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# Comparison of Spectroscopic Strategies to Determine Molecular Geometries and the Impact of Nuclear versus Atomic Masses: The Example of $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$ 

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#### Abstract

We compare a recently proposed mixed experimental/theoretical procedure for the derivation of molecular equilibrium structures with several commonly used spectroscopic approaches using experimental data for several isotopologues. We also examine the sensitivity of the results from these approaches to the replacement of the commonly employed atomic masses with nuclear masses. This point is of particular importance for ionic species like $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$which serve as numerical reference cases. The scatter of molecular equilibrium geometries derived by different approaches is found to exceed stated statistical uncertainties by about an order of magnitude.


## I. INTRODUCTION

Recently, standard spectroscopic parameters for the isotopologues of $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$were determined by means of a theoretical two-step procedure. ${ }^{1}$ In the first step, the $a b$ initio computed threedimensional CCSD $(T) / c c-p V Q Z$ potential energy surface of Mladenović and Schmatz ${ }^{2}$ for the isomerizing system $\mathrm{HCO}^{+} / \mathrm{HOC}^{+}$was used to obtain the energies of the rotational levels up to $J=15$ in the ground vibrational state and in the singly excited vibrational $\nu_{i}$ states for $i=1-3$. The computed rovibrational energies were fitted to the conventional spectroscopic expressions. The rotational constants $B_{i}^{\text {th }}$ obtained in this manner for $i=0-3$ deviate by approximately $0.005 \mathrm{~cm}^{-1}(200 \mathrm{MHz})$ from their experimentally derived counterparts $B_{i}^{\exp }$. This disagreement was corrected in the second step. Ascribing the difference between $B_{i}^{\text {th }}$ and $B_{i}^{\exp }$ to the restricted accuracy of the equilibrium bond lengths, a new set of structural parameters was derived by combining the experimental ground-state rotational constants $B_{0}^{\exp }$ with theoretical rotation-vibration corrections $\Delta B_{0}^{\text {th }}=B_{e}^{\text {th }}-B_{0}^{\text {th }}$ computed variationally (beyond a perturbational approach), where $B_{e}^{\text {th }}$ is the equilibrium rotational constant for the $\operatorname{CCSD}(\mathrm{T}) / \mathrm{cc}-\mathrm{pVQZ}$ potential energy surface. The procedure introduced in Ref. 1, hereafter Paper I, led to estimates of the spectroscopic constants and equilibrium bond distances in excellent agreement with values derived from experiments.

All stable isotopologues of $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$involving $\mathrm{H}, \mathrm{D},{ }^{16} \mathrm{O},{ }^{17} \mathrm{O},{ }^{18} \mathrm{O},{ }^{12} \mathrm{C}$, and ${ }^{13} \mathrm{C}$ were considered in Paper I, in total 24 molecular cations. The rovibrational calculations and the equilibrium structure determinations were carried out using atomic masses. This is common practice in nuclear dynamics computations, but there is an obvious problem for ionic species.

Within the Born-Oppenheimer approximation, interacting nuclei are described by a mass-independent potential energy (hyper)surface, providing the energy of the electronic subsystem, so that a single potential energy surface is used for all isotopic variants of a molecular system under investigation. The equilibrium geometry,
defined as usual as the minimum of the potential energy surface, is thus mass-independent and unique for all isotopologues. The Born-Oppenheimer approximation and its electronic-structure implementations rely on a perfect separation of the electronic and nuclear coordinates. This, in turn, implies that the nuclear (rotationvibrational) motion should, rigorously speaking, involve the nuclear masses. Whereas the use of atomic masses in connection with nuclear dynamics computations is considered to be a pragmatic approach, the use of nuclear masses is conceptually more correct and the only sound basis for systematic improvements.
The purpose of the present paper is to investigate the influence of the nuclear versus atomic masses on the spectroscopic and structural parameters of the molecular cations $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$. Since the masses explicitly enter in the process of extracting the geometric parameters from the observed spectral (rotational) transitions, this issue is of relevance also for estimating equilibrium structures from experimental ground-state rotational constants. In practice, rotational transitions observed for linear triatomic molecules are compactly represented by a polynomial expansion in terms of $J(J+1)$ and $K^{2}$, where $J$ and $K$ are the quantum numbers specifying the state of the total rotational angular momentum and of its projection onto the body-fixed (molecular) $z$ axis, respectively. The frequency $\nu$ of the rotational transition $J \rightarrow J+1$ in a vibrational $\Sigma$ state $v$ is, for instance, given by

$$
\begin{align*}
\nu= & 2 B_{v}(J+1)-4 D_{v}(J+1)^{3} \\
& +2 H_{v}(J+1)^{3}\left(3 J^{2}+6 J+4\right) \ldots, \tag{1}
\end{align*}
$$

where $B_{v}, D_{v}, H_{v}$, and so on, are expansion parameters appropriate for the vibrational state $v$. The dominant term in Eq. (1) is the first term involving the rotational constant $B_{v}$. For a given geometric arrangement or a rigid body, the rotational constant is a quantity wellestablished from the mathematical and physical point of view. In the actual circumstances of rotating and vibrating systems, we do, however, encounter problems of both conceptual and practical nature. ${ }^{3-8}$ In the present work, we revisit some of these issues on the example of $\mathrm{HCO}^{+}$

TABLE I: Atomic $m_{a}$ and nuclear $m_{n}$ masses in the unified atomic mass unit $u .{ }^{a}$ The reduced masses $\mu_{r}, \mu_{R}$ (in u) for the main form of $\mathrm{HCO}^{+}$are calculated by means of Eq. (8).

| species | $m_{a}$ | $m_{n}$ |
| :---: | ---: | ---: |
| H | 1.007825035 | 1.0072765 |
| D | 2.014101779 | 2.0135532 |
| ${ }^{12} \mathrm{C}$ | 12.0 | 11.9967085 |
| ${ }^{13} \mathrm{C}$ | 13.003354826 | 13.0000633 |
| ${ }^{16} \mathrm{O}$ | 15.99491463 | 15.9905260 |
| ${ }^{18} \mathrm{O}$ | 17.99916030 | 17.9947717 |
| $\mu_{r}$ | 6.856209 | 6.854328 |
| $\mu_{R}$ | 0.972804 | 0.972283 |

${ }^{a}$ The atomic masses are taken from Ref. 9.
and $\mathrm{HOC}^{+}$. We first give a brief overview of the theoretical approach which we employed (Section II). Our quantum-mechanical calculations provide a consistent set of data, which we use to study the mass effect on spectral and geometric parameters (Section II A). Different commonly used approaches to obtain information about the equilibrium structure are considered (Section III), leading us to a certain number of conclusions (Section IV).

## II. SPECTRAL AND GEOMETRIC PARAMETERS

The theoretical approach of Paper I is used in combination with nuclear masses in this work. Masses are given in the unified atomic mass unit $u$, which stands for $\mathrm{u}=\mathrm{m}_{a}\left({ }^{12} \mathrm{C}\right) / 12=10^{-3} \mathrm{~kg} \mathrm{~mol}^{-1} / \mathrm{N}_{A}$, where $\mathrm{m}_{a}\left({ }^{12} \mathrm{C}\right)$ is the mass of the atom ${ }^{12} \mathrm{C}$ and $N_{A}$ Avogadro's number. ${ }^{9}$ The atomic (nuclear plus electronic) masses $m_{a}$ and the nuclear masses $m_{n}$ of the isotopes of hydrogen, carbon, and oxygen are summarized in Table I for convenience. A glance at Table I shows that $m_{n}$ are approximately $5 \times 10^{-4}$ to $5 \times 10^{-3}$ u $(0.05 \%)$ smaller than $m_{a}$. In our computations, the nuclear masses are internally evaluated from the atomic masses $m_{a}$ provided in Table I. The conversion factor used for the electron mass is 1822.888515 , so that $m_{e}=1 / 1822.888515 \mathrm{u}$.

In our present analysis, we include only those isotopologues of $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$, for which experimental values of the ground-state rotational constant $B_{0}^{\exp }$ are known. This is the case for eight substituted isotopic forms of $\mathrm{HCO}^{+}\left(\mathrm{HCO}^{+},{ }^{10} \mathrm{HC}^{17} \mathrm{O}^{+},{ }^{11} \mathrm{HC}^{18} \mathrm{O}^{+},{ }^{12}\right.$ $\mathrm{H}^{13} \mathrm{CO}^{+},,^{13} \quad \mathrm{H}^{13} \mathrm{C}^{18} \mathrm{O}^{+},{ }^{14} \quad \mathrm{DCO}^{+},{ }^{15} \quad \mathrm{DC}^{18} \mathrm{O}^{+},{ }^{16}$ and $\mathrm{D}^{13} \mathrm{CO}^{+13}$ ) and four isotopologues of $\mathrm{HOC}^{+}\left(\mathrm{HOC}^{+},{ }^{17}\right.$ $\mathrm{H}^{18} \mathrm{OC}^{+},{ }^{18} \mathrm{HO}^{13} \mathrm{C}^{+},{ }^{18,19}$ and $\left.\mathrm{DOC}^{+17}\right)$.

## squeezetable

The rovibrational energies computed using the nuclear masses for the total rotational angular momentum up to $J=15$ are fitted to appropriate spectroscopic formu-
lae, following the procedure of Paper I. To facilitate the comparison, the same effective spectroscopic Hamiltonians are used for the atomic-mass and nuclear-mass cases for each of the species studied. The spectroscopic parameters obtained in the fits for the vibrational ground state and for the singly excited $\nu_{1}, \nu_{2}$, and $\nu_{3}$ states are listed in Tables II and III for the isotopic variants of $\mathrm{HCO}^{+}$and in Table IV for the isotopic variants of $\mathrm{HOC}^{+}$. There, $T_{v}$ is the term energy and $B_{v}$ the effective rotational constant for the vibrational state $v$, where we use $v=0-3$ to denote the vibrational states $\left(0,0^{0}, 0\right)$, $\left(1,0^{0}, 0\right),\left(0,1^{1}, 0\right)$, and $\left(0,0^{0}, 1\right)$, respectively. The centrifugal distortion contribution is expressed in terms of the quartic centrifugal distortion constant $D_{v}$ and higher order constants, such as $H_{v}$ (sextic), $L_{v}$ (octic), and so on. The $\ell$-type doubling contribution is described by the $\ell$-type doubling constant $q_{v}$ and the parameters $q_{v}^{J}, q_{v}^{J J}$, and so on for its centrifugal distortion corrections. The results obtained using the atomic masses are taken from Paper I. The equilibrium rotational constants and effective rotational constants computed using atomic masses are denoted by $B_{e}^{a}$ and $B_{v}^{a}$, respectively. Qualities of the fits obtained in the atomic-mass and nuclear-mass cases are comparable, as seen by similar fit standard deviations $\sigma$ of approximately 10 Hz in Tables II-IV.

The replacement of the atomic masses with the nuclear masses affects the effective rotational constants $B_{0}, B_{1}, B_{2}, B_{3}$ by $9-15 \mathrm{MHz}$, as seen by inspection of the difference $B_{v}-B_{v}^{a}$ in Tables II-IV. There we also see that the difference $B_{e}-B_{e}^{a}$ is equal to $9-15 \mathrm{MHz}$, too, where $B_{e}$ and $B_{e}^{a}$ are the equilibrium rotational constants computed for the $\operatorname{CCSD}(\mathrm{T}) / \mathrm{cc}-\mathrm{pVQZ}$ equilibrium structure using nuclear masses and atomic masses, respectively. This thus shows that the difference $B_{e}-B_{e}^{a}$ is a major contributor to the difference $B_{v}-B_{v}^{a}$ for $v=0-3$. The centrifugal distortion constants $D_{v}$ are approximately $0.05 \mathrm{kHz}(0.05 \%)$ larger than $D_{v}^{a}$. The $\ell$-type doubling constants $q_{v}$ are larger than $q_{v}^{a}$ by 0.1 MHz for the isotopic forms of $\mathrm{HCO}^{+}$and by 0.2 MHz for the isotopic forms of $\mathrm{HOC}^{+}$, which represent a change of approximately $0.05 \%$. The vibration-rotation interaction constants $\alpha_{v}=B_{0}-B_{v}$ are larger than $\alpha_{v}^{a}=B_{0}^{a}-B_{v}^{a}$ by at most $0.2 \mathrm{MHz}(0.05 \%)$. Finally, the wavenumbers of the $\nu_{1}, \nu_{2}, \nu_{3}$ vibrations are increased by at most $1 \mathrm{~cm}^{-1}$ upon replacement of atomic masses with nuclear masses.

