higher polarization indicates production deeper within the scattering region. As illustrated in Figure 12, photons emitted in the direction of the observer are blocked by the dust disk.

We note the remarkable similarity of this RT geometry to the dynamical outflow model proposed by Soker (2005). Soker proposed that there are intermittent periods of enhanced high stellar mass loss that can produce bipolar jets localized within $\sim1$ to $2$ AU of the binary system, with a velocity of $\sim300$ km s$^{-1}$. However, most of the conical volume in the disk-obscured region is filled with much colder and slower velocity gas, $<10$ km s$^{-1}$, from periods of “normal” mass loss.

5. OTHER CONSEQUENCES

5.1. The CO Fourth Positive (4P) Absorption System

The entire spectrum of HD 44179 is a complex mixture of overlapping emission and absorption features. As was demonstrated above, these arise from a complex of highly excited electronic, vibration, and rotation levels, all of which are interconnected by allowed and (normally) “forbidden” $a'\rightarrow X$, $d\rightarrow X$, and $e\rightarrow X$ transitions, which are seen in absorption. The presence of these features in significant strength indicates yet one more possible mechanism capable of exciting the $a^3\Pi$ state that gives rise to the Cameron bands: radiative cascades from the $a'\Sigma^+, d'\Delta$, and $e'\Sigma$ states to the $a^3\Pi$ state (usually referred to as the CO Triplet bands). A fully self-consistent model will need to include a complete treatment of the radiative transfer that includes all of these transitions, but that is beyond the scope of this investigation.

One of the consequences of the tremendous optical thickness of the 4P bands near the low rotational states that dominate the band cores is that transitions back down from the excited $A$ electronic state to the ground $X$ electronic state produce photons that can easily excite nearby CO molecules right back up into the $A$ state. Like $Ly_{\alpha}$ photons in a gas containing neutral H, these photons will simply “rattle around” the gas until there is another way to remove the photon: absorption by dust or another molecule, or by branching into another less-populated electronic state. The presence of a plethora of spin-forbidden absorption features indicates that CO is probably capable of converting some fraction of the energy absorbed in the 4P system into transitions among a variety of other electronic states.

5.2. The Maze

In Figures 2 and 3 we show the entire UV spectrum of HD 44179. At 2200 Å there is a distinct change in slope of the spectrum, coincident with the beginning of the CO emission blends, beginning with the Cameron bands. The dashed line in the figure indicates the continuum level used for the models of the fluorescence emission. The flux shortward of 2200 Å is increasingly dominated by CO emission as one goes to shorter wavelengths. Were it not for the very strong absorption features below 1600 Å, the CO emission complex would continue to much shorter wavelengths. Because of this, one needs to think of the entire region below 1600 Å as one where we are not simply seeing absorption of a stellar photosphere by CO, but rather a complex blend of both emission and absorption by CO overlying a (probably hidden) stellar photosphere. That is, much or all of the emitted photons seen in this region are due to CO itself.
6. THE BIG PICTURE

We have demonstrated that the UV spectrum of HD 441679 is completely dominated by a complex network of absorption and emission due to CO. There are two important consequences of this.

First, little or no stellar photosphere is apparent in the flux shortward of 2200 Å. While it seems likely that exciting UV flux may be coming from an obscured (along our line of sight) hot compact companion to the luminous A–F primary star of HD 44179, or an accretion disk deeply embedded in the system, at present we cannot detect it unambiguously in the UV spectrum. Unfortunately, this means that one simply cannot derive the dust extinction at these wavelengths using the standard “pair+” method, as was first done by Sitko et al. (1981) and again by Vrijh et al. (2005). The wavelength region between 1550 and 2200 Å is clearly affected by a vast blend of overlapping emission bands of CO. This would make any extinction curve derived to have too little extinction at these wavelengths. Shortward of 1550 Å, the continuation of this emission is joined by even stronger absorption due mostly to the A–X bands, which will distort the extinction curve further, most likely producing too much extinction at these wavelengths.

Second, virtually everything shortward of 2200 Å is due to CO. This means that in order to be able to unambiguously identify other molecular species in this region of the spectrum of HD 44179, it is necessary to first understand the intrinsic spectral signature laid down by CO. In principle, many interesting organic molecules may have absorption and emission features here, but without adequate removal of the effects of CO, it will not be an easy task to identify these other species.

