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Experimental characterization of the weakly anisotropic CN $X^2\Sigma^+$ + Ne potential from IR-UV double resonance studies of the CN-Ne complex

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IR-UV double resonance spectroscopy has been used to characterize hindered internal rotor states ($n^R = 0^R$, $1^R$, and $2^R$) of the CN-Ne complex in its ground electronic state with various degrees of CN stretch ($v_{CN}$) excitation. Rotationally resolved infrared overtone spectra of the CN-Ne complex exhibit perturbations arising from Coriolis coupling between the closely spaced hindered rotor states ($1^R$ and $0^R$) with two quanta of CN stretch ($v_{CN} = 2$). A deperturbation analysis is used to obtain accurate rotational constants and associated average CN center-of-mass to Ne separation distances as well as the coupling strength. The energetic ordering and spacings of the hindered internal rotor states provide a direct reflection of the weakly anisotropic internmolecular potential between CN $X^2\Sigma^+$ and Ne, with only an 8 cm$^{-1}$ barrier to CN internal rotation, from which radially averaged anisotropy parameters ($V_{10}$ and $V_{20}$) are extracted that are consistent for $v_{CN} = 0-3$. Complementary ab initio calculation of the CN $X^2\Sigma^+$ + Ne potential using MRCI+Q extrapolated to the complete one-electron basis set limit is compared with the experimentally derived anisotropy by optimizing the radial potential at each angle. Experiment and theory are in excellent accord, both indicating a bent minimum energy configuration and nearly free rotor behavior. Analogous experimental and theoretical studies of the CN-Ne complex upon electronic excitation to the CN $B^2\Sigma^+$ state indicate a slightly more anisotropic potential with a linear CN-Ne minimum energy configuration. The results from these IR-UV double resonance studies are compared with prior electronic spectroscopy and theoretical studies of the CN-Ne system. © 2011 American Institute of Physics. [doi:10.1063/1.3586810]

I. INTRODUCTION

Many reactions of CN radicals exhibit unusual kinetic behavior indicative of association of the reactants in a weakly bound radical-molecule complex. This is evident in the reaction kinetics of the CN radical with hydrocarbons,¹–⁴ which have been studied extensively due to the prevalence of cyanogen in the combustion products of fossil fuels and its resulting role in NOx chemistry.⁵ In such environments, the CN radical acts as a pseudo-halogen, reacting rapidly with the gaseous hydrocarbons through addition or hydrogen abstraction. Large rate coefficients in both of these cases suggest that the reactions are barrierless with negative activation energies indicating that pre-reactive complexes are formed, albeit with short lifetimes.² Crossed molecular beam studies of CN radicals with unsaturated hydrocarbons⁶ have shown that the first step in the reaction pathway is the production of a collision complex, in which the CN radical adds to the π electron density of the hydrocarbon. Although this type of addition mechanism would not be possible with saturated hydrocarbons, theoretical work by Georgievskii and Klippenstein⁷ has predicted a weakly bound van der Waals complex and submerger barrier, separating the pre-reactive complex from product formation, to explain the “strange kinetics” and unusual temperature dependence of the C₂H₆ + CN hydrogen abstraction reaction. Isolating and studying CN radical-rare gas (CN-Rg) complexes provides a starting point to study CN radical-molecule complexes, which have not yet been observed directly.

Thus far, the electronic spectroscopy and excited state predissociation dynamics of CN-Rg and CN-H₂ van der Waals complexes have been examined by Heaven and coworkers.⁸–¹¹ CN-Ne, CN-Ar, and CN-H₂ were investigated using laser induced fluorescence (LIF) with initial work on each system focusing on electronic excitation and fluorescence detection in the CN $B^2\Sigma^+ - X^2\Sigma^+$ region, hereafter referred to as $B-X$ transitions of the complexes. The notation used to denote electronic states of these complexes is not rigorous, but has been adopted in this work and all previous studies to denote states which correlate closely with those of free-CN. Rotationally structured CN-Ne $B (v_{CN} = 0) - X (v_{CN} = 0)$ excitation spectra were observed by collection of $B$-state fluorescence from the complex itself, with the electronic origin lying close in energy to that of free-CN. This is in stark contrast to the dynamics of CN-Ar and CN-H₂ complexes upon $B$-X excitation: CN-Ar rapidly predissociates into CN $A^2\Pi +$ Ar, such that rovibrionic spectra could be observed only by collecting CN A-X emission. The CN-H₂ $B$-X excitation spectrum displays no resolvable structure and was attributed to bound-free excitation with subsequent emission from CN $B^2\Sigma^+$ products. These studies provide a significant amount of information on the excited $B$ (and $A$) electronic states of these
CN complexes; however, little spectroscopic information was obtained on these CN complexes in their ground electronic state. The present study provides the first infrared spectroscopic study of a CN radical complex, specifically CN-Ne, which is utilized to characterize the anisotropy of the CN \( X^2\Sigma^+ + Ne \) intermolecular potential. Similar studies of CN-Ar and CN-H\(_2\) will be reported elsewhere.

Alexander and co-workers have generated high level \textit{ab initio} potential energy surfaces (PES) for CN-Ne, CN-Ar, and CN-H\(_2\), and these were subsequently used in analysis of spectroscopic data, giving insight on the structure and dynamics of the complexes.\(^{12-15}\) The CN \( X^2\Sigma^+ + Ne \) MRCI+Q surface indicates a long-range van der Waals type interaction with Ne bound to the CN \( X^2\Sigma^+ \) radical by just 31.3 \(cm^{-1} (D_e)\) at an intermolecular separation \( (R_e) \) of 3.75 Å. The intermolecular potential exhibits minimal anisotropy, with a barrier to CN internal rotation of less than 2 \(cm^{-1}\). Another \textit{ab initio} study of the CN \( X^2\Sigma^+ + Ne \) potential by Vrabel \textit{et al.} yielded a similar angular dependence for the interaction energy using restricted (and unrestricted) MP2 theory,\(^{16}\) with \( D_e = 32.0 \text{ cm}^{-1}, R_e = 3.75 \text{ Å}, \) and a barrier to internal rotation of less than 5 \(cm^{-1}\). The anisotropy of the ground state PES changes for different partners; for example, the CN \( X^2\Sigma^+ + Ne \) and H\(_2\) potentials are relatively flat,\(^{14,15}\) with CN likely to undergo almost free internal rotation, while the CN \( X^2\Sigma^+ + Ar \) potential is predicted to be more anisotropic with a bent minimum energy structure.\(^{12}\) The minimum energy configuration also changes upon electronic excitation of CN to the \( B^2\Sigma^+ \) state, which reverses the direction of the CN dipole moment,\(^{17}\) with CN-Ar favoring a linear minimum energy configuration.