After publication of Paper I, we have learned that Warner in his Ph.D. thesis ${ }^{14}$ had measured the rotational transitions $J=2 \rightarrow 3$ and $J=3 \rightarrow 4$ in the ground vibrational state of $\mathrm{HOC}^{+}, \mathrm{H}^{18} \mathrm{OC}^{+}$, and $\mathrm{HO}^{13} \mathrm{C}^{+}$, from which he had derived ( $\left.B_{0}^{\text {exp }}, D_{0}^{\text {exp }}\right)$ of $(44743.943,116.75)$, (43 305.969,108.32), (42 876.559,107.25) in (MHz,kHz) for these systems, respectively. These results are in good agreement with the experimental data of Gudeman et al. ${ }^{18,19}$ and Amano and Maeda, ${ }^{17}$ which were used in Paper I (Table III) and here in Table IV. Warner had also measured the rotational transition $J=2 \rightarrow 3$ at 248459.964 MHz in the ground vibrational state of $\mathrm{H}^{18} \mathrm{O}^{13} \mathrm{C}^{+}$. From this value using Eq. (1), one calcu-

TABLE II: Spectroscopic parameters derived for the isotopic variants of $\mathrm{HCO}^{+}$using nuclear masses. Values shown in brackets were obtained in Paper I using atomic masses.

| Parameter | $\mathrm{H}^{12} \mathrm{C}^{16} \mathrm{O}^{+}$ | $\mathrm{H}^{12} \mathrm{C}^{17} \mathrm{O}^{+}$ | $\mathrm{H}^{12} \mathrm{C}^{18} \mathrm{O}^{+}$ | $\mathrm{H}^{13} \mathrm{C}^{16} \mathrm{O}^{+}$ | $\mathrm{H}^{13} \mathrm{C}^{18} \mathrm{O}^{+}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| $B_{e} / \mathrm{MHz}$ | 44635.23 | 43565.63 | 42614.46 | 43412.78 | 41361.54 |
| $B_{e}-B_{e}^{a} / \mathrm{MHz}$ | 15.22 | 14.58 | 14.02 | 14.35 | 13.13 |
| $B_{0} / \mathrm{MHz}$ | 44391.37 | 43330.07 | 42386.20 | 43178.57 | 41142.93 |
| $B_{0}-B_{0}^{a} / \mathrm{MHz}$ | 15.09 | 14.45 | 13.91 | 14.23 | 13.02 |
| $D_{0} / \mathrm{kHz}$ | 82.01 [81.95] | 78.15 [78.10] | 74.81 [74.76] | 77.64 [77.59] | 70.56 [70.51] |
| $H_{0} / \mathrm{Hz}$ | 0.071 [0.071] | 0.065 [0.065] | 0.061 [0.061] | 0.063 [0.063] | 0.053 [0.053] |
| $\sigma_{0} / \mathrm{Hz}$ | 9.70 [8.45] | 8.93 [8.81] | 9.34 [7.63] | 8.17 [8.93] | 5.82 [6.90] |
| $B_{0}^{\text {est }} / \mathrm{MHz}$ | 44594.46 [44594.42] | 43528.43 [43528.77] | 42580.37 [42580.99] | 43376.53 [43377.18] | 41331.84 [41333.14] |
| $B_{0}^{\text {exp }} / \mathrm{MHz}$ | $44594.42866(16)^{\text {a }}$ | $43528.9253(19)^{\text {b }}$ | $42581.21(4)^{\text {c }}$ | $43377.3019(17)^{\text {d }}$ | $41333.587(5)^{\text {e }}$ |
| $T_{2} / \mathrm{cm}^{-1}$ | 829.44 [829.25] | 828.40 [828.21] | 827.48 [827.29] | 821.98 [821.80] | 820.01 [819.82] |
| $B_{2} / \mathrm{MHz}$ | 44471.56 | 43407.86 | 42461.87 | 43249.54 | 41209.65 |
| $B_{2}-B_{2}^{a} / \mathrm{MHz}$ | 15.11 | 14.47 | 13.93 | 14.25 | 13.03 |
| $D_{2} / \mathrm{kHz}$ | 83.62 [83.57] | 79.67 [79.62] | 76.24 [76.19] | 79.06 [79.01] | 71.81 [71.76] |
| $\mathrm{H}_{2} / \mathrm{Hz}$ | 0.084 [0.084] | 0.077 [0.077] | 0.071 [0.071] | 0.074 [0.074] | 0.062 [0.062] |
| $\sigma_{2} / \mathrm{Hz}$ | 12.33 [11.45] | 10.17 [9.81] | 9.02 [8.45] | 9.60 [9.64] | 7.95 [8.02] |
| $q_{2} / \mathrm{MHz}$ | 209.95 [209.85] | 200.30 [200.21] | 191.90 [191.82] | 200.44 [200.36] | 182.47 [182.40] |
| $q_{2}^{J} / \mathrm{kHz}$ | -1.80 [-1.79] | -1.67 [-1.67] | -1.56 [-1.56] | -1.65 [-1.65] | -1.42 [-1.42] |
| $q_{2}^{J J} / \mathrm{Hz}$ | 0.024 [0.024] | 0.022 [0.022] | 0.019 [0.019] | 0.020 [0.020] | 0.016 [0.016] |
| $\sigma_{q} / \mathrm{Hz}$ | 9.68 [9.14] | 9.65 [9.69] | 7.68 [6.15] | 7.89 [9.07] | 6.68 [6.39] |
| $\alpha_{2} / \mathrm{MHz}$ | -80.19 [-80.17] | -77.79 [-77.77] | -75.68 [-75.66] | -70.97 [-70.95] | -66.72 [-66.70] |
| $B_{2}^{\text {est }} / \mathrm{MHz}$ | 44674.65 [44674.59] | 43606.24 [43606.54] | 42656.06 [42656.65] | 43447.52 [43448.13] | 41398.57 [41399.84] |
| $B_{2}^{\exp } / \mathrm{MHz}$ | $44677.1489(18)^{\text {d }}$ |  | $42659.160(8)^{\text {e }}$ | $43450.524(8)^{\mathrm{e}}$ | $41402.389(8)^{\mathrm{e}}$ |
| $T_{3} / \mathrm{cm}^{-1}$ | 2179.44 [2179.09] | 2152.65 [2152.31] | 2128.52 [2128.19] | 2145.68 [2145.35] | 2092.82 [2092.51] |
| $B_{3} / \mathrm{MHz}$ | 44098.63 | 43047.00 | 42111.71 | 42895.00 | 40877.89 |
| $B_{3}-B_{3}^{a} / \mathrm{MHz}$ | 14.94 | 14.30 | 13.76 | 14.08 | 12.89 |
| $D_{3} / \mathrm{kHz}$ | 82.10 [82.04] | 78.24 [78.18] | 74.88 [74.83] | 77.74 [77.69] | 70.64 [70.59] |
| $\mathrm{H}_{3} / \mathrm{Hz}$ | 0.070 [0.070] | 0.065 [0.065] | 0.060 [0.060] | 0.062 [0.062] | 0.053 [0.053] |
| $\sigma_{3} / \mathrm{Hz}$ | 7.88 [7.51] | 6.60 [7.25] | 6.40 [5.62] | 7.44 [7.10] | 5.39 [5.57] |
| $\alpha_{3} / \mathrm{MHz}$ | 292.75 [292.59] | 283.07 [282.91] | 274.49 [274.34] | 283.57 [283.42] | 265.04 [264.90] |
| $B_{3}^{\text {est }} / \mathrm{MHz}$ | 44301.71 [44301.83] | 43245.38 [43245.85] | 42305.90 [42306.65] | 43092.98 [43093.76] | 41066.81 [41068.24] |
| $B_{3}^{\exp } / \mathrm{MHz}$ | $44299.8687(78)^{\text {d }}$ |  | 42305.024(8) ${ }^{\text {e }}$ | $43091.852(8)^{\mathrm{e}}$ |  |
| $T_{1} / \mathrm{cm}^{-1}$ | 3086.29 [3085.58] | 3083.79 [3083.08] | 3081.67 [3080.96] | 3063.70 [3062.99] | 3059.76 [3059.05] |
| $B_{1} / \mathrm{MHz}$ | 44038.81 | 42986.42 | 42051.39 | 42850.77 | 40834.08 |
| $B_{1}-B_{1}^{a} / \mathrm{MHz}$ | 14.94 | 14.31 | 13.77 | 14.09 | 12.91 |
| $D_{1} / \mathrm{kHz}$ | 81.27 [81.22] | 77.48 [77.42] | 74.16 [74.12] | 77.03 [76.98] | 69.98 [69.93] |
| $H_{1} / \mathrm{Hz}$ | 0.068 [0.068] | 0.065 [0.065] | 0.061 [0.061] | 0.062 [0.062] | 0.054 [0.054] |
| $\sigma_{1} / \mathrm{Hz}$ | 15.18 [14.70] | 13.25 [13.28] | 10.83 [9.41] | 9.88 [9.38] | 8.97 [9.49] |
| $\alpha_{1} / \mathrm{MHz}$ | 352.57 [352.41] | 343.65 [343.51] | 334.81 [334.67] | 327.80 [327.66] | 308.86 [308.74] |
| $B_{1}^{\text {est }} / \mathrm{MHz}$ | 44241.89 [44242.01] | 43184.80 [43185.26] | 42245.58 [42246.32] | 43048.75 [43049.52] | 41023.00 [41024.39] |
| $B_{1}^{\text {exp }} / \mathrm{MHz}$ | $44240.53309(56)^{\text {e }}$ |  | $42244.862(8)^{\mathrm{e}}$ | $43048.158(8)^{\text {e }}$ |  |
| ${ }^{(n)} B_{e}^{\text {est }}\left[{ }^{(a)} B_{e}^{\text {est }}\right] / \mathrm{MHz}$ | 44838.30 [44838.15] | 43763.99 [43764.21] | 42808.63 [42809.14] | 43610.75 [43611.28] | 41550.44 [41551.63] |
| ${ }^{(a)} B_{e}^{\exp } / \mathrm{MHz}$ | $44840.1330(161)^{\mathrm{g}}$ |  | $42811.0181(161)^{\mathrm{g}}$ | $43613.2505(161)^{\mathrm{g}}$ |  |
| $S_{0}\left[S_{0}^{a}\right] / \mathrm{MHz}$ | 242.47 [242.33] | 235.57 [235.44] | 228.97 [228.85] | 234.72 [234.59] | 220.23 [220.12] |
| $\Delta B_{0} / \mathrm{MHz}$ | 243.86 [243.73] | 235.56 [235.44] | 228.27 [228.15] | 234.22 [234.10] | 218.61 [218.50] |
| $\Delta B_{0}-S_{0} / \mathrm{MHz}$ | 1.39 | -0.01 | -0.70 | -0.50 | -1.62 |

[^0]TABLE III: Spectroscopic parameters derived for the isotopic variants of $\mathrm{DCO}^{+}$using nuclear masses. Values shown in brackets were obtained in Paper I using atomic masses.