In order to properly characterize the entire wealth of hundreds of lines that are present in the spectrum, it will be necessary to undertake a full radiative transfer treatment of the CO-containing region. Unfortunately, we still have no firm information about the intrinsic (stellar?) photon field that presumably excites the CO, nor are all the transition rates among all of the relevant states of CO completely known. Nevertheless, there may be some important general consequences of the tremendous excitation seen in the CO.

One of these is the possibility that at least some of the unidentified features seen in the Red Rectangle might arise from hitherto unobserved transitions in CO. The high optical depth of the 4P system of CO has the potential to excite a variety of other emission processes. These would include the heating of dust grains, photoexcitation of other molecules, and fluorescent pumping of other transition in CO itself. For the latter two mechanisms, highly nonequilibrium population states will result.

It has been frequently speculated that this UV-pumping might be responsible for some of the other features seen in the Red Rectangle. Sitko (1983) had tentatively identified CO₂ as a possible absorber shortward of 1600 Å, and noted that the width of the feature in energy (as determined from laboratory spectra) and the total energy absorbed (after removal of a few magnitudes of extinction) were comparable to the ERE feature. Unfortunately, the massive complexity of the CO spectrum has so far thwarted any attempt to positively identify the presence of CO₂ in the ultraviolet, although it has been detected at infrared wavelengths (Waters et al. 1998).

Glinski et al. (1996) suggested that other excited emission bands (such as the Asundi a′–a bands of CO, or the comet-tail A–X bands of CO⁺) might be observable at visible wavelengths, although they have not been detected yet. Bennett et al. (1999) suggested the radiatively pumped excited triplet states of CO itself as the possible source of the ERE in the Red Rectangle. But the ERE is now known to be present in many astronomical sources lacking the Red Rectangle’s unusual CO spectrum. It has been suggested that the ERE might represent emission from silicon nanoparticles, PAHs (both neutral and ionized), PAH dimers (charged), hydrogenated amorphous carbons (HACs), and a variety of other organic and inorganic species (Witt et al. 1998, 2006; Ledoux et al. 2001; Witt & Vrijh 2004; Rhee et al. 2007; Berné et al. 2008). Interestingly, the 3.3 μm feature in the Red rectangle is not well correlated with the ERE, but with the blue luminescence (BL; Witt et al. 2006). (Note that thin films of hydrogenated amorphous silicon carbide [Ma et al. 1998] may be capable of producing both, simply by changing the C/Si ratio. These bands arise not from SiC, but a variety of aromatic, Si-H and C-H stretching bands, and are also narrower than the SiC band—one characteristic used by Vrijh et al. [2004] to discount SiC as a source of the BL.)

The unusual environment near HD 44179 may also be relevant to the unidentified narrowband features seen at visible wavelengths. These bands are found throughout much of the Red Rectangle, and their structure is dependent on distance from the star itself (van Winckel et al. 2002). Many suggestions have been made for the source of these bands, such as carbynes (Warren-Smith et al. 1981) and PAHs (d’Hendecourt et al. 1986). Their wavelengths are also tantalizingly close to those of the diffuse interstellar bands (DIBs), and a connection between the two has been suggested many times (Scarrott et al. 1992; van Winckel et al. 2002; Sarre 2006), although a definitive connection has proved elusive. The DIBs are ubiquitous; the optical bands of the Red Rectangle are not. Is CO of any relevance?

In Figure 2 we are able to trace the A–X absorption system to at least (9, 0), indicating that the v′ = 9 level of the A 1Π electronic state is being populated. The presence of the (1, 3) and (1, 4) fluorescence bands of the A–X system shown in Figure 4 means that the v′ = 1 energy state (∼66,000 cm⁻¹) is populated sufficiently to allow downward transitions sufficiently strong to produce the fluorescent emission. The spin-forbidden absorptions observed in the a′–X, d–X, and e–X are also populating energy levels close to this. We also know that the lowest energy states in the a′ 1Π state (∼48,000 cm⁻¹) are populated. If these states support transitions among one another, they may produce a complex blend of features at energies starting at ∼5600 Å (18,000 cm⁻¹) and continuing to longer wavelengths. This may be the origin of the narrow features seen in the Red Rectangle, at least within a few arcseconds of the central stars, where CO seems to dominate all other identified species. Whether it is capable of producing emission far from the stars, including the change in spectral shape with distance, is unknown and possibly problematic.