Given the shallow nature of the ground and excited state potentials for CN-Rg complexes, both infrared and vibronic spectra will likely be dominated by progressions in the low frequency “bending” vibration associated with hindered internal rotor motion. For this reason, this work follows the guide to open shell van der Waals complexes by Dubernet \textit{et al.} which reduces the effective Hamiltonian for the system to that of the bending (or internal hindered rotor) mode alone.\(^ {18,19}\) For complexes of CN in \( ^2\Sigma^+ \) states, which includes its \( X^2\Sigma^+ \) and \( B^2\Sigma^+ \) electronic states, this reduced Hamiltonian follows Hund’s case (b) approximations and is similar to that of a closed shell system because the unpaired electron spin is only weakly coupled to the nuclear axes. It should be noted that the quantum labels used in such approximations in previous literature are not consistent, and therefore it is useful to define the notation that will be used throughout this work. In such atom-diatom complexes, the diatomic monomer rotational quantum number (denoted as \( n \)) is quantized along the intermolecular axis.\(^ {18,20}\) This projection quantum number is labeled \( K. L \) is the pseudo diatomic quantum number for the end-over-end rotation of the complex, and this can couple with \( K \) to form the resultant \( N \), which is the total angular momentum (excluding electron spin) and is a good quantum number. Electron spin can only weakly couple with \( N \), and therefore is neglected. As a result, CN-Ne states will be labeled by their hindered rotor state \((nK)\), CN vibrational state \((\nu_{CN})\), and CN electronic state \((X \text{ or } B)\), as needed; the intermolecular stretch \((\nu_s)\) is not observed and therefore is generally not included in state labels.

This paper focuses on the CN-Ne complex, which has previously been investigated through electronic spectroscopy in the \( B-X \) region by Heaven and co-workers.\(^ {9,11}\) In the present work, an IR-UV double resonance technique is employed to probe the ground electronic state of the CN-Ne complex, principally with two quanta of CN stretch, leading to the observation of several rotationally resolved bands. The rotational structure of these bands and spacings between associated angular states are used to obtain new information on the anisotropy of the CN \( X^2\Sigma^+ + Ne \) potential, which is complemented by high level \textit{ab initio} calculations and compared with previous experimental work on this system.

\[\text{CN} \quad X^2\Sigma^+ + Ne \]

\[\text{CN} \quad B^2\Sigma^+ (\nu = 0) + Ne \]

\[\text{CN} \quad X^2\Sigma^+ (\nu = 2) + Ne \]

\[\text{CN} \quad X^2\Sigma^+ (\nu = 0) + Ne \]

\[\text{R} / \text{Å} \]

\[3.0 \quad 3.5 \quad 4.0 \quad 4.5 \quad 5.0 \quad 5.5 \]

\[\text{FIG. 1. IR-UV double resonance scheme is shown on schematic radial potentials for CN-Ne in the CN stretch overtone region as well as UV spectra in the CN } B^2\Sigma^+ (\nu = 0)-X^2\Sigma^+ (\nu = 2) \text{ spectral region. As depicted schematically in Fig. 1 the IR laser prepares the CN-Ne complexes with two quanta of CN vibrational excitation. The UV laser then promotes the vibrationally excited CN-Ne complexes to the electronically excited } B \text{ state, from which } B-X \text{ fluorescence is collected. IR spectra are recorded by fixing the UV laser on a CN-Ne electronic transition and scanning the IR laser,} \]

\[\text{UV}^+ \quad \text{UV} \]

\[\text{CN} \quad X^2\Sigma^+ (\nu = 2) + Ne \]

\[\text{CN} \quad X^2\Sigma^+ (\nu = 0) + Ne \]
while UV enhancement spectra are obtained by fixing the IR laser on a CN-Ne vibrational transition and scanning the UV laser.

Cyanogen iodide (ICN), which served as the CN radical precursor, is synthesized following methods described previously.\textsuperscript{21,22} ICN vapor (1 torr at 25°C (Ref. 23)) is entrained in a 75% Ne/He carrier gas (30 psi) by flowing the carrier gas over solid ICN contained in a polytetrafluoroethylene sample holder.\textsuperscript{9,11} CN radicals are generated by 193 nm (Ref. 24) or 248 nm (Ref. 25) photolysis (Coherent, LPX 105i excimer laser) of the ICN vapor in the throat of a pulsed supersonic expansion (General Valve, Parker Hannifin Series 9). The photolytically generated CN radicals associate with Ne in the collisional region of the expansion to form CN-Ne complexes. At a distance approximately 10 nozzle diameters downstream from the orifice, the gas mixture is interrogated with the spatially overlapped, counter-propagating IR and UV laser beams.