| Parameter | $\mathrm{D}^{12} \mathrm{C}^{16} \mathrm{O}^{+}$ | $\mathrm{D}^{12} \mathrm{C}^{18} \mathrm{O}^{+}$ | $\mathrm{D}^{13} \mathrm{C}^{16} \mathrm{O}^{+}$ |
| :---: | :---: | :---: | :---: |
| $B_{e} / \mathrm{MHz}$ | 36035.19 | 34424.09 | 35379.38 |
| $B_{e}-B_{e}^{a} / \mathrm{MHz}$ | 9.86 | 9.02 | 9.52 |
| $B_{0} / \mathrm{MHZ}$ | 35861.59 | 34261.63 | 35210.56 |
| $B_{0}-B_{0}^{a} / \mathrm{MHz}$ | 9.78 | 8.96 | 9.45 |
| $D_{0} / \mathrm{kHz}$ | 55.24 [55.21] | 50.16 [50.13] | 52.88 [52.85] |
| $H_{0} / \mathrm{Hz}$ | 0.050 [0.050] | 0.043 [0.043] | 0.046 [0.046] |
| $\sigma_{0} / \mathrm{Hz}$ | 6.76 [6.56] | 4.99 [5.54] | 6.19 [6.18] |
| $B_{0}^{\text {est }} / \mathrm{MHz}$ | 36020.07 [36019.81] | 34413.35 [34413.66] | 35366.58 [35366.70] |
| $B_{0}^{\exp } / \mathrm{MHz}$ | $36019.76765(14)^{\text {a }}$ | $34413.78556(18)^{\text {a }}$ | $35366.70968(21)^{\text {a }}$ |
| $T_{2} / \mathrm{cm}^{-1}$ | 666.39 [666.30] | 663.90 [663.81] | 656.94 [656.85] |
| $B_{2} / \mathrm{MHz}$ | 35956.82 | 34352.11 | 35297.88 |
| $B_{2}-B_{2}^{a} / \mathrm{MHz}$ | 9.83 | 9.00 | 9.48 |
| $D_{2} / \mathrm{kHz}$ | 56.85 [56.82] | 51.59 [51.56] | 54.33 [54.30] |
| $H_{2} / \mathrm{Hz}$ | 0.063 [0.063] | 0.054 [0.054] | 0.058 [0.058] |
| $\sigma_{2} / \mathrm{Hz}$ | 12.75 [12.29] | 8.55 [8.19] | 8.30 [7.76] |
| $q_{2} / \mathrm{MHz}$ | 169.63 [169.56] | 155.42 [155.36] | 165.82 [165.76] |
| $q_{2}^{J} / \mathrm{kHz}$ | -1.32 [-1.32] | -1.15 [-1.15] | -1.25 [-1.25] |
| $q_{2}^{J J} / \mathrm{Hz}$ | 0.023 [0.023] | 0.019 [0.019] | 0.021 [0.021] |
| $\sigma_{q} / \mathrm{Hz}$ | 5.18 [5.05] | 7.50 [7.81] | 9.66 [9.44] |
| $\alpha_{2} / \mathrm{MHz}$ | -95.22 [-95.18] | -90.48 [-90.45] | -87.32 [-87.29] |
| $B_{2}^{\text {est }} / \mathrm{MHz}$ | 36115.30 [36115.00] | 34503.83 [34504.10] | 35453.90 [35453.98] |
| $B_{2}^{\exp } / \mathrm{MHz}$ | $36116.66634(80)^{\text {b }}$ |  |  |
| $T_{3} / \mathrm{cm}^{-1}$ | 1901.07 [1900.81] | 1870.99 [1870.75] | 1894.10 [1893.85] |
| $B_{3} / \mathrm{MHz}$ | 35656.03 | 34071.61 | 35012.79 |
| $B_{3}-B_{3}^{a} / \mathrm{MHz}$ | 9.70 | 8.89 | 9.36 |
| $D_{3} / \mathrm{kHz}$ | 55.21 [55.18] | 50.11 [50.09] | 52.84 [52.81] |
| $H_{3} / \mathrm{Hz}$ | 0.049 [0.049] | 0.043 [0.043] | 0.044 [0.044] |
| $\sigma_{3} / \mathrm{Hz}$ | 2.83 [2.44] | 2.64 [3.00] | 2.66 [2.54] |
| $\alpha_{3} / \mathrm{MHz}$ | 205.56 [205.48] | 190.02 [189.94] | 197.77 [197.69] |
| $B_{3}^{\text {est }} / \mathrm{MHz}$ | 35814.51 [35814.34] | 34223.33 [34223.71] | 35168.81 [35169.01] |
| $B_{3}^{\exp } / \mathrm{MHz}$ | $35813.3618(31)^{\text {b }}$ |  |  |
| $T_{1} / \mathrm{cm}^{-1}$ | 2580.80 [2580.46] | 2554.57 [2554.23] | 2529.79 [2529.47] |
| $B_{1} / \mathrm{MHz}$ | 35636.66 | 34010.18 | 34844.93 |
| $B_{1}-B_{1}^{a} / \mathrm{MHz}$ | 9.74 | 8.87 | 9.31 |
| $D_{1} / \mathrm{kHz}$ | 46.08 [46.05] | 40.54 [40.52] | 40.32 [40.28] |
| $H_{1} / \mathrm{Hz}$ | -2.444 [-2.441] | 0.039 [0.039] | -4.001 [-4.004] |
| $L_{1} / \mathrm{mHz}$ | 0.328 [0.326] | -0.198 [-0.196] | 1.508 [1.509] |
| $K_{1} / \mu \mathrm{Hz}$ | 0.037 [0.037] | 0.001 [-0.002] | -0.484 [-0.484] |
| $\sigma_{1} / \mathrm{Hz}$ | 4.12 [3.92] | 8.33 [10.12] | 16.47 [15.73] |
| $\alpha_{1} / \mathrm{MHz}$ | 224.94 [224.89] | 251.45 [251.36] | 365.62 [365.50] |
| $B_{1}^{\text {est }} / \mathrm{MHz}$ | 35795.13 [35794.93] | 34161.90 [34162.30] | 35000.95 [35001.20] |
| $B_{1}^{\exp } / \mathrm{MHz}$ | $35792.3325(46)^{\text {c }}$ |  |  |
| ${ }^{(n)} B_{e}^{\text {est }}\left[{ }^{(a)} B_{e}^{\text {est }}\right] / \mathrm{MHz}$ | 36193.65 [36193.34] | 34575.79 [34576.06] | 35535.38 [35535.45] |
| ${ }^{(a)} B_{e}^{\exp } / \mathrm{MHz}$ | $36194.3538(79)^{\text {d }}$ |  |  |
| $S_{0}\left[S_{0}^{a}\right] / \mathrm{MHz}$ | 120.03 [120.00] | 130.25 [130.21] | 194.37 [194.30] |
| $\Delta B_{0} / \mathrm{MHz}$ | 173.60 [173.53] | 162.46 [162.40] | 168.82 [168.75] |
| $\Delta B_{0}-S_{0} / \mathrm{MHz}$ | 53.57 | 32.21 | -25.55 |

[^1]TABLE IV: Spectroscopic parameters derived for $\mathrm{H}^{16} \mathrm{O}^{12} \mathrm{C}^{+}, \mathrm{H}^{18} \mathrm{O}^{12} \mathrm{C}^{+}, \mathrm{H}^{16} \mathrm{O}^{13} \mathrm{C}^{+}$, and $\mathrm{D}^{16} \mathrm{O}^{12} \mathrm{C}^{+}$using nuclear masses. Values shown in brackets were obtained in Paper I using atomic masses.

| Parameter | $\mathrm{H}^{16} \mathrm{O}^{12} \mathrm{C}^{+}$ | $\mathrm{H}^{18} \mathrm{O}^{12} \mathrm{C}^{+}$ | $\mathrm{H}^{16} \mathrm{O}^{13} \mathrm{C}^{+}$ | $\mathrm{D}^{16} \mathrm{O}^{12} \mathrm{C}^{+}$ |
| :---: | :---: | :---: | :---: | :---: |
| $B_{e} / \mathrm{MHz}$ | 44567.75 | 43156.82 | 42709.61 | 37877.31 |
| $B_{e}-B_{e}^{a} / \mathrm{MHz}$ | 14.43 | 13.47 | 13.33 | 10.38 |
| $B_{0} / \mathrm{MHz}$ | 44525.06 | 43091.57 | 42665.27 | 38024.35 |
| $B_{0}-B_{0}^{a} / \mathrm{MHz}$ | 14.31 | 13.35 | 13.23 | 10.42 |
| $D_{0} / \mathrm{kHz}$ | 113.83 [113.76] | 106.02 [105.96] | 104.21 [104.14] | 93.64 [93.59] |
| $H_{0} / \mathrm{Hz}$ | 0.076 [0.760] | 0.579 [0.579] | 0.619 [0.611] | 2.306 [2.305] |
| $L_{0} / \mathrm{mHz}$ | -0.046 [-0.046] | -0.032 [-0.032] | -0.034 [-0.034] | -0.245 [-0.245] |
| $K_{0} / \mu \mathrm{Hz}$ |  |  |  | 0.025 [0.026] |
| $\sigma_{0} / \mathrm{Hz}$ | 3.40 [4.30] | 3.32 [3.77] | 3.53 [3.41] | 0.74 [0.88] |
| $B_{0}^{\text {est }} / \mathrm{MHz}$ | 44744.06 [44743.93] | 43305.20 [43305.74] | 42875.92 [42876.36] | 38193.18 [38193.20] |
| $B_{0}^{\exp } / \mathrm{MHz}$ | $44743.9141(35)^{\text {a }}$ | $43305.969(35)^{\text {b }}$ | $42876.559(8)^{\text {b }}$ | $38193.1984(18)^{\text {a }}$ |
| $T_{2} / \mathrm{cm}^{-1}$ | 243.67 [243.60] | 241.15 [241.08] | 243.30 [243.23] | 176.60 [176.57] |
| $B_{2} / \mathrm{MHz}$ | 44719.37 | 43252.36 | 42845.08 | 38349.08 |
| $B_{2}-B_{2}^{a} / \mathrm{MHz}$ | 14.32 | 13.33 | 13.22 | 10.54 |
| $D_{2} / \mathrm{kHz}$ | 116.23 [116.15] | 107.84 [107.78] | 106.39 [106.33] | 100.69 [100.63] |
| $\mathrm{H}_{2} / \mathrm{Hz}$ | 0.284 [0.284] | 0.177 [0.177] | 0.234 [0.235] | 1.738 [1.734] |
| $L_{2} / \mathrm{mHz}$ | 0.028 [0.028] | 0.026 [0.026] | 0.021 [0.021] | -0.051 [-0.049] |
| $K_{2} / \mu \mathrm{Hz}$ |  |  |  | -0.011 [-0.012] |
| $\sigma_{2} / \mathrm{Hz}$ | 8.90 [8.38] | 6.48 [6.46] | 6.16 [6.72] | 0.77 [1.02] |
| $q_{2} / \mathrm{MHz}$ | 517.25 [517.07] | 489.50 [489.34] | 475.91 [475.75] | 512.35 [512.16] |
| $q_{2}^{J} / \mathrm{kHz}$ | -20.66 [-20.66] | -17.73 [-17.72] | -17.80 [-17.79] | -31.92 [-31.90] |
| $q_{2}^{J J} / \mathrm{Hz}$ | 1.485 [1.484] | 1.205 [1.205] | 1.189 [1.188] | 3.024 [3.024] |
| $q_{2}^{J J J} / \mathrm{mHz}$ | -0.151 [-0.150] | -0.118 [-0.118] | -0.112 [-0.112] | -0.375 [-0.378] |
| $q_{2}^{J J J J} / \mu \mathrm{Hz}$ | 0.017 [0.016] | 0.012 [0.013] | 0.012 [0.011] | 0.053 [0.056] |
| $\sigma_{q} / \mathrm{Hz}$ | 1.28 [2.40] | 2.67 [1.59] | 2.32 [2.29] | 1.74 [1.74] |
| $\alpha_{2} / \mathrm{MHz}$ | -194.31 [-194.30] | -160.79 [-160.80] | -179.81 [-179.81] | -324.73 [-324.61] |
| $B_{2}^{\text {est }} / \mathrm{MHz}$ | 44938.37 [44938.23] | 43465.99 [43466.54] | 43055.73 [43056.17] | 38517.92 [38517.78] |
| $B_{2}^{\text {exp }} / \mathrm{MHz}$ | $44939.7878(38)^{\text {a }}$ |  |  |  |
| $T_{3} / \mathrm{cm}^{-1}$ | 1901.59 [1901.32] | 1862.26 [1862.01] | 1858.88 [1858.63] | 1840.02 [1839.77] |
| $B_{3} / \mathrm{MHz}$ | 44166.72 | 42747.90 | 42327.67 | 37763.65 |
| $B_{3}-B_{3}^{a} / \mathrm{MHz}$ | 14.13 | 13.18 | 13.05 | 10.31 |
| $D_{3} / \mathrm{kHz}$ | 115.47 [115.39] | 107.37 [107.30] | 105.47 [105.40] | 94.06 [94.01] |
| $\mathrm{H}_{3} / \mathrm{Hz}$ | 1.064 [1.066] | 0.724 [0.725] | 0.722 [0.722] | 2.354 [2.352] |
| $L_{3} / \mathrm{mHz}$ | -0.245 [-0.251] | -0.090 [-0.091] | -0.067 [-0.067] | -0.251 [-0.250] |
| $K_{3} / \mu \mathrm{Hz}$ | 0.112 [0.119] | 0.026 [0.026] | 0.012 [0.012] | 0.026 [0.023] |
| $\sigma_{3} / \mathrm{Hz}$ | 9.42 [8.36] | 3.82 [6.08] | 5.11 [5.39] | 1.66 [5.76] |
| $\alpha_{3} / \mathrm{MHz}$ | 358.34 [358.16] | 343.68 [343.50] | 337.60 [337.44] | 260.70 [260.59] |
| $B_{3}^{\text {est }} / \mathrm{MHz}$ | 44385.72 [44385.77] | 42961.52 [42962.23] | 42538.32 [42538.93] | 37932.49 [37932.58] |
| $T_{1} / \mathrm{cm}^{-1}$ | 3277.34 [3276.54] | 3263.63 [3262.83] | 3276.86 [3276.06] | 2483.25 [2482.93] |
| $B_{1} / \mathrm{MHz}$ | 44240.69 | 42835.12 | 42398.30 | 37728.43 |
| $B_{1}-B_{1}^{a} / \mathrm{MHz}$ | 14.21 | 13.27 | 13.13 | 10.30 |
| $D_{1} / \mathrm{kHz}$ | 115.08 [115.01] | 106.88 [106.81] | 105.21 [105.14] | 99.93 [99.88] |
| $H_{1} / \mathrm{Hz}$ | 1.046 [1.045] | 0.820 [0.820] | 0.840 [0.839] | 3.434 [3.434] |
| $L_{1} / \mathrm{mHz}$ | -0.068 [-0.067] | -0.058 [-0.058] | -0.053 [-0.053] | -0.421 [-0.425] |
| $K_{1} / \mu \mathrm{Hz}$ |  |  |  | 0.056 [0.058] |
| $\sigma_{1} / \mathrm{Hz}$ | 15.13 [14.12] | 13.43 [11.59] | 6.71 [6.71] | 2.13 [3.84] |
| $\alpha_{1} / \mathrm{MHz}$ | 284.37 [284.27] | 256.45 [256.37] | 266.97 [266.89] | 295.92 [295.80] |
| $B_{1}^{\text {est }} / \mathrm{MHz}$ | 44459.69 [44459.65] | 43048.75 [43049.36] | 42608.95 [42609.48] | 37897.27 [37897.37] |
| $B_{1}^{\text {exp }} / \mathrm{MHz}$ | $44457.10(24)^{\text {c }}$ |  |  |  |
| ${ }^{(n)} B_{e}^{\text {est }}\left[{ }^{(a)} B_{e}^{\text {est }}\right] / \mathrm{MHz}$ | 44786.69 [44786.50] | 43370.40 [43370.86] | 42920.21 [42920.59] | 38046.09 [38046.18] |
| $S_{0}\left[S_{0}^{a}\right] / \mathrm{MHz}$ | 127.04 [126.91] | 139.27 [139.13] | 122.18 [122.01] | -46.42 [-46.41] |
| $\Delta B_{0} / \mathrm{MHz}$ | 42.69 [42.57] | 65.25 [65.13] | 44.34 [44.23] | -147.04 [-146.99] |
| $\Delta B_{0}-S_{0} / \mathrm{MHz}$ | -84.35 | -74.02 | -77.84 | -100.62 |

