ABSTRACT: The cyanate anion (NCO\(^-\)) is a species of considerable astrophysical relevance. It is widely believed to be embedded in interstellar ices present in young stellar objects but has not yet been detected in the dense gas of the interstellar medium. Here we report highly accurate laboratory measurements of the rotational spectrum of the N\(^{13}\)CO\(^-\) isotopologue at submillimeter wavelengths along with the detection of three additional lines of the parent isotopologue up to 437.4 GHz. With this new data, the rotational spectrum of both isotopologues can be predicted to better 0.25 km s\(^{-1}\) in equivalent radial velocity up to 1 THz, more than adequate for an astronomical search in any source. Moreover, a semi-experimental equilibrium structure of the anion is derived by combining the experimental ground-state rotational constants of the two isotopologues with theoretical vibrational corrections, obtained by using the coupled-cluster method with inclusion of single and double excitations and perturbative inclusion of triple excitations (CCSD(T)). The estimated accuracy of the two bond distances is on the order of 5 \times 10^{-4} \text{Å}; a comparison to the values obtained by geometry optimization with the CCSD(T) method and the use of a composite scheme, including additivity and basis-set extrapolation techniques, reveals that this theoretical procedure is very accurate.

1. INTRODUCTION

The cyanate ion is the conjugate base of cyanic (HOCN) and isocyanic (HNCO) acids and serves as a well-known pseudohalide anion in salts which are employed in organic synthesis. It has also a biological role since it is a human metabolite. NCO\(^-\) is almost certainly embedded in interstellar ices with experimental and theoretical studies strongly supporting the argument that the so-called XCN\(^2\) feature at 2165 cm\(^{-1}\) (4.62 \(\mu\)m) observed toward various young stellar objects is due to this ion.\(^3\)--\(^6\) It has been proposed that NCO\(^-\) can be formed on icy grain mantles from the isocyanic acid, either by ionization of the acid interacting with NH\(_3\) solvated by H\(_2\)O molecules\(^1\) or by thermal processing.\(^8\) Additional mechanisms, involving radicals (NH\(_2\)/NH + CO) and suprathermal oxygen atoms (O + CN\(^-\)), have been also put forth,\(^5\) as has a suggestion that NCO\(^-\) is efficiently formed by the interaction of cosmic rays with CO\(_2\)-NH\(_3\) ices.\(^9\)

Experiments of proton irradiation and UV photolysis of nitriles frozen on H\(_2\)O ice, performed to explore the chemistry induced by far-UV photons and ionizing radiation, led to the formation NCO\(^-\), whereas under anhydrous conditions isonitriles were formed.\(^10\) On this basis, Hudson and Moore\(^10\) argue that the cyanate ion is a possible stable radiation product in Titan or cometary ices where both nitriles and H\(_2\)O are present. In addition, its production by electron bombardment of an ice mixture of H\(_2\)O, CO\(_2\), CH\(_4\), NH\(_3\), and CH\(_3\)OH led Bergantini et al.\(^11\) to postulate a relevance of cyanate in the chemistry of Enceladus.

Although NCO\(^-\) does not react by associative detachment with H\(_2\),\(^12\) it has yet to be detected in molecular clouds. Presumably the absence of NCO\(^-\) there points to the lack of an efficient gas-phase formation route. Yurtsever et al.\(^13\) for example, discarded the possibility of radiative electron attachment to the NCO radical because of the high electron affinity of the anion (3.609 eV\(^14\)) and its small size, which makes dissipation of the large amount of energy needed for the stabilization inefficient.