A second consequence is that even if CO is not directly responsible for any of the features observed at longer wavelengths, it will still be necessary to provide precursor molecules for the actual emitters. This would seem to require UV photons and one or more abundant molecules. Both are evident in the spectrum of HD 44179, where we see abundant CO and a source of UV photons. The UV photons that we actually observe are also due to CO, although a more deeply embedded source (stellar or accretion disk) is likely to be present.

At the present time we also have no convincing explanation for the presence of the OH bands. Certainly a source of H₂O in the form of vapor or ice would be a good source for the material. Currently, the material flowing out from the central region to form the nebula is C-rich, not O-rich. Nevertheless, the torus seems to be rich in silicate dust (Waters et al. 1998), possibly the result...
of the outflow of material from one or both stars prior to the dredge-up of the He-burning products from its (their) interiors.

In the model of the star system by Men’shchikov et al. (2002) the system began as a pair of stars of roughly $2 M_\odot$, each, separated by $126 R_\odot$. In their model, the originally more massive star has passed through the red giant phase and is now a He-rich white dwarf, while the other star has just completed a similar evolutionary path (but becoming an AGB star), and is the source for much of the current mass outflow in the system. Throughout, the separation of the pair varies, increasing from 126 to 589 $R_\odot$ and finally back to 192 $R_\odot$. In the process, a dust envelope of mass $M_{\text{dust}} \sim 0.015 M_\odot$ is produced. Under the assumption that the gas shed from the evolving stars is retained in the system, a net $1.2 M_\odot$ remains in the envelope.

Some post-AGB stars are known to possess circumstellar grains of water ice. HD 161796 seems to have gone through a phase that lasted approximately 900 yr, where a massive outflow condensed into grains that contained about 27% of the dust mass (Hoogzaad et al. 2002). If such a system had a secondary star which could later reheat this material, a ready supply of water vapor would be available for photodissociation by a hot source of photons. Many post-AGB systems possess O-rich dust, where in many cases the dust is stored in an unseen companion (van Winckel 2003).

Saavik Ford et al. (2001) have suggested that the water vapor observed in IRC +10216 could have even been produced by the vaporization of a “Kuiper Belt” cometary system surrounding the star. If located far enough from the system, water ice could survive the evolution of the star into the AGB phase. In their case, a mass of ice of $\sim 10 M_\odot$ is required. The total mass in the form of solids is likely higher, but not by a large amount. While this is significantly less than the mass of dust seen in the Red Rectangle, it does suggest that at least some fraction of the O-rich material could be “primordial” as well as from a stellar outflow.

It is known that multiple star systems are capable of retaining their pre-main-sequence disk material. Hen 3-600 is a triple star system, with two close stars and a more distant companion, with a small circumbinary disk surrounding the tightly bound components, and a more distant one that is exterior to the orbit of the outer component (Webb et al. 2001; Zuckerman 2001). We also know that some main-sequence stars with masses similar to those sometimes suggested for HD 44179 (Men’shchikov et al. 2002), $\sim 2 M_\odot$, have retained significant dust material after $10^7$ yr or more (Di Folco et al. 2004), a significant fraction of their main-sequence lifetimes. What will happen to these disks (and planetesimals, comnetesimals, and planets) when one of these stars inevitably evolves to become a red giant or AGB star as HD 44179 has?

For the Red Rectangle system, with main-sequence luminosities of $L \sim 11 L_\odot$, each star would be capable of supporting a circumstellar disk with a “snow line” for H$_2$O as close as 11 AU (using a sublimation temperature in a vacuum of 150 K), and a more distant circumbinary disk. In the outer disk, each volatile will possess its own “snow line.” The changing separation of the stars, as well as the growth of the luminosity to near 6000 $L_\odot$, virtually ensures that any objects (asteroids, comets, or planets) within the system will undergo significant changes in temperature. As the luminosities rise, the more volatile species (CO, CO$_2$, CH$_4$, NH$_3$) will vaporize first, with H$_2$O lagging significantly behind. Gravitational perturbations due to the changing separation of the wide components may stir the system further. Whether such a system can provide the OH in sufficient quantities, along with the other positively identified species seen in the system—CH, CH$^+$, and CN (Hobbs et al. 2004)—is unknown.