[^2]lates a $B_{0}^{\exp }$ of 41409.994 MHz . In Table VII on page 373 of his Ph.D. thesis, ${ }^{14}$ Warner, however, reports a ground state rotational constant of 41411.773 MHz for $\mathrm{H}^{18} \mathrm{O}^{13} \mathrm{C}^{+}$. This result can be obtained from the observed rotational $J=2 \rightarrow 3$ transition only under the assumption of $D_{0}^{\exp }=98.83 \mathrm{kHz}$. Interestingly, the latter $B_{0}^{\exp }$ and $D_{0}^{\exp }$ values agree nicely with our results of Table IV reported in Paper I, where $B_{0}^{\text {est }}=41411.39 \mathrm{MHz}$ and $D_{0}=96.64 \mathrm{kHz}$ are given for $\mathrm{H}^{18} \mathrm{O}^{13} \mathrm{C}^{+}$.

## A. Improving the equilibrium structure

The ground state vibrational correction $\Delta B_{0}$ to the equilibrium rotational constant is derived from our calculations as

$$
\begin{equation*}
\Delta B_{0}=B_{e}^{t h}-B_{0}^{t h} \tag{2}
\end{equation*}
$$

where $B_{e}^{t h}$ and $B_{0}^{t h}$ are theoretical values of the rotational constant at equilibrium and in the ground vibrational state, respectively. The equilibrium geometry used in Eq. (2) to evaluate $B_{e}^{t h}$ is the geometry corresponding to the minimum of the $\operatorname{CCSD}(\mathrm{T}) / \mathrm{cc}-\mathrm{pVQZ}$ potential energy surface. Combining $\Delta B_{0}$ of Eq. (2) with a groundstate rotational constant $B_{0}^{\text {exp }}$ known from experiment, we may derive a new estimate of the equilibrium rotational constant $B_{e}^{\text {est }}$ for a given isotopic species as

$$
\begin{equation*}
B_{e}^{\text {est }}=B_{0}^{\exp }+\Delta B_{0} . \tag{3}
\end{equation*}
$$

Knowing the experimental $B_{0}^{\exp }$ constant for several isotopologues, the values for $B_{e}^{\text {est }}$ of Eq. (3) can be used to determine the equilibrium $r_{e}$ molecular structure. The estimate $B_{v}^{\text {est }}$ for the effective rotational constant in the vibrational state $v$ is then

$$
\begin{equation*}
B_{v}^{\mathrm{est}}=B_{e}^{\mathrm{est}}-\Delta B_{0}-\alpha_{v}, \tag{4}
\end{equation*}
$$

where the vibration-rotation interaction constant $\alpha_{v}$ is

$$
\begin{equation*}
\alpha_{v}=B_{0}^{\mathrm{th}}-B_{v}^{\mathrm{th}} \quad \text { for } \quad v=1-3 \tag{5}
\end{equation*}
$$

For a linear triatomic molecule HXY, the equilibrium rotational constant $B_{e}$ is expressed in terms of Jacobi coordinates $r, R$ as

$$
\begin{equation*}
B_{e}=\frac{\hbar^{2}}{2 I_{e}}=\frac{\hbar^{2}}{2\left(\mu_{r} r_{e}^{2}+\mu_{R} R_{e}^{2}\right)}, \tag{6}
\end{equation*}
$$

where the equilibrium distances $r_{e}$ and $R_{e}$ stand for

$$
\begin{align*}
r_{e} & =r_{e}(\mathrm{XY}), \\
R_{e} & =r_{e}(\mathrm{HX})+\frac{m_{\mathrm{Y}}}{m_{\mathrm{XY}}} r_{e}(\mathrm{XY}) \tag{7}
\end{align*}
$$

using $m_{\mathrm{XY}}=m_{\mathrm{X}}+m_{\mathrm{Y}}$. The reduced masses $\mu_{r}$ and $\mu_{R}$ are defined by

$$
\begin{equation*}
\mu_{r}=\frac{m_{\mathrm{X}} m_{\mathrm{Y}}}{m_{\mathrm{XY}}} \quad \text { and } \quad \mu_{R}=\frac{m_{\mathrm{H}} m_{\mathrm{XY}}}{m_{\mathrm{HXY}}}, \tag{8}
\end{equation*}
$$

TABLE V: Equilibrium distances (in $\AA$ ) of $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$ derived using nuclear and atomic masses. Fit $U$ refers to results from an unweighted fit and DBPC refers to results based on Ref. 15. See Section II A for details. ${ }^{a}$

| $\mathrm{HCO}^{+}$ | $r_{e}(\mathrm{HC})$ | $r_{e}(\mathrm{CO})$ | $r_{e}(\mathrm{HO})$ |
| :--- | :--- | :--- | :--- |
| atomic masses | $1.091981(7)$ | $1.105615(2)$ | $2.19760(1)$ |
| nuclear masses | $1.09180(3)$ | $1.105874(9)$ | $2.19767(4)$ |
| Fit U atomic | $1.09200(1)$ | $1.105609(3)$ | $2.19761(2)$ |
| Fit U nuclear | $1.09187(5)$ | $1.10585(1)$ | $2.19772(6)$ |
| DBPC atomic | 1.09204 | 1.10558 | 2.19762 |
| DBPC nuclear | $1.09193(3)$ | $1.105816(8)$ | $2.19775(4)$ |
| HOC ${ }^{+}$ | $r_{e}(\mathrm{HO})$ | $r_{e}(\mathrm{CO})$ | $r_{e}(\mathrm{HC})$ |
| atomic masses | $0.990482(7)$ | $1.154468(2)$ | $2.14495(1)$ |
| nuclear masses | $0.99041(3)$ | $1.154692(7)$ | $2.14510(4)$ |
| Fit U atomic | $0.99050(2)$ | $1.154464(4)$ | $2.14496(2)$ |
| Fit U nuclear | $0.99046(7)$ | $1.15468(1)$ | $2.14513(7)$ |

${ }^{a}$ Values in parentheses show one standard error to the last significant digits of the distances from the least-squares procedure.
where $m_{\text {HXY }}=M$ is the total molecular mass. The explicit values of $\mu_{R}, \mu_{r}$ obtained using the atomic $m_{a}$ and nuclear $m_{n}$ masses are shown in Table I for the parent $\mathrm{HCO}^{+}$species. The atomic-mass values are larger by approximately $0.03-0.05 \%$, implying that the harmonic wavenumbers for the nuclear-mass case are larger by approximately 0.01-0.03 \% (up to $1 \mathrm{~cm}^{-1}$ ).

The $B_{e}$ value of a single isotopologue is clearly insufficient to uniquely determine $r_{e}$. The equilibrium structures of $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$are therefore extracted from a set of $B_{e}$ values for several isotopologues by means of a Levenberg-Marquardt nonlinear least-squares algorithm ${ }^{23}$ and using experimental uncertainties to compute weights. The analytical expression of Eq. (6) for the equilibrium rotational constant $B_{e}$ was used in combination with analytical expressions for the partial derivatives of $B_{e}$ with respect to the geometric parameters $r_{e}(\mathrm{HX})$ and $r_{e}(\mathrm{XY})$. The equilibrium bond lengths $r_{e}(\mathrm{HX})$ and $r_{e}(\mathrm{XY})$ calculated by this procedure for $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$are summarized in Table V, where we also show $r_{e}(\mathrm{HY})=r_{e}(\mathrm{HX})+r_{e}(\mathrm{XY})$. The results of the calculations using atomic masses are taken from Paper I. Rows denoted by DBPC show the experimental $r_{e}$ values for $\mathrm{HCO}^{+}$from the work of Dore and coworkers. ${ }^{15}$ We were able to reproduce their $r_{e}$ results, shown in Table XI of Ref. 15, by using $B_{e}$ values from their Table X and atomic masses (row DBPC atomic). ${ }^{24}$ Combining the $B_{e}$ values of Dore et al. with nuclear masses, we calculated the nuclear-mass counterparts (row DBPC nuclear).

Upon substitution of the atomic masses of the constituent species with their nuclear counterparts, the length $r_{e}(\mathrm{HX})$ of the bond involving hydrogen decreases by $1.8 \times 10^{-4} \AA$ for $\mathrm{HCO}^{+}$and by $7 \times 10^{-5} \AA$ for $\mathrm{HOC}^{+}$,
whereas the equilibrium bond length $r_{e}(\mathrm{CO})$ increases by $1.6 \times 10^{-4} \AA$ for $\mathrm{HCO}^{+}$and by $2.3 \times 10^{-4} \AA$ for $\mathrm{HOC}^{+}$in Table V. The overall length $r_{e}(\mathrm{HY})$ of the cations is larger for the nuclear masses by approximately $1 \times 10^{-4} \AA$. In other words, the effect of the substitution of the atomic masses with the nuclear masses is in order of $10^{-4} \AA$ for the equilibrium bond lengths and is thus larger than accepted statistical uncertainties for $r_{e}$ shown in Table V.

The equilibrium bond distances of Table V are used to calculate the equilibrium rotational constants, denoted by ${ }^{(n)} B_{e}^{\text {est }}$ and ${ }^{(a)} B_{e}^{\text {est }}$ in the nuclear-mass and atomicmass case, respectively. The new estimates of the effective rotational constants $B_{v}^{\text {est }}$ are derived for $v=0-3$ by means of Eqs. (3) and (4). The values of $B_{e}^{\text {est }}$ and $B_{v}^{\text {est }}$ are listed in Tables II-IV. They both agree with their atomic-mass counterparts within 1.5 MHz for $\mathrm{HCO}^{+}$and within 0.5 MHz for $\mathrm{HOC}^{+}$.