Tunable IR radiation with a 0.1 cm\textsuperscript{-1} bandwidth is generated with an optical parametric oscillator (OPO, LaserVision), pumped by an injection-seeded Nd:YAG laser (Continuum Precision II 8000), and operates in the CN overtone region at ∼2.5 μm at typical pulse powers of 10 mJ. The probe laser is the output of a Nd:YAG (Innolas, SL600) pumped dye laser (Radiant Dyes, Narrowscan) using Coumarin 460 dye, which operates in the vicinity of the CN B 2Σ\textsuperscript{+}–X 2Σ\textsuperscript{−} (0,2) transition at ∼460 nm at considerably less than 1 mJ/pulse; the fundamental dye output has a bandwidth of 0.1 cm\textsuperscript{-1}. The IR laser is focused into the center of the molecular beam using a 50 cm focal length lens, and is overlapped spatially with the collimated but unfocused UV laser. Both IR and UV lasers are calibrated using a wavelength meter (Coherent Wavemaster). The IR laser (10 Hz) is run at half the repetition rate of the UV laser (20 Hz) so that active background subtraction, |{IR + UV} – UV|, can be employed on alternating laser shots to remove any background UV-only signal from IR-UV double resonance spectra. The delay between the IR and UV lasers is typically set at 50 ns. The resulting fluorescence in the CN B (v = 0)–X (v = 0) region (with a lifetime of τ = 65 ns) is collimated using f/10 optics and passed through a 390 (FWHM = 10 nm) bandpass filter before impinging on a photomultiplier tube (ET Enterprises 9816B). The laser-induced fluorescence signal is then preamplified and displayed on a digital storage oscilloscope (LeCroy WaveRunner 44xi) attached to a PC for further analysis.

Rovibronic spectra of uncomplexed CN radicals were recorded to assess its rotational and vibrational distributions in the supersonic jet. The resulting experimental spectra were simulated using a rotational temperature of ∼4 K, but with significant vibrational population observed in higher vibrational levels including v = 2 at both photolysis wavelengths. In addition, CN-Ne complexes were readily detected with two quanta of CN vibrational excitation without IR excitation when 193 nm photolysis was utilized. In some cases, this resulted in a UV-only signal from the probe laser alone, which was subtracted in IR-UV double resonance spectra. This CN-Ne UV-only signal was eliminated with photolysis at 248 nm, resulting in nearly zero background when recording IR-UV spectra.

\textit{Ab initio} potential energy surfaces were generated for CN X 2Σ\textsuperscript{+} + Ne and CN B 2Σ\textsuperscript{+} + Ne as a function of the CN center-of-mass to Ne distance (R) and intermolecular angle (θ) at a fixed, experimentally determined, CN bond length (r = 1.1718 Å).\textsuperscript{26,27} The angle θ between R and r is defined as 0° for the linear CN-Ne configuration, adopting the convention of previous workers. The surfaces were generated using multireference configuration interaction (MRCI) energies with a Davidson correction (MRCI+Q), following the approach of Yang and Alexander\textsuperscript{15} and extending their prior work with extrapolation to the one electron complete basis set (CBS) limit. The procedures used for the MRCI calculations and extrapolation to the CBS limit are given in supplementary information.\textsuperscript{28} The potential energy landscape was generated by calculating energies every 0.25 Å for 3 Å < R < 4.5 Å and R = 6 Å for 10 values of θ between 0° (CN-Ne) and 180° (Ne-CN). All calculations were carried out in the MOLPRO suite of programs.\textsuperscript{29}

### III. EXPERIMENTAL RESULTS

Following 193 nm photolysis of ICN in Ne carrier gas, two new CN-Ne features are observed upon UV-only excitation in the CN B (v = 0)–X (v = 2) spectral region, neither of which can be attributed to CN alone. Analogous CN-Ne spectral features have been observed previously in the CN B (v = 0)–X (v = 0) region by Heaven and co-workers.\textsuperscript{9,11} The CN-Ne features in the B (v = 0)–X (v = 2) region are assigned as transitions from X (v\textsubscript{CN} = 2, n\textsuperscript{K} = 0\textsuperscript{\textsubscript{0}}) to B (0, 0\textsuperscript{\textsubscript{0}} and 1\textsuperscript{\textsubscript{1}}), with no intermolecular stretch (ν\textsubscript{r}) excitation, by comparison with the prior assignments in the B (v = 0)–X (v = 0) region. The energy difference between analogous CN-Ne features in the two regions is 4058.6(1) cm\textsuperscript{-1} and corresponds to two quanta of CN stretch with no intermolecular excitation (n\textsuperscript{r} = 0\textsuperscript{\textsubscript{0}}) in the ground electronic state of the CN-Ne complex. This value is within experimental uncertainty of the overtone transition of the CN monomer at 4058.55 cm\textsuperscript{-1}.\textsuperscript{30} The negligible change in the CN stretching frequency upon complexation is indicative of very little change in binding energy for CN-Ne upon CN stretch excitation.

IR-UV double resonance spectra of CN-Ne are also recorded with the IR laser preparing various hindered rotor states (ν\textsubscript{CN} = 2, n\textsuperscript{K}) in the ground electronic state and subsequent UV laser excitation to (ν\textsubscript{CN} = 0, n\textsuperscript{K}) levels in the excited B electronic state as shown in Fig. 1. IR-UV double resonance excitation was initially achieved with the IR laser positioned at 4061.7 cm\textsuperscript{-1}. The UV laser was then scanned revealing five features in the UV enhancement spectra as shown in Fig. 2. Upper state assignments are made by comparing the energy and rotational band structures of the features observed in the B (v = 0)–X (v = 2) region with those previously reported in the B (v = 0)–X (v = 0) region (see Table I). The sum of the IR and UV transition frequencies correspond precisely to the UV transition frequencies of CN-Ne features in the B (v = 0)–X (v = 0) region.\textsuperscript{9}

IR spectra of CN-Ne, such as the band seen at 4061.7(1) cm\textsuperscript{-1} in Fig. 3, are then recorded by scanning the IR laser with the UV laser fixed on the Q-branch of an
electronic transition, in this case at 21 750.5(1) cm$^{-1}$. Positioning the UV laser on the $Q$-branch is advantageous for observing rotationally structured IR spectra without scanning the UV laser. Simulation of the UV transition indicates that the $Q$-branch encompasses $N \leq 7$ rotational levels as a result of power broadening, which adds a Lorentzian component to the linewidth of 0.5 cm$^{-1}$. Spectra of other IR bands of CN-Ne are recorded in a similar fashion, however, in each case using different UV transitions since the IR transitions terminate on different hindered rotor states. The band origins of the IR and UV transitions used to obtain IR-UV double-resonance spectra are listed in Table I. Varying the IR-UV time delay shows that the vibrationally activated CN-Ne complexes are long lived, with lifetimes longer than several microseconds. The long lifetime of CN-Ne upon CN overtone excitation is consistent with slow vibrational relaxation of CN ($v = 2$) with nonreactive partners studied previously.$^{31,32}$

**FIG. 3.** Rotationally structured CN-Ne bands in the CN overtone region observed by IR-UV double resonance spectroscopy. The $n^b$ hindered rotor states (with $\Delta E$ spacings) associated with the IR transitions are indicated in the energy level diagram and spectroscopic labels. The $0^0$-$0^0$ transition (gray) is not observed in IR spectra because of $\Delta n = \pm 1$ selection rules, but is deduced from electronic transitions (see text and Table I).