The rotational spectrum of NCO\(^-\) was first reported by Lattanzi et al.\(^15\), who measured its spectrum both in a supersonic molecular beam and in a low-pressure glow discharge between 23 and 368 GHz. In addition to the fundamental (\(J' = 1 \rightarrow 0\)) transition at 23 GHz, which displayed the characteristic triplet of the nitrogen hyperfine...
structure, nine harmonically related lines with \( J = 6−15 \) were also detected. The carrier of these lines can be assigned with certainty to NCO\(^-\) since the derived rotational constant, albeit much more precise, is in very good agreement with early infrared measurements of Gruebele et al.\(^{16}\) In that study, a total of 132 IR transitions in the \( \nu_1 \) (CN stretching) fundamental and corresponding bending and stretching hot bands were reported, and the equilibrium rotational constant derived on the basis of the all three normal coordinates. However, assignments of two hot bands were subsequently questioned, first by Botschwina et al.\(^{17}\) and later by Pak et al.,\(^{18}\) who proposed a reassignment. As a result, the only reliable vibration—rotation interaction constant from the IR analysis is \( \alpha_{bf} \), which arises from the CN stretch.

The present work was undertaken with the intention of determining a precise molecular structure of NCO\(^-\). Our efforts have focused on the millimeter-wave spectrum of the C-13 isotopologue, in addition to extending measurements of the parent isotopologue into the submillimeter-wave region. With the rotational constants available for two isotopologues, a semiexperimental equilibrium structure of the anion was obtained by using vibrational corrections from quantum-chemical calculations.

### 2. METHODS

#### 2.1. Experimental Section. 2.1.1. Parent Isotopologue.

The submillimeter-wave rotational lines of the cyanate ion reported in this work were observed in Bologna with a frequency-modulated spectrometer\(^{19}\) equipped with a discharge cell made of a Pyrex tube, 3.25 m long and of 5 cm diameter, with two cylindrical hollow electrodes 25 cm in length at either end. The radiation source was a frequency quadrupler, which consists of two doublers in cascade (RPG—Radiometer Physics GmbH) driven by a Gunn diode oscillator (J. E. Carlstrom Co.) operating between 75 and 115 GHz. Two phase-lock loops allow the stabilization of the Gunn oscillator with respect to a frequency synthesizer, which is locked to a 5 MHz rubidium standard. Frequency modulation of the radiation is obtained by sine-wave modulating at 6 kHz, the reference signal of the wide-band Gunn synchronizer (total harmonic distortion less than 0.01%). The signal, detected by a liquid-helium-cooled InSb hot electron bolometer (QMC Instr. Ltd. type QF1/2), is demodulated at 2f by a lock-in amplifier.

The best conditions for NCO\(^-\) were found by maximizing the production of HNCO via its \( J_{K,K} = 53_{0,53} ← 52_{1,52} \) transition near 368 GHz.\(^{20}\) Using a DC-discharge operating at room temperature, we combined a 5:2 mixture (7 mTorr, 0.9 Pa) of humid air and CO with He buffer gas for a total pressure of about 15 mTorr (2.0 Pa), and the discharge current varied between 20 and 40 mA. Anion production was slightly enhanced with respect to that achieved in the positive column.\(^{21}\) Not surprisingly, the formation mechanisms of cations and anions are markedly different: for cations, ionizing collisions of the magnetically confined electrons are important, while NCO\(^-\) is more likely produced by dissociative electron attachment to HNCO.\(^{22}\) Our measurements, although indirect, support this hypothesis, in that there is a close correlation between the production of HNCO and NCO\(^-\), regardless of the precursor combination used. A sample spectrum of the \( J = 19 ← 18 \) transition is shown in Figure 1. Because a signal-to-noise ratio of less than 10 was achieved in 10 m of integration and the integrated line strength of the transition is fairly high (5 × 10\(^{-2}\) nm\(^2\) MHz), the steady-state abundance of the anion under our most favorable discharge conditions is low.