Higher quality spectra of the OH bands, and a search for rovibrational transitions in the near-infrared are needed for further clarification.

7. WHERE IS THE HOT EXCITING STAR?

The high degree of excitation in the CO bands requires a source of ultraviolet flux. The more evolved star, now the “secondary” star whose light has not been detected at visible wavelengths, is generally assumed to be either a white dwarf or on its way to becoming one. In the model of Men’shchikov et al. (2002) the UV tail in the spectrum is identified with this object. However, we have demonstrated that in fact this part of the spectrum is dominated by a complex blend of CO emission, not a stellar photosphere. Nevertheless, it would be important to be able to know something about this star.

In Figure 14 we show the spectrum in the vicinity of 1930 Å. In addition to the narrow C I band structure discussed by Glinski et al. (1996) HD 44179 also exhibits a broad feature best seen in the low-resolution FOS spectra (Fig. 3). Also shown in Figure 14 is the UV spectrum of the white dwarf star in the LDS 678 binary star system, taken from the IUE archives, and whose spectrum has been discussed by Wegner (1981). To simulate the effects of the added emission line blanketing due to the $A-X$ and $a-X$ bands of CO in this region, and possible mismatch in true C I band strength, we have diluted the spectrum of LDS 678 by a factor of 4.5 and added a constant flux of $4.4 \times 10^{-14}$ erg cm$^{-2}$ s$^{-1}$ Å$^{-1}$.

While Wegner (1981) described LDS 678B as a cool ($T \sim 10,000$ K) DC type white dwarf, more recent work by Oswalt et al. (1991) classify it as DBQA5, and also refer to it as LDS 678A to be consistent with Luyten’s original designation. The parallax of the system (Harrington & Dahn 1980) is consistent with a distance of 10 pc. The distance to HD 44179 that would be implied by the scaling of LDS 678A used in Figure 14 would only be 21 pc (closer if suffering extinction)! This is, however, far closer than any other estimate of the distance to the Red Rectangle.
The spectrum of LDS 678A also peaks near 3500 Å in these flux units, also not consistent with what is seen. Likewise, we do not see the other strong C i bands exhibited by LDS 678A at shorter wavelengths, although these might be hopelessly blended with the CO bands. Finally, if the hot component in HD 44179 were a very luminous hot star, as suggested by Men’shchikov et al., its surface gravity would be insufficient to produce the broad asymmetric C i band that due to normal pressure effects, unless the atmosphere were unusually transparent, allowing us to see deep high-pressure layers.

Other possible explanations for the shape of this feature is that it is the result of a wind containing C i, or that it is a red-degraded band of some as yet unknown molecular species. If the former, it would require an outflow speed $v \approx 8000 \text{ km s}^{-1} = 0.02c$, which seems unlikely. The alternative is a blue-degraded band structure of some molecule which just happens to coincide with the rest wavelength of the C i line. This explanation cannot be ruled out, but hardly seems satisfactory.

Without further information, it is difficult to assign any specific origin for this feature.

### 8. CONCLUSIONS

The UV spectrum of HD 44179 provides an unusual window into the energetics of the innermost regions of the Red Rectangle nebula. There are more identified electronic-vibration bands in this portion of the spectrum than are known anywhere else in the nebula. Modeling the features has revealed information on the vibrational, rotational, and kinetic temperatures of the gas. The high optical depth of the dominant species, CO, is bound to have a profound effect on the emission of all other molecules that coexist with it.

The entire spectrum at wavelengths shortward of 2200 Å is dominated by a complex of emission and absorption bands due to CO with high rotational and vibrational temperatures. The column density is high enough that spin-forbidden transitions are observed in absorption. The strength and complexity of the spectrum makes deriving the extinction characteristics in this region of the spectrum virtually impossible.

We have tentatively identified emission bands due to OH between 2800 and 3300 Å. Such bands might be the result of the sublimation and subsequent photodissociation of H₂O ice, yielding OH in an excited state. Additional emission is due to photoinduced fluorescence.

Further progress will require detailed radiative transfer calculations for CO. Additional observational progress in the UV will require the use of new instruments, such as the Cosmic Origins Spectrograph (COS) planned for the HST.

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