**TABLE I.** Experimental CN-Ne band origins (cm$^{-1}$) with corresponding CN stretch and hindered rotor assignments ($v_{CN}$, $n^b$) for IR and UV transitions in IR-UV double resonance spectra and one-photon $B$-$X$ electronic transitions. Uncertainties in the band positions are consistent with the IR and UV laser bandwidths of 0.10 cm$^{-1}$.

<table>
<thead>
<tr>
<th>IR transition</th>
<th>UV transition</th>
<th>$B$-$X$ transition$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>2045.6 (1,1$^1$-0,0$^0$)</td>
<td>23 766.7 (0,2$^1$-1,1$^1$)</td>
<td>25 812.3 (0,2$^2$-0,0$^0$)</td>
</tr>
<tr>
<td>4055.55 (2,0$^1$-1,1$^0$)</td>
<td>21 745.0 (0,1$^0$-2,1$^1$)</td>
<td>25 800.6 (0,1$^1$-1,1$^1$)</td>
</tr>
<tr>
<td>4058.6 (2,0$^0$-2,0$^0$)$^b$</td>
<td>21 739.2 (0,0$^2$-2,0$^0$)</td>
<td>25 797.8 (0,0$^0$-0,0$^0$)</td>
</tr>
<tr>
<td>4061.74 (2,1$^1$-0,0$^0$)</td>
<td>21 736.0 (0,0$^0$-2,1$^1$)</td>
<td>25 797.8 (0,0$^0$-0,0$^0$)</td>
</tr>
<tr>
<td>4061.74 (2,1$^1$-0,0$^0$)</td>
<td>21 741.9 (0,1$^1$-2,1$^1$)</td>
<td>25 803.7 (0,1$^1$-1,1$^1$)</td>
</tr>
<tr>
<td>4064.3 (2,1$^0$-0,0$^0$)</td>
<td>21 744.8 (0,1$^0$-2,1$^1$)</td>
<td>25 806.5 (0,1$^0$-0,1$^1$)</td>
</tr>
<tr>
<td>4064.3 (2,1$^0$-0,0$^0$)</td>
<td>21 750.5 (0,2$^2$-2,1$^1$)</td>
<td>25 812.3 (0,2$^2$-0,0$^0$)</td>
</tr>
<tr>
<td>4064.3 (2,1$^0$-0,0$^0$)</td>
<td>21 751.9 (0,2$^2$-2,1$^1$)</td>
<td>25 813.6 (0,2$^2$-0,1$^1$)</td>
</tr>
<tr>
<td>6051.6 (3,1$^1$-0,0$^0$)</td>
<td>21 884.3 (1,2$^3$-3,1$^1$)</td>
<td>27 936.0 (1,2$^3$-0,0$^0$)</td>
</tr>
</tbody>
</table>

$^a$This IR transition is not allowed and not observed directly in the present work. See text.

**FIG. 2.** UV transitions observed for CN-Ne complexes in the $B$ ($v_{CN} = 0$)-$X$ ($v_{CN} = 2$) region following IR excitation at 4061.7 cm$^{-1}$, which prepares the $X$ ($v_{CN}, n^b = (2,1)^1$) state. The UV transitions terminate on various $n^b$ hindered rotor states (with $\Delta E$ spacings) in the excited $B$ ($v_{CN} = 0$) electronic state as indicated by the energy level diagram and spectral band labels.
For a few experiments, the OPO is operated in the CN fundamental region at 4.9 μm (0.5 mJ/pulse) and the second overtone region at 1.6 μm (15 mJ/pulse). Strong IR transitions are observed for CN-Ne at 2045.6(1) and 6051.6(1) cm⁻¹, shifted 3.2(1) and 3.3(1) cm⁻¹ to higher energy of the corresponding band origins for free CN, respectively. For these experiments, the UV laser is positioned on CN-Ne B (v_{CN} = 0)–X (v_{CN} = 1) and B (v_{CN} = 1)–X (v_{CN} = 3) transitions at 23 766.7(1) and 21 884.2(1) cm⁻¹, respectively. The sums of the IR and UV frequencies in these instances correspond to known CN-Ne B (v_{CN} = 0)–X (v_{CN} = 0) and B (v_{CN} = 1)–X (v_{CN} = 0) transitions, respectively. The similar shifts of the CN-Ne fundamental, overtone (see Fig. 3), and second overtone transitions from those in free CN leads to their assignment as (v_{CN} = 1, 2, or 3,1)←(0,0) transitions. The similar shift in three spectral regions suggests that there is little change in the hindered rotor states with various degrees of CN vibrational excitation.

The rotational band structures of the IR features at 4061.7 cm⁻¹ and 4064.3 cm⁻¹ are not straightforward to analyze. The close proximity of the n² = 1¹ and 1⁰ upper states (<3 cm⁻¹) will likely result in sizable Coriolis coupling or other perturbations that would lead to complicated rotational structures and intensity patterns. Initial rotational analyses of the 4061.7 cm⁻¹ and 4064.3 cm⁻¹ bands are carried out using the PGOPHER program for simulating rotational structure.33 The line positions of the P and R branches in both bands are fit by floating a common ground state (0,0⁰) rotational constant, B₀₀, the rotational constants for the two upper states (2,1¹ and 2,1⁰), B₁₁ and B₁₀, and band origins, ν₁₁ and ν₁₀, for both transitions. The unresolved Q-branch of the former transition is not included in the fit. The resultant spectral simulations with rotational temperature of 10 K and laser linewidth of 0.1 cm⁻¹ are shown in Fig. 4 and the derived constants are listed in Table II.