Finally, the accuracy of frequency measurements of NCO\(^-\) was carefully checked since line centers can be shifted by the Doppler effect due to the drift velocity of molecular ions produced in the positive column of a DC glow discharge.\(^{23}\) Frequency measurements of the test transition \( J = 16 ← 15 \) at 368.372 GHz, previously observed by Lattanzi et al.,\(^{15}\) did not yield a statistically significant difference in the line frequency when the polarity of the cell electrodes was reversed. This does not preclude the possibility of a small Doppler shift; rather, it may simply be obscured by the poor signal-to-noise ratio.

#### 2.1.2. C-13 Isotopologue. The same free space absorption spectrometer in Cambridge\(^{15}\) that was used to observe the millimeter-wave spectrum of NCO\(^-\) was also employed to detect the C-13-containing isotopologue, N\(^{13}\)CO\(^-\). Radiation was produced by frequency multiplication from one of several phase-locked Gunn oscillator that operate in the region 70–145 GHz and fed into a double-pass absorption cell 3 m in length. The signal, modulated at \( f = 95 \) kHz, was detected by a liquid-helium-cooled InSb hot-electron bolometer and then demodulated at 2f with a lock-in amplifier.

Unlike the original study, \(^{13}\)CO was used instead of CO and was combined with N, and Ar in a 1:2:2 mixture at a total pressure of 10 mTorr (1.3 Pa); the cell was kept at about 290 K, and the discharge current was set at 30 mA. Although no H-containing precursors were used, enough trace hydrogen was present to readily detect HN\(^{13}\)CO, which in turn could dissociate to N\(^{13}\)CO\(^-\) after electron attachment in the low-density plasma. Evidence implicating this production scheme came from detection of anion signal even after the precursor flow was switched off but while the discharge was maintained with Ar buffer gas alone. The presence of N\(^{13}\)CO\(^-\) under these conditions likely comes about simply because isocyanic acid is metastable and persists in the cell for some time after its formation.

Because of the small change on the moment of inertia upon substitution of the C atom, a reliable prediction of the

![Figure 1. J = 19 ← 18 rotational transition of NCO\(^-\) with the background subtracted; the total integration time was 565 s for a scan at 3.24 MHz/s with a time constant of 3 ms.](https://doi.org/10.1021/acs.jpca.2c00313)
rotational constant of the C-13 isotopologue was calculated by scaling the theoretical B constant of N\textsuperscript{13}CO\textsuperscript{-} by the ratio of the theoretical and experimental rotational constants for the parent species. The first transition sought was the \( J = 14 \leftarrow 13 \) at 322.323 GHz, and indeed a discharge-dependent line was soon observed (see Figure 2). Nevertheless, the detection of additional harmonically related lines at higher and lower frequency, which was needed to confirm the identification, proved challenging because even modestly strong lines from unrelated species easily obscure the relative weak lines of N\textsuperscript{13}CO\textsuperscript{-}. For example, the \( J = 15 \leftarrow 14 \) of N\textsuperscript{13}CO\textsuperscript{-} lies very close in frequency to a strong transition of H\textsuperscript{13}CN and could not be measured.

2.2. Computational Details. The CFOUR program package\textsuperscript{25} has been employed to obtain an ab initio estimate of the equilibrium \( r_e \) structure and the vibrational corrections needed to derive the semiequilibrium equilibrium \( r_{eq} \) structure. The coupled-cluster method with inclusion of single and double excitations and perturbative inclusion of triple excitations, CCSD(T),\textsuperscript{23} has been used. For the electrons of the three atoms, correlation-consistent basis sets of Dunning (cc-pV\textsubscript{n}Z)\textsuperscript{24} have been applied, augmented by diffuse functions (aug-cc-pV\textsubscript{n}Z)\textsuperscript{25} for a proper description of the anion.