Our estimates $r_{e}(\mathrm{HX}), r_{e}(\mathrm{CO})$ of Table V reproduce the fitted $B_{e}^{\text {est }}$ values with root-mean-square (rms) deviations of $0.8[0.2] \mathrm{MHz}$ for $\mathrm{HCO}^{+}$isotopologues and of $0.5[0.2] \mathrm{MHz}$ for $\mathrm{HOC}^{+}$isotopologues in the calculations using nuclear [atomic] masses, with a largest individual difference of $1.7[0.4] \mathrm{MHz}$ for $\mathrm{H}^{13} \mathrm{C}^{18} \mathrm{O}^{+}$. In our fitting procedure, experimental data coming from different sources were employed, with experimental standard deviations for $B_{0}^{\text {exp }}$ varying from $0.00016 \mathrm{MHz}\left(\mathrm{HCO}^{+}\right)$ to $0.040 \mathrm{MHz}\left(\mathrm{HC}^{18} \mathrm{O}^{+}\right)$in the case of the $\mathrm{HCO}^{+}$cation. To test the sensititity of our results to these values, we also carried out unweighted nonlinear least-squares fits, denoted by Fit U in Table V. The equilibrium distances found in the original and Fit U differ by approximately $1 \times 10^{-5} \AA$. This difference is thus comparable with statistical uncertainties of $r_{e}$ within one to two standard deviations. In Fit U, the $B_{e}^{\text {est }}$ values of $\mathrm{HCO}^{+}$isotopologues are reproduced with a rms deviation of 0.5 [0.1] MHz and a maximum individual deviation of 0.9 [0.3] MHz for $\mathrm{H}^{13} \mathrm{C}^{18} \mathrm{O}^{+}$.

## III. ROTATIONAL CONSTANTS AND THE DETERMINATION OF MOLECULAR STRUCTURE

The equilibrium geometry is the nuclear arrangement corresponding to the minimum of the potential energy surface for a given electronic state, computed in the framework of the Born-Oppenheimer approximation. The experimental counterpart is the $r_{e}$ structure. This is a single geometric structure, which reproduces with high accuracy the experimental equilibrium constants derived by correcting the zero-point rotational constants for vibrational effects. ${ }^{5}$ In the case of linear triatomic molecules, whose geometric arrangements are described by two bond distances, two isotopologues are required for the determination of the geometric parameters from the experimental (rotational) data. In real vibrating-rotating molecules (real experimental sit-
uation), the ground vibrational state is described by a wave function of some (sometimes also considerable) extent over close-to-equilibrium arrangements. Such a situation leads to an effective vibrationally averaged rotational constant $B_{0}$ different from the equilibrium $B_{e}$ value. The zero-point rotational constants $B_{0}$ are experimentally available, but not $B_{e}$.

Two traditional approaches of experimental spectroscopy for the determination of the molecular structure are the $r_{0}$ approach (the $r_{0}$ structure), which directly uses the $B_{0}$ values, and the $r_{s}$ approach (the substitution structure), which employs the analytical solutions of Kraitchman's equations. ${ }^{4,25}$

The so-called $r_{0}$ structure is computed from the zeropoint rotational constants $B_{0}$. Whereas the equilibrium structure is well-defined and unique within the Born-Oppenheimer approximation, this is not the case with the concept of the $r_{0}$ geometry. To exemplify this issue, we provide the radial amplitude $\Delta_{r}$ for the ground vibrational state and several $J$ values in Table VI. There the radial amplitude $\Delta_{r}$ is the difference between the expectation value of the coordinate $r=r(\mathrm{XY})$ in the rovibrational state $|n\rangle$ and the equilibrium distance $r_{e}=r_{e}(\mathrm{XY})$,

$$
\begin{equation*}
\Delta_{r}^{(n)}=\Delta_{r}=\langle n| r|n\rangle-r_{e} . \tag{9}
\end{equation*}
$$

A quick glance at Table VI shows that $\Delta_{r}$ is isotopologuedependent and $J$-dependent. The same also applies for the expectation values $\langle r\rangle$. For a given isotopic variant, $\Delta_{r}$ increases by approximately $5 \times 10^{-4} \AA$ for $J=15$ with respect to the $J=0$ result and decreases by approximately $5 \times 10^{-4} \AA$ with respect to the result for the parent molecule. In our rovibrational calculations, the expectation values $\langle r\rangle$ are found to be almost unaffected by the substitution of atomic masses by nuclear masses, such that Table VI applies to both the atomic-mass and nuclear-mass case.
In addition to the fact that the effective $r_{0}$ structural parameters are different for different isotopic forms, we may also remember that

$$
\begin{equation*}
\langle B\rangle=\left\langle\frac{\hbar^{2}}{2 I}\right\rangle \neq \frac{\hbar^{2}}{2\langle I\rangle}=R_{I} \tag{10}
\end{equation*}
$$

from a mathematical point of view. This is relevant for theoretical approaches. Whereas the integral $\langle I\rangle$ can be solved analytically, the integral $\langle B\rangle$ is solvable only by numerical means. To substantiate this issue, we compare in Table VII the ground state rotational constant $B_{0}$ with the vibrationally averaged value $\langle B\rangle$ and with $R_{I}$ for $\mathrm{HCO}^{+}$and $\mathrm{DCO}^{+}$. The values of $R_{I}$ are computed using analytical integrals and the basis set expansion of the full-dimensional wavefunction for $\mathrm{J}=0$. In Table VII, the quantity $B_{0 \rightarrow 1}$ is additionally shown, which is computed from the ground-state energies $E_{0}(J, p)$ as

$$
\begin{equation*}
B_{0 \rightarrow 1}=\frac{1}{2}\left[E_{0}(J=1, p=1)-E_{0}(J=0, p=0)\right] \tag{11}
\end{equation*}
$$

where $p$ stands for parity. The results for $B_{0}$ are taken from Tables II and III, respectively.

TABLE VI: Amplitude $\Delta_{r}$ (in $\AA$ ) in the ground vibrational state of the isotopic variants of $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$with the total rotational angular momentum $J=0,5,10$, and 15 .

|  | $\mathrm{HCO}^{+}$ |  |  |  | $\mathrm{HOC}^{+}$ |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| species | $J=0$ | $J=5$ | $J=10$ | $J=15$ | species | $J=0$ | $J=5$ | $J=10$ | $J=15$ |
| $\mathrm{HCO}^{+}$ | 0.00469 | 0.00475 | 0.00490 | 0.00517 | $\mathrm{HOC}^{+}$ | 0.00853 | 0.00864 | 0.00885 | 0.00922 |
| $\mathrm{HC}^{17} \mathrm{O}^{+}$ | 0.00464 | 0.00469 | 0.00485 | 0.00512 | $\mathrm{H}^{17} \mathrm{OC}^{+}$ | 0.00848 | 0.00858 | 0.00880 | 0.00911 |
| $\mathrm{HC}^{18} \mathrm{O}^{+}$ | 0.00459 | 0.00464 | 0.00480 | 0.00506 | $\mathrm{H}^{18} \mathrm{OC}^{+}$ | 0.00842 | 0.00848 | 0.00869 | 0.00906 |
| $\mathrm{H}^{13} \mathrm{CO}^{+}$ | 0.00453 | 0.00459 | 0.00475 | 0.00501 | $\mathrm{HO}^{13} \mathrm{C}^{+}$ | 0.00842 | 0.00853 | 0.0087 | 0.00906 |
| $\mathrm{H}^{13} \mathrm{C}^{17} \mathrm{O}^{+}$ | 0.00459 | 0.00464 | 0.00480 | 0.00506 | $\mathrm{H}^{17} \mathrm{O}^{13} \mathrm{C}^{+}$ | 0.00837 | 0.00842 | 0.0086 | 0.00901 |
| $\mathrm{H}^{13} \mathrm{C}^{18} \mathrm{O}^{+}$ | 0.00448 | 0.00453 | 0.00469 | 0.00496 | $\mathrm{H}^{18} \mathrm{O}^{13} \mathrm{C}^{+}$ | 0.00832 | 0.00837 | 0.00858 | 0.00895 |
| $\mathrm{DCO}^{+}$ | 0.00453 | 0.00459 | 0.00469 | 0.00485 | $\mathrm{DOC}^{+}$ | 0.00795 | 0.00800 | 0.00816 | 0.00842 |
| $\mathrm{DC}^{17} \mathrm{O}^{+}$ | 0.00448 | 0.00453 | 0.00464 | 0.00480 | $\mathrm{D}^{17} \mathrm{OC}^{+}$ | 0.00784 | 0.00790 | 0.00811 | 0.00837 |
| $\mathrm{DC}^{18} \mathrm{O}^{+}$ | 0.00443 | 0.00448 | 0.00459 | 0.00475 | $\mathrm{D}^{18} \mathrm{OC}^{+}$ | 0.00779 | 0.0078 | 0.00800 | 0.00827 |
| $\mathrm{D}^{13} \mathrm{CO}^{+}$ | 0.00443 | 0.00448 | 0.00459 | 0.00480 | $\mathrm{DO}^{13} \mathrm{C}^{+}$ | 0.00784 | 0.00790 | 0.00805 | 0.00832 |
| $\mathrm{D}^{13} \mathrm{C}^{17} \mathrm{O}^{+}$ | 0.00438 | 0.00443 | 0.00453 | 0.00469 | $\mathrm{D}^{17} \mathrm{O}^{13} \mathrm{C}^{+}$ | 0.00774 | 0.00779 | 0.00795 | 0.00821 |
| $\mathrm{D}^{13} \mathrm{C}^{18} \mathrm{O}^{+}$ | 0.00432 | 0.00438 | 0.00448 | 0.00464 | $\mathrm{D}^{18} \mathrm{O}^{13} \mathrm{C}^{+}$ | 0.00768 | 0.00774 | 0.00790 | 0.00816 |

TABLE VII: Various representations of the ground state rotational constants computed for $\mathrm{HCO}^{+}$and $\mathrm{DCO}^{+}$using nuclear masses.

|  | $B_{0}$ | $B_{0 \rightarrow 1}$ | $\langle B\rangle$ | $R_{I}$ |
| :--- | :---: | :---: | :---: | :---: |
| $\mathrm{HCO}^{+}$ |  |  |  |  |
| $(\mathrm{MHz})$ | 44391.37 | 44391.21 | 44245.40 | 44155.26 |
| $\left(\mathrm{~cm}^{-1}\right)$ | 1.48074 | 1.48073 | 1.47587 | 1.47286 |
| $\mathrm{DCO}^{+}$ |  |  |  |  |
| $(\mathrm{MHz})$ | 35861.59 | 35861.48 | 35743.61 | 35675.60 |
| $\left(\mathrm{~cm}^{-1}\right)$ | 1.19621 | 1.19621 | 1.19228 | 1.19001 |

For the vibrational ground state of $\mathrm{HCO}^{+}$, the vibrationally averaged moments of inertia is $14.166 \mathrm{u}^{2}$. Compared to $B_{0}$, the values for $\langle B\rangle$ and $R_{I}$ are smaller by approximately 100 and 200 MHz , respectively, what amounts to $0.5 \%$. On the other hand, the value for $B_{0 \rightarrow 1}$, computed according to Eq. (11) from the ground state energies for $J=0,1$ only, differ by 0.16 and 0.11 MHz from the $B_{0}$ result for $\mathrm{HCO}^{+}$and $\mathrm{DCO}^{+}$, respectively. This finding leads us to the conclusion that Coriolis coupling does not play an important role in the internal dynamics of both $\mathrm{HCO}^{+}$and $\mathrm{DCO}^{+}$. The comparison of the values presented in Table VII also indicates that rovibrational calculations, carried out even for only a modest number of $J$ values, may provide more useful results for $B$ than their vibrationally averaged counterparts obtained from solely $J=0$ computations.