Several aspects of the experimental and simulated rotational band structures shown in Fig. 4 are worthy of note. The R-branch of the (2,1¹)←(0,0⁰) band at 4061.7 cm⁻¹ appears congested, indicative of an R-branch bandhead (B₁₁ < B₀₀), suggesting that a perturbing state is pushing the rotational levels of the (2,1¹) state downward. In addition, the position and shading of the Q-branch of the (2,1¹)←(0,0⁰) band is dramatically different from that predicted (dashed curve in upper panel) by spectroscopic constants (Table II) generated from analysis of the P and R branches. Furthermore, the relative intensities of P and R branch lines of the (2,1⁰)←(0,0⁰) band at 4064.3 cm⁻¹ do not match those anticipated from HönI-London factors in the simulation (lower panel).

The preliminary rotational analysis of P- and R-branch lines yields a ground state rotational constant, B₀₀, of 0.106(2) cm⁻¹, in good accord with that reported previously based on electronic spectroscopy studies.9,11 On the other hand, the derived upper state rotational constants, B₁₁ = 0.093(2) cm⁻¹ and B₁₀ = 0.111(2) cm⁻¹, differ significantly from the ground state value. The difference could arise from coupling between the two upper states and/or changes in the geometry of the complex as discussed below.

IV. ANALYSIS

A. Coriolis coupling effects and deperturbation analysis

As presented in Sec. III, several aspects of the rotationally structured CN-Ne bands suggest a perturbation between the (v_{CN}, n²) = (2,1¹) and (2,1⁰) upper states. A significant body of work has been published on infrared spectra of closed-shell HX and DX (X = F, Cl)-Rg complexes in which Coriolis coupling between closely spaced states has been used to explain very similar experimental findings.34–36 These HX-Rg systems exhibit J-dependent shifts in energy levels and deviations in rotational transition strengths, which were analyzed as an l-doubling perturbation between intermolecular bending and/or stretching levels. A similar analysis is performed here for CN-Ne to elucidate “deperturbed” rotational constants for the bands in question, and to account for the line positions and band shapes observed. However, no attempt is made to explicitly treat the line intensities, which are influenced by both Coriolis coupling and the double resonance nature of the experiments.

FIG. 4. Experimental and simulated rotational band contours for the (ν_{CN}, n²) = (2,1¹) ← (0,0⁰) (upper panel) and (2,1⁰) ← (0,0⁰) (lower panel) transitions. The red/blue traces are experimental spectra from IR-UV double resonance measurements, while the black traces are simulations with rotational temperature of 10 K and laser linewidth of 0.1 cm⁻¹. The simulations are based on fits to P- and R-branch lines only. Rotational assignments are shown as ticks. The dashed gray line illustrates the contour for the Q-branch anticipated using the same (perturbed) rotational constant for the (2,1¹) upper state. The experimental Q-branch structure is well represented (red) using the deperturbed rotational constant derived for the (2,1¹) upper state. The simulated line intensities are based on HönI-London factors.
The HX-Rg systems studied by Nesbitt and co-workers were treated as linear complexes, in which the lowest \( \Sigma \cdot \Sigma \) bending transition terminates on \( e \) symmetry upper rotational levels (with equivalent notation of \( n^K = 1^0 \) for the upper state); this is also the case for \( P \)- and \( R \)-branch lines of the \( \Pi \cdot \Sigma \) band (with \( n^K = 1^1 \) upper state). The \( Q \)-branch lines of the \( \Pi \cdot \Sigma \) bending transition, however, terminate on \( f \) symmetry upper levels. The close proximity of the two upper states can give rise to noticeable Coriolis coupling between \( e \) symmetry rotational levels, which is treated by modeling the upper state energy levels through a modified rotational Hamiltonian. The \( Q \)-branch of the \( \Pi \cdot \Sigma \) transition is not perturbed.

Analogous Coriolis coupling is immediately evident in CN-Ne overtone spectra. Preliminary analysis of the transition to the \((2,1)^{1}\) state indicates that the upper state rotational constant for \( e \) symmetry levels, \( B_{11} \), derived from \( P \) and \( R \) branch lines, is reduced relative to the unperturbed ground state value. The \( Q \) branch of this transition, which terminates solely on \( f \) symmetry levels, is not resolved and therefore cannot be used to obtain an unperturbed rotational constant for the \((2,1)^{1}\) state; thus, we utilize \( B_{00} \) for comparisons. Assuming the reduction in rotational constant for the \( 1^1 \) state is a result of Coriolis coupling, we deduce that the perturbing state lies higher in energy and is presumably the nearby \((2,1)^{0}\) state.

A complete deperturbation analysis was then undertaken using a modified rotational Hamiltonian

\[
\hat{H} = \begin{pmatrix}
H_{11} & H_{12} \\
H_{21} & H_{22}
\end{pmatrix},
\]

with matrix elements

\[
H_{11} = v_{10} + B_{10}[N(N + 1) - K^2],
\]

\[
H_{22} = v_{11} + B_{11}[N(N + 1) - K^2],
\]

\[
H_{12} = H_{21} = \beta [N(N + 1)]^{1/2},
\]

where \( B_{10} \) and \( B_{11} \) are the deperturbed rotational constants for the \((2,1)^{1}\) and \((2,1)^{1}\) states, \( v_{10} \) and \( v_{11} \) are their respective band origins listed in Table II. The constant \( \beta \) is related to the magnitude of the Coriolis coupling. The eigenvalues for this \( N \)-dependent Hamiltonian are fit using a least-squares method to rotational energy levels for the \((2,1)^{0}\) and \((2,1)^{1}\) states, which were obtained from simulation of \( P \) and \( R \) branch lines described in Sec. III. All parameters except the band origins are allowed to float in the fitting procedure. Fitting the rotational energy levels in this fashion reveals a strong Coriolis coupling between the \((2,1)^{0}\) and \((2,1)^{1}\) hindered internal rotor states of CN-Ne, which are separated by only \( 2.6(1) \text{ cm}^{-1} \), with \( \beta \) of \( 0.195 \text{ cm}^{-1} \) and deperturbed rotational constants also listed in Table II. The deperturbation analysis shows that Coriolis coupling increases \( / \) decreases the magnitude of the experimentally observed rotational constant for the \((2,1)^{0}(2,1)^{1} \) \( (f \) symmetry) state by nearly \( 20\% \) compared to its unperturbed value.