An automatized composite scheme implemented in CFOUR has been adopted for the geometry optimization.\textsuperscript{26} Briefly, the method first uses basis-set extrapolation at the Hartree–Fock (HF) level in which extrapolation of the energy gradient to the complete basis set limit is achieved by using three basis sets of the aug-cc-pV\textsubscript{n}Z hierarchy with \( n = T, Q, S \). Then, extrapolation of the correlation-energy contribution was performed with the two basis sets aug-cc-pVTZ and aug-cc-pVQZ. In the framework of the additivity scheme, the correlation energy gradient is added to the HF term. With the previous two terms obtained in the frozen-core (fc) approximation, a third energy gradient accounting for inner-shell correlation contributions has to be added, and for this purpose we performed all-electron and the corresponding fc calculations at the CCSD(T) level with the basis set aug-cc-pVTZ.

Vibrational corrections to the equilibrium rotational constants have been computed within a second-order vibra-
tional perturbation theory (VPT2) treatment using the full cubic force field together with the semidiagonal part of the quartic force field evaluated at the CCSD(T)/aug-cc-pV\textsubscript{n}Z level where three calculations with \( n = T, Q, S \) have been performed to check for accuracy. The presence of a Fermi resonance between the \( v_1 = 1 \) and \( v_2 = 2 \) states has been considered by removing the resonance denominator in the sum of the first-order vibration–rotation interaction constants.

3. RESULTS AND DISCUSSION

3.1. Rotational Spectra. Using the spectroscopic constants of Lattanzi et al.,\textsuperscript{15} we observed three higher-frequency rotational lines of the main isotopologue at 391.4, 414.14, and 437.4 GHz, with \( J = 16, 17, \) and 18 (see Table 1). Attempts to detect still higher frequency lines (e.g., the \( J = 20 \leftarrow 19 \) transition) was unsuccessful, in large part because of a rapid drop-off of the millimeter-wave power at shorter wavelengths. At high rotation quantum numbers, hyperfine structure due to the \( ^{14}N \) nucleus was blended in a single peak, and the unperturbed line frequency was recovered by a line shape analysis of the spectral profile modeled as a single Voigt profile.

Using the standard frequency expression of the rotational transition \( J + 1 \leftarrow J \)

\[
\nu_0 = 2B_0(J + 1) - 4D_J(J + 1)^3
\]

and the experimental transition frequencies, we determined \( B_0 \) and \( D_J \), the ground-state rotational and quartic centrifugal distortion constants, respectively, using a weighted least-squares procedure.\textsuperscript{27} Because hyperfine structure was resolved in the \( J = 1 \leftarrow 0 \) measurements in ref 15, the quadrupole coupling (eqQ) constant of the N nucleus (spin quantum number \( I = 1 \) can also be derived according to the expression

\[
\nu_{eq} = \nu_0 - eqQ[Y(J + 1, I, F') - Y(J, I, F)]
\]

where \( F \) is the quantum number associated with the total angular momentum \( \vec{F} = \vec{J} + \vec{I} \), and the Casimir function \( Y(J, I, F) \) is given by

![Figure 2. J = 14 ← 13 rotational transition of N\textsuperscript{13}CO\textsuperscript{-} with the background subtracted; the total integration time was 341 s for the average of six sweeps in the forward and reverse directions. The feature at lower frequency is from an unassigned molecule present in the glow discharge.](image-url)
Table 2. Spectroscopic Constants of NCO

| constants          | Lattanzi et al. | this work \(^{a}\) | correlation matrix |
|--------------------|-----------------|---------------------|-------------------|
| \(B_0\) (MHz)      | 11513.9683(8)   | 11513.9677(43)      | 1.000             |
| \(D_J\) (kHz)      | 4.561(2)        | 4.55907(87)         | 0.0885            |
| \(\sigma Q\) (MHz) |                  |                     | 1.000             |
| \(r_{nm}\) b (kHz) | 12.7            | 0.724               |

“Errors, as listed in the output of the SPFIT program,” are reported in parentheses in units of the last quoted digit; standard errors of the fit parameters are obtained multiplying by \(\sigma\). \(^{b}\)Rms error of residuals: \(\sqrt{\sum_{i=1}^{N_{\text{obs}}} \frac{(\text{residual}_i)^2}{\text{uncert}_i}}\). \(^{a}\)Fit standard deviation: \(\sqrt{\sum_{i=1}^{N_{\text{residual}}} (\text{residual}_i / \text{uncert}_i)^2 / \text{degrees of freedom}}\).