The determination of structural parameters from the experimental data is very often carried out by the em-
ployment of the moments of inertia, which are reciprocals of the rotational constants, Eq. (6). The positions $z_{1}, z_{2}, z_{3}$ of the aligned atoms $\mathrm{H}, \mathrm{X}, \mathrm{Y}$ in the reference system with the origin in the center of mass of the parent molecule HXY are explicitly

$$
\begin{align*}
M z_{1} & =-m_{\mathrm{XY}} x-m_{\mathrm{Y}} y, \\
M z_{2} & =m_{\mathrm{H}} x-m_{\mathrm{Y}} y,  \tag{12}\\
M z_{3} & =m_{\mathrm{H}} x+m_{\mathrm{HX}} y,
\end{align*}
$$

where $x=r(\mathrm{HX})$ and $y=r(\mathrm{XY})$. The difference $\Delta I$ between the moments of inertia of two different isotopic forms is then

$$
\begin{align*}
\Delta I=I_{2}-I_{1}= & \left(a_{2}-a_{1}\right) x^{2} \\
& +\left(b_{2}-b_{1}\right) x y+\left(c_{2}-c_{1}\right) y^{2} . \tag{13}
\end{align*}
$$

For explicit values of the parameters $a_{i}, b_{i}, c_{i}$, see Eq. (A4) of the Appendix. It is an easy matter to derive for a simple substitution $m_{i} \rightarrow m_{i}^{\prime}$ and $i=1-3$ that

$$
\begin{align*}
& \Delta I\left(m_{\mathrm{H}} \rightarrow m_{\mathrm{H}}^{\prime}\right)=\frac{m_{\mathrm{H}}^{\prime}-m_{\mathrm{H}}}{M M^{\prime}}\left(m_{\mathrm{XY}} x+m_{\mathrm{Y}} y\right)^{2} \\
& \Delta I\left(m_{\mathrm{X}} \rightarrow m_{\mathrm{X}}^{\prime}\right)=\frac{m_{\mathrm{X}}^{\prime}-m_{\mathrm{X}}}{M M^{\prime}}\left(m_{\mathrm{H}} x-m_{\mathrm{Y}} y\right)^{2}  \tag{14}\\
& \Delta I\left(m_{\mathrm{Y}} \rightarrow m_{\mathrm{Y}}^{\prime}\right)=\frac{m_{\mathrm{Y}}^{\prime}-m_{\mathrm{Y}}}{M M^{\prime}}\left(m_{\mathrm{H}} x+m_{\mathrm{HY}} y\right)^{2}
\end{align*}
$$

implying

$$
\begin{equation*}
z_{i}^{2}=\frac{M^{\prime}}{M\left(m_{\mathrm{i}}^{\prime}-m_{\mathrm{i}}\right)} \Delta I\left(m_{\mathrm{i}} \rightarrow m_{\mathrm{i}}^{\prime}\right) . \tag{15}
\end{equation*}
$$

The latter equation is known as Kraitchman's relation for linear triatomic molecules. ${ }^{4}$ It provides the position
$z_{i}$ of the $i$ th atom with respect to the center of mass of the parent molecule as a function of the moment-of-inertia difference $\Delta I\left(m_{\mathrm{i}} \rightarrow m_{\mathrm{i}}^{\prime}\right)$. From three known $z_{1}, z_{2}, z_{3}$ values, the bond distances are calculated as $r(\mathrm{HX})=x=z_{2}-z_{1}$ and $r(\mathrm{XY})=y=z_{3}-z_{2}$. To compute two distances of a linear triatomic molecule by means of Eq. (15), we effectively need four $B$ values available for the parent molecules HXY and three singly substituted species DXY, HX'Y, and HXY'.

Kraitchman's equation of Eq. (15) is frequently used for the estimation of the experimental equilibrium structure. Letting $I_{1}$ and $I_{2}$ be the moment of inertia of the parent molecule and of the substituted species, respectively, where

$$
\begin{equation*}
I_{i}=I_{i}^{(0)}=\frac{\hbar^{2}}{2 B_{0}^{(i)}} \quad \text { for } \quad i=1,2 \tag{16}
\end{equation*}
$$

then

$$
\begin{align*}
\Delta I^{(0)} & =I_{1}^{(0)}-I_{2}^{(0)}=I_{1}^{(e)}-I_{2}^{(e)}+D_{I}  \tag{17}\\
& =\Delta I^{(e)}+D_{I}=\Delta I .
\end{align*}
$$

In the case that the vibrational corrections for two isotopic species in the ground vibrational state are similar, the deviation $D_{I}$ in Eq. (17) will be small, so that $\Delta I^{(0)} \approx \Delta I^{(e)}$, implying that $\Delta I^{(0)}$ may be used instead of $\Delta I^{(e)}$. The structure obtained in this fashion is known as the substituted $r_{s}$ structure. Only under the condition of $\Delta I^{(0)} \approx \Delta I^{(e)}$, the $r_{s}$ structure will be a good approximation of the $r_{e}$ structure. The $r_{m}$ structure introduced by Watson, derived by approximating the equilibrium moment of inertia with $2 I_{s}-I_{0}$, is found less satisfactory for hydrogen containing species, ${ }^{5}$ where $I_{s}$ is the moment of inertia computed using the $r_{s}$ geometry.

Since two bond lengths of a linear triatomic molecule can be always determined from two moments of inertia available for two different isotopologues, we finally also examine a pair determination of the molecular structure. An analytic solution for the corresponding mathematical problem is provided in the Appendix. In the case when experimental data are known for $n$ isotopic species, we can pair them in $N=n(n-1) / 2$ different ways, yielding $N$ pair solutions $\left(x_{i}, y_{i}\right)$, where $x=r(\mathrm{HX})$ and $y=r(\mathrm{XY})$ for $i=1, \ldots, N$. The mean pair values are

$$
\begin{equation*}
x^{\mathrm{ap}}=\frac{1}{N} \sum_{i=1}^{N} x_{i} \quad \text { and } \quad y^{\mathrm{ap}}=\frac{1}{N} \sum_{i=1}^{N} y_{i} . \tag{18}
\end{equation*}
$$

To quantify the spread of the pair solutions $\left(x_{i}, y_{i}\right)$ around the mean value ( $x^{\text {ap }}, y^{\text {ap }}$ ), we introduce the average radius of the pair distribution $d_{\text {ap }}$ as

$$
\begin{equation*}
d_{\mathrm{ap}}=\sqrt{\sum_{i=1}^{N} \frac{1}{N}\left[\left(x_{i}-x_{1}^{\mathrm{ap}}\right)^{2}+\left(y_{i}-y_{2}^{\mathrm{ap}}\right)^{2}\right]} . \tag{19}
\end{equation*}
$$

We will call the solution $\left(x^{\text {ap }}, y^{\text {ap }}\right)$ a $r_{\text {ap }}$ structure.

To correct for the rovibrational effects in $B_{0}$, one needs the effective rotational constants for the singly excited vibrational states and two isotopic forms in experimental spectroscopy. Knowing the effective rotational constants $B_{1}, B_{2}, B_{3}$, one may derive the vibration-rotation interaction constants $\alpha_{1}, \alpha_{2}, \alpha_{3}$, yielding the spectroscopic zeropoint vibrational correction as

$$
\begin{equation*}
S_{0}=\frac{1}{2}\left(\alpha_{1}+2 \alpha_{2}+\alpha_{3}\right) \tag{20}
\end{equation*}
$$

for linear triatomic molecules in the traditional spectroscopic approach. Therefrom the experimental equilibrium rotational constant follows as

$$
\begin{equation*}
{ }^{(\alpha)} B_{e}^{\exp }=B_{0}^{\exp }+S_{0} \tag{21}
\end{equation*}
$$

We will denote by $r_{\alpha}$ the equilibrium structure derived from ${ }^{(\alpha)} B_{e}^{\exp }$. In electronic-structure program packages, the expression of Eq. (21) has become a standard tool to compute the zero-point rotational constant $B_{0}$ from the calculated equilibrium geometry (providing $B_{e}$ ) and the zero-point vibrational correction $S_{0}$ computed according to Eq. (20) by means of second order vibrational perturbation theory. ${ }^{26}$ This spectroscopic model is effectively based on the harmonic-oscillator-rigid-rotor description as a zero-order picture.

In our approach used here and in Paper I, the ground state vibrational corrections $\Delta B_{0}$ are computed according to Eq. (2) as a difference between the rotational constant $B_{e}^{t h}$ at equilibrium of the $\operatorname{CCSD}(\mathrm{T}) / \mathrm{cc}-\mathrm{pVQZ}$ potential energy surface and the rotational constant $B_{0}^{t h}$ in the ground vibrational state. The $B_{0}^{t h}$ values are derived from rovibrational energies computed for $J=0-15$ by a numerically exact full-dimensional quantum-mechanical method. In addition to $\Delta B_{0}$, we also derived the vibration-rotation interaction constants $\alpha_{v}=B_{0}^{t h}-B_{v}^{t h}$ of Eq. (5) and used them to calculate the spectroscopic correction $S_{0}$ according to Eq. (20). The quantities $\Delta B_{0}$ and $S_{0}$ are both shown in Tables II-IV. The difference between $\Delta B_{0}$ and $S_{0}$ is in order of 1 MHz for the hydrogen containing forms of $\mathrm{HCO}^{+}$in Table II. However, $\Delta B_{0}-S_{0}$ is as large as 50 and 100 MHz for $\mathrm{DCO}^{+}$and $\mathrm{DOC}^{+}$in Tables III and IV, respectively. In our calculations, the mass effect on $\Delta B_{0}$ and $S_{0}$ is smaller than 0.2 MHz .

## A. Results

In the case when experimental data are available for a series of isotopic variants, the determination of the geometric parameters may proceed employing different mathematical procedures. Nonlinear fitting algorithms are expected to give the best over-all fit to all of the rotational constants. Analytical solutions using Kraitchman's relations are applicable for the parent molecule and three singly substituted species. The geometry problem can be also solved analytically for any pair of isotopic variants. The rotational constants not included in the

TABLE VIII: Structural parameters (in $\AA$ ) obtained for $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$by means of Kraitchman's relations. Root-meansquare (rms) deviations are in MHz . Values shown in brackets for $\mathrm{HCO}^{+}$are rms deviations for the rotational constants of all eight isotopologues.

| source | structure | $\mathrm{HCO}^{+}$ |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | atomic mass |  |  | nuclear mass |  |  |
|  |  | $r(\mathrm{HC})$ | $r(\mathrm{CO})$ | rms | $r(\mathrm{HC})$ | $r(\mathrm{CO})$ | rms |
| $B_{0}^{\text {exp }}$ | $r_{s}^{(0)}$ | 1.092921 | 1.107228 | 108.1 [101.3] | 1.092926 | 1.107236 | 121.2 [113.5] |
| $B_{0}^{\text {exp }}+S_{0}$ | $r_{s}^{(\alpha)}$ | 1.098720 | 1.104852 | 32.5 [46.5] | 1.098732 | 1.104862 | 19.8 [37.1] |
| $B_{0}^{\exp }+\Delta B_{0}$ | $r_{s}^{(e)}$ | 1.092014 | 1.105560 | 2.9 [2.7] | 1.092020 | 1.105568 | 15.8 [14.8] |
| $\mathrm{HOC}^{+}$ |  |  |  |  |  |  |  |
| source | structure | atomic mass |  |  | nuclear mass |  |  |
|  |  | $r(\mathrm{HO})$ | $r(\mathrm{CO})$ | rms | $r(\mathrm{HO})$ | $r(\mathrm{CO})$ | rms |
| $B_{0}^{\text {exp }}$ | $r_{s}^{(0)}$ | 0.964099 | 1.159471 | 7.5 | 0.964102 | 1.159478 | 18.4 |
| $B_{0}^{\text {exp }}+S_{0}$ | $r_{s}^{(\alpha)}$ | 0.984741 | 1.155069 | 52.8 | 0.984759 | 1.155071 | 40.2 |
| $B_{0}^{\exp }+\Delta B_{0}$ | $r_{s}^{(e)}$ | 0.990522 | 1.154383 | 4.8 | 0.990543 | 1.154387 | 17.3 |

process of determining the geometry by analytical means will generally be less well predicted, such that the resulting structure will be a less balanced representation compared to the structure from the nonlinear fits of all available isotopologues.

A more serious obstacle to the derivation of an equilibrium structure is related to uncertainties which occur as a necessary consequence of the effects due to zero-point vibrations. We investigated this issue in practical terms by considering the ground state vibrational corrections given as $\Delta B_{0}$ of Eq. (2) and as $S_{0}$ of Eq. (20).

The structural parameters of $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$derived in different approaches are summarized in Tables VIII and IX. The results are presented for both the atomic-mass and nuclear-mass cases. The root-meansquare (rms) deviation is employed as a measure of goodness of the derived bond lengths to predict the corresponding set of rotational constants. Eight isotopic variants of $\mathrm{HCO}^{+}$and four of $\mathrm{HOC}^{+}$are considered.

Table VIII summarizes distances derived using Kraitchman's relations. The main isotopic forms of $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$were chosen as parent species. The four isotopic variants considered, for instance, for the formyl cation were $\mathrm{HCO}^{+}, \mathrm{DCO}^{+}, \mathrm{H}^{13} \mathrm{CO}^{+}$, and $\mathrm{HC}^{18} \mathrm{O}^{+}$. Three sets of data are studied using the experimental zero-point rotational constants $B_{0}^{\exp }$ shown in Tables II-IV. The values of $B_{0}^{\exp }$ uncorrected for vibrational effects form the first set, yielding the substitution structure denoted by $r_{s}^{(0)}$ in Table VIII. The second set composed of $B_{0}^{\exp }+S_{0}$ produces the $r_{s}^{(\alpha)}$ structure, whereas the third set based on the values of $B_{0}^{\exp }+\Delta B_{0}$ gives the $r_{s}^{(e)}$ structure. For $\Delta B_{0}$ and $S_{0}$, we use our the-
oretical values. Note that the structure $r_{s}^{(0)}$ is commonly called the $r_{s}$ structure in the spectroscopic literature.