The rotational constants for the ground state and hindered internal rotor states, the latter following deperturbation, are then used to evaluate the average intermolecular distance, \( \langle R^2 \rangle^{1/2} \), between the center-of-mass of CN and Ne using a pseudo-diatomic model. This yields an average separation distance in the ground \((0,0)^{0}\) state of \( 3.70(4) \text{ Å} \). The average separation in the vibrationally excited \((2,1)^{1}\) state is essentially unchanged at \( 3.71(4) \text{ Å} \), whereas the \((2,1)^{0}\) state shows a substantial increase in the average distance of \( 3.92(4) \text{ Å} \). This indicates a significant change in the properties of hindered rotor states with \( n^K = 1^1 \) and \( 1^0 \) as discussed below.

### B. Hindered rotor analysis

The IR-UV double resonance experiments reveal several hindered rotor states supported by the CN \( X \ 2\Sigma^+ \ (v = 2) \ + \ Ne \) intermolecular potential, namely states with \( n^K = 0^0 \), \( 1^1 \), and \( 1^0 \). No intermolecular stretching states have been observed and thus we focus our theoretical analysis exclusively on the angular states. In this section, we use the assignments and energy spacings of these hindered internal rotor states to deduce information about the anisotropy of the CN \( X \ 2\Sigma^+ \ (v = 2) \ + \ Ne \) potential based solely on the experimental results, which is then extended to CN \( X \ 2\Sigma^+ \ (v = 0) + Ne \) and compared with prior results on CN \( B \ 2\Sigma^+ \ (v = 0) + Ne \). The resultant angular potentials for CN \( X \ 2\Sigma^+ \ + \ B \ 2\Sigma^+ \) are then compared with those obtained from high level \textit{ab initio} theory in Sec. V.

The hindered rotor analysis is based on the theoretical treatment for open-shell diatom-rare gas complexes developed by Dubernet et al.\textsuperscript{18} (and briefly summarized in supplementary information\textsuperscript{26}), utilizing two parameters, \( V_{10} \) and \( V_{20} \), to describe the anisotropy of the system. The parameters can be determined directly from the hindered rotor state spacings observed experimentally, and can then be inserted into the truncated Legendre polynomial

\[
V(\theta) = V_{10} \cos(\theta) + \frac{V_{20}(3 \cos^2(\theta) - 1)}{2},
\]

in order to model the angular intermolecular potential while neglecting any isotropic contribution.

Dubernet et al.\textsuperscript{18} provide a discussion of the limits imposed on these parameters; however, certain restrictions can be derived from the experimental data alone. For example, a simple inspection of the CN-Ne \( X \ (v_{CN} = 2) \) hindered rotor energy level spacings shows that \( V_{10} \) is closer to the limit of \( V_{10} = 0 \) than \( V_{10} = 2 \text{h} \) (illustrated by Fig. 2 of Ref. 18). Furthermore, the energy ordering of the \((2,1)^{1}\) and \((2,1)^{0}\) states indicates that \( V_{20} \) is positive, while the larger spacing between the \((2,0)^{0}\) and \((2,1)^{1}\) states as compared to the \((2,1)^{1}\) and \((2,1)^{0}\) states limits the magnitude of \( V_{20} \) to less than \( 5.1 \text{ cm}^{-1} \). A unique solution for \( V_{10} \) and \( V_{20} \) is found within these constraints yielding values of \( -1.6 \text{ cm}^{-1} \) and \( 4.3 \text{ cm}^{-1} \), respectively, which reproduce the experimentally observed energy spacings precisely. To account for the experimental uncertainty associated with measurement of the IR band origins \((\pm 0.1 \text{ cm}^{-1})\), we also derive a corresponding range of \( V_{10} \) and \( V_{20} \) values, specifically \( -2.1 \) to \(-0.8 \text{ cm}^{-1} \) for \( V_{10} \) and \( 4.3 \pm 0.1 \text{ cm}^{-1} \) for \( V_{20} \). It should be noted that sign of \( V_{10} \), indicating a preference for Ne binding to a specific end of the CN radical, cannot be determined from the experimental methods or associated theoretical analysis used in this work and as
such the sign of $V_{10}$ is arbitrary; we assume $V_{10}$ is negative following prior work.

The corresponding energy spacings for CN-Ne $X$ ($\nu_{CN} = 0$) were then calculated using the same values for the potential anisotropy parameters ($V_{10} = -1.6$ cm$^{-1}$, $V_{20} = 4.3$ cm$^{-1}$) and the appropriate monomer rotational constant for CN $X^2 \Sigma^+$ ($\nu = 0$). This yields an energy spacing between CN-Ne $X$ (0,0$^0$) and (0,1$^1$) states, which is in excellent agreement with the only experimentally observed spacing of 3.1(1) cm$^{-1}$. These anisotropy parameters for the $X$ ($\nu_{CN} = 0$) state of CN-Ne differ somewhat from those reported previously, particularly for $V_{20}$.

Repeating this analysis for CN-Ne $B$ ($\nu_{CN} = 0$), based on the $nK^2 = 0^1$, 1$^1$, 1$^0$, 2$^2$, and 2$^1$ hindered rotor states observed in the $B$ ($\nu = 0$)$-X$ ($\nu = 2$) and $B$ ($\nu = 0$)$-X$ ($\nu = 0$) spectral regions, we obtain values of 8.2 cm$^{-1}$ and 0.6 cm$^{-1}$ for the potential anisotropy parameters $V_{10}$ and $V_{20}$, respectively. These parameters are in good accord with results from a prior analysis by Fei and Heaven.