Table 3. Measured Rotational Transition Frequencies of \(\text{N}^{13}\text{CO}\)

| transition \(J' \leftarrow J\) | frequency \(^{a}\) (MHz) | residual (kHz) |
|-----------------------------|------------------------|---------------|
| 8 \leftarrow 7             | 184204.055             | 0             |
| 9 \leftarrow 8             | 207226.785             | 15            |
| 10 \leftarrow 9            | 230248.488             | 11            |
| 11 \leftarrow 10           | 253299.142             | 8             |
| 12 \leftarrow 11           | 276388.544             | 20            |
| 14 \leftarrow 13           | 322323.382             | 8             |

“The estimated 1σ uncertainty is 20 kHz.

determined in an analogus procedure as that used for the parent species sans the nitrogen hyperfine structure since the fundamental rotational transition of \(\text{N}^{13}\text{CO}\) was not measured in ref 15. Table 3 provides the residuals of the fit while the best-fit spectroscopic constants are given in Table 4 for \(\text{N}^{13}\text{CO}\).

3.2. Vibrational Corrections. Table 5 reports the contributions of zero-point vibration motion to the rotational constant for both \(\text{NCO}^-\) and \(\text{N}^{13}\text{CO}^-\) by using three correlation-consistent Dunning’s basis sets. The differences in the derived values for the three basis sets is quite small, amounting to at most 1.8%. A particularly good test of the accuracy of the calculations is the theoretical values of \(\alpha_3\), the largest vibration–rotation interaction constant, relative to the

Table 4. Spectroscopic Constants of \(\text{N}^{13}\text{CO}^-\) \(^{a}\)

| constant          | value                  | correlation matrix |
|-------------------|------------------------|-------------------|
| \(B_0\) (MHz)     | 11513.33735(117)       | 1.000             |
| \(D_J\) (kHz)     | 4.5620(42)             | −0.947            |
| \(r_{nm}\) b (kHz)| 12.3                   | 0.613             |

“Notes of Table 2 apply to this table as well.

Table 5. Calculated Vibrational Corrections for \(\text{NCO}^-\) and \(\text{N}^{13}\text{CO}^-\)

| basis set         | \(B_e - B_0\) (MHz) | \(\alpha_3\) (cm\(^{-1}\)) | \(\text{diff}^b\) (%) | \(B_e - B_0\) (MHz) |
|-------------------|---------------------|-----------------------------|-----------------------|---------------------|
| aug-cc-pVTZ       | 46.5658             | 0.00296761                  | −4.4                  | 45.9527             |
| aug-cc-pVQZ       | 46.6154             | 0.00297521                  | −1.8                  | 45.9990             |
| aug-cc-pVSZ       | 47.3810             | 0.00297935                  | −0.4                  | 46.7460             |

“Difference with respect to the experimental value of the vibration–rotation interaction constant \(\alpha_3 = 0.0029806(22)\) cm\(^{-1}\) from ref 16.

Table 6. Semiexperimential Equilibrium Rotational Constants of \(\text{NCO}^-\) and \(\text{N}^{13}\text{CO}^-\)

|              | \(\text{NCO}^-\) | \(\text{N}^{13}\text{CO}^-\) |
|--------------|-----------------|-----------------------------|
| \(B_0\) (MHz) | 11561.3488      | 11560.0833                  |
| rel accuracy\(^a\) | 6 × 10\(^{-6}\) |                             |

“Conservative value, estimated assuming an accuracy of 0.15% for vibrational corrections.

is derived from the experimental ground state constant, \(B_0\), plus the \(ab\) \(\text{initio}\) calculated vibrational correction, \(B_e - B_0\). Although the latter term contributes only for 0.4% to \(B_0\), its precision with respect to that of \(B_0\) is large and therefore dominates the resulting uncertainty of \(B_e\).