The distances in Table VIII show negligible variations upon the replacement of atomic masses with their nuclear counterparts. This is to be expected from the mathematical form of Kraitchman's relation in Eq. (15), which involves explicit mass dependence only through the factor $f_{m}=M^{\prime} / M\left(m_{\mathrm{i}}^{\prime}-m_{\mathrm{i}}\right)$. The difference between the $f_{m}$ values computed using atomic and nuclear masses is somewhat smaller than $1 \times 10^{-5} \mathrm{u}^{-1}$ for the studied systems. To facilitate comparison between distances obtained using the atomic and nuclear masses, the values in Table VIII are given with six decimal places.

The $r_{s}$ distances in Table VIII exhibit a broad spread of 0.006 and $0.026 \AA$ for $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$, respectively. The large difference of approximately $6-7 \times 10^{-3} \AA$ between the $r_{s}^{(\alpha)}(\mathrm{HX})$ and $r_{s}^{(e)}(\mathrm{HX})$ distances arises from a large difference of 54 and 100 MHz between the $S_{0}$ and $\Delta B_{0}$ values for $\mathrm{DCO}^{+}$and $\mathrm{DOC}^{+}$in Tables III and IV, respectively. Compared to the equilibrium distances of Table V , the $r_{s}^{(e)}(\mathrm{HX})$ bond length is longer by $3-4 \times 10^{-5} \AA$ and $r_{s}^{(e)}(\mathrm{XY})$ shorter by $4-8 \times 10^{-5} \AA$ in the atomic-mass case. These differences are in order of $10^{-4} \AA$ for the nuclear-mass case since the $r_{s}$ distances are nearly insensitive to the replacement of atomic masses with their nuclear counterparts.

For given $r_{s}$ distances, rotational constants for all isotopologues were evaluated according to Eq. (6). The rms deviations in Table VIII are thus a measure of goodness of the approximation of the equilibrium rotational constant by $B_{0}^{\exp }, B_{0}^{\exp }+S_{0}$, and $B_{0}^{\exp }+\Delta B_{0}$ in the case of the $r_{s}^{(0)}, r_{s}^{(\alpha)}$, and $r_{s}^{(e)}$ structure, respectively. Among


FIG. 1: Pair solutions $\left(x_{i}, y_{i}\right)_{a}$ and $\left(x_{i}, y_{i}\right)_{n}$ for $\mathrm{HCO}^{+}$obtained for the atomic and nuclear masses, respectively, from $B_{0}^{\exp }+\Delta B_{0}$. Explicit values of $r_{e}(\mathrm{a}), r_{e}^{\text {ap }}(\mathrm{a})$ for the atomic masses and $r_{e}(\mathrm{n}), r_{e}^{\mathrm{ap}}(\mathrm{n})$ for the nuclear masses are found in Table IX. For the definition of $r_{e}^{\mathrm{ap}}(\mathrm{n})_{\text {red }}$, see the main text.
the three representations in Table VIII, the $r_{s}^{(e)}$ distances perform the best, providing the smallest rms deviation. Root-mean-square deviations are somewhat larger for the nuclear-mass case than for the atomic-mass case.

The three sets of data involving the $B_{0}^{\exp }, B_{0}^{\exp }+S_{0}$, and $B_{0}^{\exp }+\Delta B_{0}$ values are now used in combination with a nonlinear least-squares procedure, yielding the $r_{0}, r_{\alpha}$, and $r_{e}$ distances, respectively. These results are summarized in Table IX. In Fit 4 there, we employ the data for the four $\mathrm{HCO}^{+}$isotopologues used to derive Kraitchman's solutions of Table VIII. The $r_{e}$ distances of Table V are repeated in bold face in Table IX. For the $r_{0}$ and $r_{e}$ structures, we also give the mean pair solutions of Eq. (18), denoted by $r_{0}^{\text {ap }}$ and $r_{e}^{\text {ap }}$, respectively. The mean radius of the pair distribution is defined by Eq. (19). Inspired by the approach of Kraitchman, we also tested a fitting procedure for the differences of rotational constants. These results are labelled with $r_{0}^{\text {diff }}$ and $r_{e}^{\text {diff }}$. In addition to the rms deviations for the rotational constants, Table IX also shows rms errors for the differences of the rotational constants in brackets. For our data sets, the total number of pairs and differences is 28 for $\mathrm{HCO}^{+}$ and 6 for $\mathrm{HOC}^{+}$.
All of the pair solutions for $B_{0}^{\exp }+\Delta B_{0}$ of $\mathrm{HCO}^{+}$are graphically displayed in Fig. 1. The pair solutions derived in the atomic-mass case lie in $x_{i} \in(1.0910 \AA, 1.0929 \AA)$ and $y_{i} \in(1.1054 \AA, 1.1059 \AA)$. The pair solutions obtained using nuclear masses are distributed over a larger space, occupying $x_{i} \in(1.0892 A, 1.0944 \AA$ ) and $y_{i} \in(1.1054 A, 1.1067 \AA)$. Most of the solutions are aligned or nearly aligned for both the atomic-mass and nuclear-mass cases in Fig. 1. Two $\left(x_{i}, y_{i}\right)_{n}$ pairs for the
nuclear masses falling out of line are circled in Fig. 1. Eliminating them from the averaging process, the mean value $\left(x_{e}^{\text {ap }}, y_{e}^{\mathrm{ap}}\right)$ is changed from $(1.09239 \AA, 1.10575 \AA$ ) to a value of $(1.09213 \AA, 1.10558 \AA)$. This solution is denoted by $r_{e}^{\text {ap }}(n)_{\text {red }}$ in Fig. 1.

The estimates $r_{e}$ for the equilibrium bond distances reproduce the rotational constants $B_{0}^{\exp }+\Delta B_{0}$ with a rms deviation of about $0.2[0.8] \mathrm{MHz}$ in the atomic-mass [nuclear-mass] case, as seen in Table IX. This implies that $B_{0}^{\exp }+\Delta B_{0}$ approximates fairly well the equilibrium rotational constant $B_{e}^{\exp }$. Rms deviations of approximately 5 MHz are seen for the $r_{0}$ parameters, which are computed from $B_{0}^{\exp }$. In the case of the $r_{\alpha}$ structure, the results for $B_{0}^{\exp }+S_{0}$ are reproduced by the linearmolecule model of Eq. (6) with rms deviations of about 20 MHz for $\mathrm{HCO}^{+}$and 2 MHz for $\mathrm{HOC}^{+}$.

The three $r_{0}, r_{\alpha}, r_{e}$ geometries in Table IX are different. For $\mathrm{HCO}^{+}$, the distance $r_{0}(\mathrm{HC})$ is shorter by $0.0006 \AA$ and $r_{\alpha}(\mathrm{HC})$ longer by $0.004 \AA$ than $r_{e}(\mathrm{HC})$, whereas $r_{0}(\mathrm{CO})$ is longer by $0.004 \AA$ and $r_{\alpha}(\mathrm{CO})$ shorter by $0.0008 \AA$ than $r_{e}(\mathrm{CO})$. In the case of $\mathrm{HOC}^{+}$, the $r_{\alpha}$ geometry deviates somewhat less than the $r_{0}$ geometry from the $r_{e}$ results. Interestingly, the pair solution $r_{e}^{\text {ap }}$ and the solution $r_{e}^{\text {diff }}$, obtained by fitting the differences of the rotational constants, agree with each other within $1-2 \times 10^{-5} \AA$. They both, however, differ by $1-2 \times 10^{-4} \AA$ from the $r_{e}$ structure.

Another salient feature is observed by comparing the results for $r_{0}, r_{\alpha}, r_{e}$ from Fit 4 of Table IX with the values of $r_{s}^{(0)}, r_{s}^{(\alpha)}, r_{s}^{(e)}$ derived from Kraitchman's relations in Table VIII. In all three cases, rms deviations are smaller for the geometric parameters obtained by nonlinear leastsquares fits, which thus appear more appropriate for the determination of molecular structures.

## IV. FINAL REMARKS

Accurate determination of the equilibrium geometry of molecules is considered as one of the most important goals of spectroscopy. Geometric parameters are unambiguously determined only from actual equilibrium rotational constants $B_{e}$, which, however, are not accessible experimentally. Experimental spectroscopy thus takes on other views of the situation. If there is no data on excited vibrational states, two traditional approaches to determining the molecular structure are the $r_{0}$ approach, which uses the zero-point rotational constants $B_{0}$ directly, and the $r_{s}$ approach, which uses Kraitchman's relations in combination with $B_{0}$. In the case when information is available on singly excited vibrational states, the zero-point rotational constants $B_{0}$ are corrected for the vibrational effects with the help of the spectroscopic correction $S_{0}$ of Eq. (20) to give ${ }^{(\alpha)} B_{e}^{\exp }$ of Eq. (21), yielding the $r_{\alpha}$ structure. The three sets $r_{0}, r_{s}, r_{\alpha}$ of geometric parameters are generally different. Modifications of data sets, such as a change of a parent molecule in the Kraitchman's approach, may introduce additional inconsisten-

TABLE IX: Various structural parameters (in $\AA$ ) obtained for $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$by means of a nonlinear least-squares technique. Root-mean-square (rms) deviations are shown in MHz. For more details, see the main text. ${ }^{a}$

| $\mathrm{HCO}^{+}$ |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| structure | atomic mass |  |  |  | nuclear mass |  |  |  |
|  | $r$ ( HC ) | $r(\mathrm{CO})$ | rms | $d_{\text {ap }}$ | $r$ ( HC ) | $r$ (CO) | rms | $d_{\text {ap }}$ |
| $r_{0}$ | $1.09142(22)$ | $1.10932(6)$ | 5.0 [5.7] |  | $1.09123(24)$ | 1.109 59(7) | 5.6 [6.4] |  |
| $r_{0}$ (Fit 4) | $1.09172(5)$ | 1.109 27(1) | 3.9 [4.5] |  | $1.09156(6)$ | $1.10952(2)$ | 4.3 [5.0] |  |
| $r_{0}^{\text {ap }}$ | 1.09547 | 1.10846 | 15.1 [19] | $7.1 \times 10^{-3}$ | 1.09577 | 1.10862 | 16.9 [21] | $7.8 \times 10^{-3}$ |
| $r_{0}^{\text {diff }}$ | 1.095 30(10) | 1.109 92(2) | 92 [1.8] |  | $1.09554(11)$ | $1.11025(2)$ | 102 [2.1] |  |
| $r_{\alpha}$ | 1.0961 (30) | 1.1048 (9) | 20.4 [31] |  | 1.0959 (30) | 1.1051(9) | 20.5 [31] |  |
| $r_{\alpha}$ (Fit 4) | 1.099 29(2) | $1.104197(7)$ | 1.2 [1.6] |  | $1.09915(2)$ | 1.104449 (5) | 0.9 [1.2] |  |
| $\mathrm{r}_{\mathrm{e}}$ | $1.091981(7)$ | $1.105615(2)$ | 0.2 [0.2] |  | 1.09180 (3) | $1.105874(9)$ | 0.8 [0.9] |  |
| $r_{e}$ (Fit 4) | $1.091989(1)$ | $1.105614(3)$ | 0.1 [0.1] |  | $1.091839(8)$ | $1.105867(2)$ | 0.6 [0.7] |  |
| $r_{e}^{\text {ap }}$ | 1.09209 | 1.10559 | 0.5 [0.5] | $3.5 \times 10^{-4}$ | 1.09239 | 1.10575 | 2.4 [2.7] | $1.1 \times 10^{-3}$ |
| $r_{e}^{\text {diff }}$ | 1.092 104(7) | $1.105634(1)$ | 2.9 [0.1] |  | $1.09234(1)$ | $1.105957(2)$ | 13 [0.4] |  |