The effective angular dependence of the CN $X^2 \Sigma^+$ ($\nu$) + Ne potential is then plotted using the truncated Legendre polynomial expansion shown in Eq. (5). The $V_{10}$ and $V_{20}$ anisotropy parameters are radially averaged over the intermolecular distances sampled experimentally for CN $X$ ($\nu = 2$) and are consistent with the data available for CN $X$ ($\nu = 0$), where rotational analysis indicates an average CN-Ne separation of $\langle R^2 \rangle^{1/2} = 3.70(4)$ Å for the ground state. $V(\theta)$ as well as the individual terms in Eq. (5) with associated uncertainties (shown as shaded regions) are plotted in the upper panel of Fig. 5. A similar graph shown in the upper panel of Fig. 6 depicts the effective angular dependence of the CN $B^2 \Sigma^+$ ($\nu = 0$) + Ne potential for comparison, again neglecting isotropic contributions. The angular dependence of the potentials derived from this analysis of hindered internal rotor states of the CN-Ne complex are discussed in the following section.

V. DISCUSSION

A. Experimental determination of potential anisotropy

The angular potential $V(\theta)$ for CN $X^2 \Sigma^+$ + Ne, derived from the analysis of observed hindered rotor states (Sec. IV and supplementary information) and shown in the upper panel of Fig. 5, indicates that the minimum energy geometry of CN $X^2 \Sigma^+$ + Ne is nonlinear with a shallow minimum at $\theta \sim 80^\circ$. The ground state angular potential is dominated by the $V_{20}$ anisotropy term, although the smaller $V_{10}$ term tips the minimum away from 90$^\circ$. The barrier ($\theta \sim 0^\circ$, 180$^\circ$) to internal CN rotation is only ~8 cm$^{-1}$, indicating a weakly anisotropic potential close to the free rotor limit. As a result, the ground state of the CN-Ne complex will have a bent configuration on average, while undergoing wide amplitude motion in the internal bending coordinate. By contrast, the CN $B^2 \Sigma^+$ + Ne potential, plotted in the upper panel of Fig. 6, shows that the minimum energy configuration in the excited $B$ electronic state is linear with a somewhat higher barrier ($\theta \sim 180^\circ$) to internal rotation of ~16 cm$^{-1}$. In this case, the potential anisotropy is dominated by the $V_{10}$ term.

The anisotropy parameters and correlation diagram shown in supplementary Fig. S1 were generated using the CN $X$ ($\nu = 2$) monomer rotational constant. Regeneration of the correlation diagram for CN $X^2 \Sigma^+$ ($\nu = 0$, 1, and 3) + Ne, using the appropriate CN monomer rotational constants with identical $V_{10}$ and $V_{20}$ anisotropy parameters, predicts 0$^0$ to 1$^1$ hindered rotor spacings in each CN spectral region. The values produced in this fashion are consistent with the spacing in each region between the experimentally observed CN-Ne 1$^1$ feature and the anticipated position of the 0$^0$ band, which is assumed to be unchanged from the CN monomer stretch. Thus, the potential anisotropy for CN $X^2 \Sigma^+$ + Ne shown in Fig. 5 appears to be essentially unchanged for CN ($\nu = 0$-4).

B. Ab initio MRCI potentials for CN ($X^2 \Sigma^+$, $B^2 \Sigma^+$) + Ne

Figure 7 shows 2D $ab$ initio potential energy surfaces (PES) for the ground CN $X^2 \Sigma^+$ + Ne and excited CN $B^2 \Sigma^+$ + Ne electronic states, generated in the fashion described in Sec. II and supplementary information. The ground state surface obtained using the MRCI+Q/CBS method builds on that previously reported by Yang and Alexander based on MRCI+Q/aVTZ calculations with both counterpoise and size-consistency corrections. The $ab$ initio PES generated in

FIG. 5. Potential anisotropy for CN $X^2 \Sigma^+$ ($\nu_{CN} = 2$) + Ne as a function of intermolecular angle $\theta$. Upper panel: The solid line indicates the effective angular potential, represented by Eq. (5), while the dotted and dashed lines give the individual contributions from the smaller $V_{10}$ and dominant $V_{20}$ terms derived from the experimental hindered rotor spacings. The gray shaded regions indicate the experimental uncertainties associated with $V_{10}$ and $V_{20}$. Lower panel: The $ab$ initio data points are illustrated by red crosses. The black trace is a fit to the $ab$ initio data using the Legendre expansion with an additional isotropic component.
this work exhibits a shallow global minimum at $\theta = 80^\circ$, consistent with the experimentally determined anisotropy for the CN $X^2\Sigma^+ + \text{Ne}$ potential, but differs from the nearly isotropic potential with very shallow minima at linear $\theta = 0^\circ$ and $180^\circ$ configurations predicted using an AVTZ basis set.\textsuperscript{15} Fitting a radial slice of the potential landscape to a Morse potential at the minimum energy geometry ($\theta = 80^\circ$) yields an intermolecular distance of $R_e = 3.51$ Å and a dissociation energy of $D_e \sim 41$ cm$^{-1}$.

The lower panel of Fig. 5 shows the angular dependence of the \textit{ab initio} potential for CN $X^2\Sigma^+ + \text{Ne}$. The theoretical anisotropy parameters and those determined experimentally yield $V_{10}$ and $V_{20}$ values of $-0.8 \pm 0.1$ cm$^{-1}$ and $3.2 \pm 0.2$ cm$^{-1}$, respectively, for CN $X^2\Sigma^+ + \text{Ne}$, where the uncertainty represents $1\sigma$ in the Legendre fitting procedure.