3.3. Semiexperimential Equilibrium Structure. Assuming a linear geometry for \(\text{NCO}^-\), the relationship between the moment of inertia and the two bond distances, \(r_{\text{NC}}\) and \(r_{\text{CO}}\), is the following (see Table 13.1 of ref 28):

\[
I = \frac{1}{m_N + m_C + m_O} \left[ m_N m_C r_{\text{NC}}^2 + m_C m_O r_{\text{CO}}^2 + m_N m_O (r_{\text{NC}} + r_{\text{CO}})^2 \right]
\]
where \( m \) denotes the mass of the atom. From the values of the moments of inertia of NCO\(^-\) and N\(^{13}\)CO\(^-\) it is thus possible to determine the two bond lengths. However, this solution is poorly constrained when substitution is very close to the center of mass, and the value of the moment of inertia changes very little. Nevertheless, the stability of the solution to the center of mass, and the value of the moment of inertia of NCO\(^-\) is poorly constrained when substitution is very close to determine the two bond lengths. However, this semiexperimental method is the most accurate in calculating bond lengths of a triatomic linear molecule containing three atoms, presumably because of its ability to treat electron correlation effects. In addition, basis-set extrapolation, combined with additivity schemes, improves accuracy in geometry optimization. For NCO\(^-\), differences between the theoretical and semiexperimental bond lengths are about 1 mÅ or less: −1.3 × 10⁻³ and +0.3 × 10⁻³ Å for \( r_{NC} \) and \( r_{CO} \), respectively.

4. CONCLUSIONS

High-resolution spectroscopic observations of the cyanate anion are limited to the ground-state rotational spectrum and the \( \nu_2 \) fundamental and corresponding bending and stretching hot bands of NCO\(^-\). In addition to higher-frequency rotational measurements of the isotope species, this study adds the rotational spectrum of the isotopologue N\(^{13}\)CO\(^-\) to this body of work. From the two rotational constants, the two bond lengths of this linear triatomic can be derived. Going beyond a vibrationally averaged structure, quantum-chemical calculations have to be performed to obtain the vibrational contributions to the \( \beta \) constants, which, otherwise, would require extensive spectroscopic work. The molecular structure retrieved by such a combination of experimental and theoretical data, termed the semiexperimental structure, is the main result of this paper. The accuracy of the two derived bond distances is estimated in the order of 5 × 10⁻⁴ Å, based on an assumed underestimate of the vibrational correction to the rotational constants of about 0.15%. In addition, geometry optimization with the CCSD(T) method and the use of a composite scheme, including additivity and basis-set extrapolation techniques, are very accurate for this simple, closed-shell linear molecule.

Table 7. Molecular Structure of NCO\(^-\)

| level of theory | bond lengths (Å) | N–C | C–O |
|----------------|------------------|-----|-----|
| semiexperimental theory |                    |     |     |
| Botschwa et al., 1995 CCSD(T)/138cGTOs | 1.9110 | 1.22625 |
| Pak et al., 1997 CCSD(T)/6s5p3d2f | 1.9134 | 1.2306 |
| Prasad, 2004 f-CASSCF/6s4p2df | 1.199 | 1.231 |
| Borsboom et al., 2004 CCSD(T)/aug-cc-pV5Z | 1.1928 | 1.2293 |
| this work CCSD(T)/aug-cc-pV5Z | 1.1898 | 1.2266 |

“Reference 17, with all electrons correlated. †Reference 18, 150 cGTOs. ‡Reference 30. ††Reference 31. For the composite scheme employed see text.”

and those calculated theoretically are closely consistent. Based on these comparisons, the bond distances derived from the semiexperimental \( B_e \) constants appear to be determined to high accuracy.

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