$\mathrm{HOC}^{+}$

| structure | atomic mass |  |  |  | nuclear mass |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $r$ (HO) | $r(\mathrm{CO})$ | rms | $d_{\text {ap }}$ | $r$ ( HO ) | $r(\mathrm{CO})$ | rms | $d_{\text {ap }}$ |
| $r_{0}$ | $0.96532(20)$ | 1.15931 (5) | 4.5 [7.1] |  | 0.965 23(21) | $1.15953(6)$ | 4.4 [7.0] |  |
| $r_{0}^{\text {ap }}$ | 0.95980 | 1.16018 | 21 [28] | 0.016 | 0.96024 | 1.16032 | 19 [26] | 0.017 |
| $r_{0}^{\text {diff }}$ | 0.9680 (12) | $1.1602(4)$ | 87 [10] |  | 0.9683 (12) | $1.1605(4)$ | 99 [10] |  |
| $r_{\alpha}$ | 0.98548 (6) | $1.15407(2)$ | 2.3 [3.1] |  | 0.98540 (4) | 1.15430 (1) | 2.0 [2.7] |  |
| $\mathrm{r}_{\mathrm{e}}$ | 0.990 482(7) | 1.154 468(2) | 0.14 [0.18] |  | 0.99041 (3) | 1.154 692(7) | 0.5 [0.6] |  |
| $r_{e}^{\text {ap }}$ | 0.99069 | 1.15443 | 0.7 [1.1] | $2.5 \times 10^{-4}$ | 0.99114 | 1.15456 | 2.5 [3.6] | $9.1 \times 10^{-4}$ |
| $r_{e}^{\text {diff }}$ | 0.99062 | 1.15451 | 4.5 [0.06] |  | 0.99092 | 1.15485 | 16 [0.1] |  |

${ }^{a}$ Values in parentheses show one standard error to the last significant digits of the distances from the least-squares procedure.
cies among the values predicted by a chosen method. ${ }^{14}$
The concepts of the equilibrium geometry and the vibrational correction to the zero-point rotational constant are well founded in theoretical approaches, such that theory may provide useful assistance to experiment in the evaluation of the equilibrium structure. To achieve as accurate as possible corrections to $B_{e}$ in the real situation of a vibrating and rotating molecule, theoretical methods for numerically exact quantum-mechanical calculations are desirable since only then we can have a proper full-dimensional physical answer for a given potential energy surface in the Born-Oppenheimer approximation. In the present work and previously in Paper I, we point to the difference between the spectroscopic correction $S_{0}$ of Eq. (20) and the zero-point correction $\Delta B_{0}$, defined in Eq. (2) literally as a difference between the rotational constant at equilibrium and the rotational constant in the ground vibrational state. In connection with this, we also observed that rovibrational calculations, carried out even for lowest $J$ values, may provide useful values of
$B_{0}$. Unlike variational methods, vibrational second-order perturbational approaches are capable of providing only the spectroscopic correction $S_{0}$, which is not always sufficient to evaluate the molecular structure. The concept of the $r_{e}$ structure is generally useful. Its applicability in the case of quasi-linear molecules and weakly bonded complexes deserves, however, some attention.

Another issue relevant for the determination of molecular geometries is related to the answer on the question 'which masses are vibrating or rotating in a molecule?', to cite the title of a paper by Kutzelnigg. ${ }^{6}$ This question appears natural in connection with charged molecular systems, like $\mathrm{HCO}^{+}$and $\mathrm{HOC}^{+}$, which lack an electron.

Several choices of mass are commonly used in nuclear dynamics computations. The atomic masses have became standard in connection with the BornOppenheimer approximation because the atomic masses are expected to minimize the deviation of the employed electronic-structure approach with respect to complete theory. For the Born-Oppenheimer approximation plus
adiabatic contributions, the nuclear masses are considered more adequate physically. Rescaled masses or distance-dependent masses are also encountered in some applications. ${ }^{6,27,28}$ Systematic improvements of the computational approaches are possible only in conjunction with the nuclear masses.

In experimental studies, the atomic masses are tacitly in use, ${ }^{5}$ so that an explicit reference regarding the employed masses is rarely given. In the case of $\mathrm{HCO}^{+}$, we may note that Bogey et al. ${ }^{12}$ made use of the atomic masses. In addition to the atomic masses, Warner ${ }^{14}$ also performed several tests aiming at electron mass corrections in the framework of the $r_{s}$ structure.
In the present work, we found that the replacement of the atomic masses with the nuclear masses affects the equilibrium bond lengths in order of $10^{-4} \AA$. This effect is larger than the accepted statistical uncertainties of $10^{-5}$ $\AA$ for the $r_{e}$ parameters, such that the mass issue deserves some attention in molecular studies. We, however, also
observed that the equilibrium structure obtained by the employment of the atomic masses is statistically somewhat better than the nuclear-mass counterpart. Regarding concrete mathematical approaches to solve for the equilibrium structure, a nonlinear fit of all available data is recommened as more trustworthy than other possible variants.

## Dedication

This work is dedicated to William Klemperer. In 1970, Klemperer proposed the assignment of the X-ogen transition to $\mathrm{HCO}^{+}$, leading to the identification of this species as the first cation in the interstellar environment and marking the beginning of the fascinating field of interstellar chemistry.

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## APPENDIX A: ANALYTICAL SOLUTION FOR <br> BOND LENGTHS FROM TWO KNOWN MOMENTS OF INERTIA FOR THE CASE OF LINEAR TRIATOMIC MOLECULES

For linear triatomic molecules HXY, the equilibrium moment of inertia is explicitly

$$
\begin{equation*}
I_{e}=\mu_{r} r_{e}^{2}+\mu_{R} R_{e}^{2} \tag{A1}
\end{equation*}
$$

where the bond lengths $r_{e}, R_{e}$ are given by Eq. (7) and the reduced masses $\mu_{r}, \mu_{R}$ by Eq. (8). Letting $x=r_{e}($ HX) and $y=r_{e}$ (XY), Eq. (A1) is rewritten to

$$
\begin{align*}
I_{e} & =\mu_{r} x^{2}+\mu_{R}\left[x+\frac{m_{\mathrm{Y}}}{m_{\mathrm{XY}}} y\right]^{2}  \tag{A2}\\
& =\frac{m_{\mathrm{H}} m_{\mathrm{XY}}}{M} x^{2}+2 \frac{m_{\mathrm{H}} m_{\mathrm{Y}}}{M} x y+\frac{m_{\mathrm{Y}} m_{\mathrm{HX}}}{M} y^{2},
\end{align*}
$$

in fact

$$
\begin{equation*}
I=a x^{2}+b x y+c y^{2}, \tag{A3}
\end{equation*}
$$

using

$$
\begin{equation*}
a=\frac{m_{\mathrm{H}} m_{\mathrm{XY}}}{M}, \quad b=2 \frac{m_{\mathrm{H}} m_{\mathrm{Y}}}{M}, \quad c=\frac{m_{\mathrm{Y}} m_{\mathrm{HX}}}{M}, \tag{A4}
\end{equation*}
$$

and $M=m_{\mathrm{H}}+m_{\mathrm{X}}+m_{\mathrm{Y}}$.
The moment of inertia is the reciprocal of the rotational constant, Eq. (6). Knowing $B_{e}$ for two different isotopic forms of a linear triatomic molecule, two values $I_{e}^{(1)}$ and $I_{e}^{(2)}$ become available. The equilibrium distances $x$ and $y$ can then be calculated by solving the following system of equations

$$
\begin{align*}
& d_{1}=a_{1} x^{2}+b_{1} x y+c_{1} y^{2}, \\
& d_{2}=a_{2} x^{2}+b_{2} x y+c_{2} y^{2}, \tag{A5}
\end{align*}
$$

where $d_{1}=I_{e}^{(1)}$ and $d_{2}=I_{e}^{(2)}$. Multiplying the first equation by $b_{2}$ and substracting the second equation multiplied by $b_{1}$, we obtain
$\left(a_{1} b_{2}-a_{2} b_{1}\right) x^{2}=d_{1} b_{2}-d_{2} b_{1}+\left(b_{1} c_{2}-b_{2} c_{1}\right) y^{2}$.
Substituting into Eq. (A5), we find that

$$
\begin{align*}
\left(a_{1} b_{2}-a_{2} b_{1}\right) x y= & a_{1} d_{2}-a_{2} d_{1}  \tag{A7}\\
& -\left(a_{1} c_{2}-a_{2} c_{1}\right) y^{2} .
\end{align*}
$$

We square both sides of the equation and use Eq. (A6) to replace $x^{2}$, yielding

$$
\begin{equation*}
A y^{4}+B y^{2}+C=0, \tag{A8}
\end{equation*}
$$

where

$$
\begin{align*}
A= & \left(a_{1} b_{2}-a_{2} b_{1}\right)\left(b_{2} c_{1}-b_{1} c_{2}\right)+\left(a_{1} c_{2}-a_{2} c_{1}\right)^{2}, \\
B= & \left(b_{1} d_{2}-b_{2} d_{1}\right)\left(a_{1} b_{2}-a_{2} b_{1}\right) \\
& +2\left(a_{1} d_{2}-a_{2} d_{1}\right)\left(a_{2} c_{1}-a_{1} c_{2}\right)  \tag{A9}\\
C= & \left(a_{1} d_{2}-a_{2} d_{1}\right)^{2}
\end{align*}
$$

The two solutions for $y^{2}$ are thus

$$
\begin{equation*}
y_{1 / 2}^{2}=\frac{-B \pm \sqrt{D}}{2 A} \tag{A10}
\end{equation*}
$$

with the determinant $D=B^{2}-4 A C$ given explicitly by

$$
\begin{align*}
D=\left(a_{1} b_{2}-a_{2} b_{1}\right)^{2} & {\left[\left(b_{1} d_{2}-b_{2} d_{1}\right)^{2}\right.}  \tag{A11}\\
& \left.+4\left(a_{1} d_{2}-a_{2} d_{1}\right)\left(c_{1} d_{2}-c_{2} d_{1}\right)\right] .
\end{align*}
$$

The two solutions of Eq. (A10) for $y^{2}$ provide two results for $y$, taken as positive square roots since $y$ is a distance. For each possible $y$, we use Eq. (A6) to determine $x$ as a positive square root of $x^{2}$. The procedure may thus lead to two different solution pairs $(x, y)$. This is a consequence of the squaring of Eq. (A7), which may introduce new solutions. Direct substitution in Eq. (A5) should be used to verify which of the two pairs is appropriate as a solution. In the case that both solutions are eligible with respect to Eq. (A5), additional criteria are needed. For instance, resulting values for $x$ and $y$ may physically be inacceptable when $x$ and/or $y$ are too short or too long from the chemical point of view.

The front term $\left(a_{1} b_{2}-a_{2} b_{1}\right)$ in the determinant $D$ of Eq. (A11) is explicitly

$$
\begin{equation*}
a_{1} b_{2}-a_{2} b_{1}=2 \frac{m_{\mathrm{H}} m_{\mathrm{H}}^{\prime}}{M M^{\prime}}\left(m_{\mathrm{X}} m_{\mathrm{Y}}^{\prime}-m_{\mathrm{Y}} m_{\mathrm{X}}^{\prime}\right) \tag{A12}
\end{equation*}
$$

In the case of a terminal substitution $m_{\mathrm{H}} \rightarrow m_{\mathrm{H}}^{\prime}$, when $m_{\mathrm{X}}=m_{\mathrm{X}}^{\prime}$ and $m_{\mathrm{Y}}=m_{\mathrm{Y}}^{\prime}$, the term $\left(a_{1} b_{2}-a_{2} b_{1}\right)$ becomes zero, implying the solution for $y^{2}$ as

$$
\begin{equation*}
y_{1 / 2}^{2}=\frac{B}{2 A}=\frac{a_{1} d_{2}-a_{2} d_{1}}{a_{2} c_{1}-a_{1} c_{2}} . \tag{A13}
\end{equation*}
$$

The corresponding solution for $x^{2}$ follows from Eq. (A5). Taking the positive square roots of $x^{2}$ and $y^{2}$, we obtain a unique solution of Eq. (A5) for the case of the terminal substitution $m_{\mathrm{H}} \rightarrow m_{\mathrm{H}}^{\prime}$.


[^0]:    ${ }^{a}$ Fit 2 of Cazzoli et al. ${ }^{10}$
    ${ }^{b}$ Dore et al. ${ }^{11}$
    ${ }^{c}$ Bogey et al. ${ }^{12}$
    ${ }^{d}$ Lattanzi et al. ${ }^{13}$
    ${ }^{e}$ Warner. ${ }^{14}$
    ${ }^{f}$ Neese et al. ${ }^{20}$
    ${ }^{g}$ Dore et al. ${ }^{15}$

[^1]:    ${ }^{a}$ Caselli and Dore. ${ }^{16}$
    ${ }^{b}$ Hirao et al. ${ }^{21}$
    ${ }^{c}$ Lattanzi et al. ${ }^{13}$
    ${ }^{d}$ Dore et al. ${ }^{15}$

[^2]:    ${ }^{a}$ Amano and Maeda. ${ }^{17}$
    ${ }^{b}$ calculated from the transition $J=0 \rightarrow 1$ observed by Gudeman and Woods. ${ }^{18,19}$
    ${ }^{c}$ Nakanaga and Amano. ${ }^{22}$