The \textit{ab initio} CN $B^2\Sigma^+ + \text{Ne}$ PES (upper panel of Fig. 7) exhibits a much higher degree of anisotropy than the ground state, favoring a linear geometry ($\theta = 0^\circ$) with $R_e = 3.76$ Å and $D_e \sim 44$ cm$^{-1}$ evaluated in the same manner as above. Once again, a minimum energy angular path through the potential energy landscape was generated and fit to a truncated Legendre polynomial, shown in the lower panel of Fig. 6. The CN $B^2\Sigma^+ + \text{Ne}$ potential anisotropy is dominated by a $V_{10}$ term of $-7.6 \pm 0.3$ cm$^{-1}$, with significantly less dependence on $V_{20}$ ($-0.5 \pm 0.4$ cm$^{-1}$).

Figures 5 and 6 allow comparisons between the theoretical anisotropy parameters and those determined experimentally. The \textit{ab initio} calculations reproduce the angular dependence of the CN $X^2\Sigma^+ + \text{Ne}$ and CN $B^2\Sigma^+ + \text{Ne}$ potentials remarkably well. The relative signs of
the $V_{10}$ anisotropy parameters can also be determined from these \textit{ab initio} potentials. The excellent agreement between experiment and theory validates these new \textit{ab initio} potentials and the need for complete basis set extrapolation, rather than utilizing counterpoise correction and basis set superposition error approximations with small basis sets.

\section{C. Generation of body fixed wavefunctions and implications}

Wavefunctions for the hindered rotor state are produced through the diagonalization of the experimental angular potential for CN $X^2\Sigma^+ + Ne$ (Fig. 5). As in previous studies of HX complexes, the potential is diagonalized in a body-fixed free rotor basis to give the angular eigenfunctions as a function of $\theta$. The probability distributions for the $n^K = 0^0$, $1^1$ and $1^0$ levels are shown in Fig. 8. In each case, the broad distributions are indicative of wide amplitude hindered internal rotor motion. The probability distribution for the $0^0$ level is peaked in bent configurations slightly favoring the N-side of CN, whereas free rotor behavior samples all configurations with equal probability. The $1^0$ level has maximum probability at linear Ne-CN and CN-Ne orientations, with greater integrated likelihood for Ne to be located on the N-side of CN compared to free rotor behavior. The $1^1$ level is nearly free rotor-like with a most probable bent configuration.

The probability distributions can also be used to explain the different rotational constants (deperturbed) obtained for the CN-Ne (2,10) hindered rotor state as compared to the (2,11) and (0,00) levels (see Table II). The rotational constant for the CN-Ne (2,10) level is some 10\% smaller than those for the other two levels, indicating a corresponding increase in the Ne to CN (center-of-mass) distance for the CN-Ne (2,10) level. The longer effective bond length for the $1^1$ level appears to stem from its most probable linear configuration, which differs from the $0^0$ and $1^0$ levels where nonlinear configurations are most probable. This can be understood by examining the CN $X^2\Sigma^+ + Ne$ \textit{ab initio} potential (Fig. 7), which shows that the optimized internuclear bond length changes from 3.5 Å at the global minimum with $\theta = 80^\circ$ to 3.9 Å for linear CN-Ne ($\theta = 0^\circ$). The van der Waals interaction allows the Ne atom to get closer to C≡N center-of-mass on a sideways approach as compared to a head-on approach to either end of the CN radical. The experimentally derived average bond lengths follow this trend with 3.71(4) Å for the (2,11) level and 3.70(4) Å for the (0,00) level, which have bent configurations, and 3.92(4) Å for the (2,10) level with a linear most probable configuration.

\section{D. Comparison with previous work}

Rotational analysis of the IR-UV double resonance spectra reported in this work has afforded the re-evaluation of the CN-Ne ground state rotational constant as 0.106(1) cm$^{-1}$, which is in excellent agreement with the value previously reported by Fei and Heaven. The weak coupling of the CN stretch to intermolecular degrees of freedom is evident in the very small shift in CN stretching frequency upon complexation, which was also noted in the previous electronic spectroscopy. The IR spectra obtained in the fundamental, overtone, and second overtone regions (for analogous hindered rotor states) show that the CN-Ne hindered rotor band positions change very little with increasing quanta of vibration in the C-N stretching coordinate as compared with free CN stretch. The experimental spectra have also allowed the assignment of accurate anisotropy parameters for CN
$X^2\Sigma^+ + \text{Ne}$, where $V_{10}$ and $V_{20}$ are $-1.6$ cm$^{-1}$ and 4.3 cm$^{-1}$, respectively, and for CN $B^2\Sigma^+ + \text{Ne}$ where $V_{10}$ and $V_{20}$ are $-8.2$ cm$^{-1}$ and 0.6 cm$^{-1}$. These values have been determined directly from the energy spacings of hindered internal rotor states, with $V_{20}$ for CN $X^2\Sigma^+ + \text{Ne}$ close to the value estimated (5.1 cm$^{-1}$), and with the excited state values in excellent agreement with those reported by Lin and Heaven.\(^{11}\)

The new \textit{ab initio} potential energy surface for CN $X^2\Sigma^+ + \text{Ne}$ shows the weakly anisotropic ground state potential energy surface to have a nonlinear minimum energy configuration, consistent with the bent minimum energy configuration predicted with MP2 methods,\(^{16}\) yet contrary to the linear configuration, consistent with the bent minimum energy surface to have a nonlinear minimum energy configuration on average with only an 8 cm$^{-1}$ barrier to internal rotation of CN, giving rise to wide amplitude motion. High-level \textit{ab initio} calculations of the CN $X^2\Sigma^+ + \text{Ne}$ potential were carried out for comparison with the experimentally derived potential. The angular \textit{ab initio} potential for CN-Ne, obtained by optimizing the radial potential at each angle, is in remarkably good agreement with experiment.

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28 See supplementary material at http://dx.doi.org/10.1063/1.3586810 for details of the theoretical methods and hindered rotor analysis including the correlation diagram.


33 *P*Gopher, a program for simulating rotational structure, C. M. Western, University of Bristol, available at http://pgopher.chm.bris.ac.uk.


