Anomalous isotopic effect on vibrational properties of HXeOH

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(Received 8 January 2002; accepted 22 January 2002)

We study IR absorption spectra of various isotopomers of the HXeOH molecules isolated in solid Xe at 7 K. Upon the $^{16}$O to $^{18}$O isotopic substitution, we found experimentally an anomalous shift of the H–Xe stretching vibration frequency of HXeOD and HXeOH. This spectral behavior, when the fundamental frequency shifts up in energy for the heavier oxygen isotope, cannot be understood on the basis of the harmonic approximation for fundamental vibrations, and the anharmonic coupling between normal modes is presumably responsible for this anomalous effect. Our anharmonic CC-VSCF calculations describe correctly the experimental observations. © 2002 American Institute of Physics. [DOI: 10.1063/1.1459703]

Vibrational spectroscopy is a sensitive probe of potential energy surfaces (PES) that determine dynamics of nuclear motion in molecules. The analysis of vibrational spectra allows prediction of structural and thermodynamical properties of matter. In addition to experimental data, theoretical analysis based on electronic structure theory is valuable here. For small molecules, one can obtain an accurate analytical fit of a multidimensional PES by using extensive ab initio electronic structure calculations and apply it then in theoretical vibrational spectroscopy. For more complicated systems, the harmonic vibrational approximation is usually employed, but this does not provide often deep enough insight. In order to extend the applicability of theoretical analysis, an anharmonic method has recently been developed, which combines electronic ab initio electronic structure calculations and its extension employing perturbation theory (CC-VSCF). In this approach, interactions between pairs of normal modes are taken into consideration, and interactions of triples or higher are neglected.

A large number of novel charge-transfer HRgY molecules (H=hydrogen; Rg=Ar, Kr, Xe; Y=an electronegative fragment) have been characterized by using experimental vibrational spectroscopy in rare-gas solids and ab initio theoretical methods. From the energetic point of view, these molecules are quite unusual due to the low dissociation energy (~0.4–1.5 eV), and we were able to photodecompose one of them (HxeI) directly by exciting the second H–Xe stretching overtone at ~3000 cm$^{-1}$ with intense narrow-band IR radiation. The shallowness of the potential well suggests the importance of anharmonic effects in their ground-state dynamics. It follows that one should carefully account the anharmonicity of various normal modes and the anharmonic couplings between the modes when treating these molecules theoretically. We have recently applied the CC-VSCF approach to HxeI, and a number of exceptionally strong combination and overtone bands predicted theoretically have been found in the experimental spectra. The HRgY molecules are interesting in many other respects as well. For instance, they constitute an important intermediate during photodissociation of HY molecules in rare-gas hosts influencing the local photodissociation dynamics, and this process has recently been modeled theoretically for photodissociation of HCl in Xe clusters.

The anharmonic character of the HRgY molecules suggests search for exceptional effects in their ground-state dynamics. On one hand, we should precisely look at changes of the vibrational properties upon isotopic substitution because the anharmonic mode coupling might be very essential here. On the other hand, the isotopic substitution can test adequateness of the theory employed to obtain the multidimensional PES. The “trivial” harmonic isotopic effect usually considered in spectroscopic studies is a downshift of the fundamental vibrational frequencies when the reduced mass increases. How general is this rule? This question is, in addition to purely fundamental interest, of importance for practical vibrational spectroscopy. In particular, the heavy-atom substitution has been successfully employed in many studies, from laboratory identification of unusual species to astrophysics analysis. One experimental gas-phase work reports a small anomalous shift (~0.05 cm$^{-1}$) of the (CIHCl)$^-$ anion fundamental frequency upon $^{35}$Cl to $^{37}$Cl substitution, and their tentative interpretation was based on symmetry changes. In the present study, we found experimentally and interpreted theoretically the anomalous shifts of H–Xe stretching vibrational frequencies of HXeOH and HXeOD upon oxygen isotopic substitution when the fundamental frequencies increased with an increase of mass. We demonstrate that it is impossible to interpret this effect on the basis of harmonic approximation, and the use of anharmonic theory is crucial for qualitatively correct insight.

The experimental procedure to prepare the HRgY molecules includes UV solid-state photodissociation of a HY precursor followed by thermal mobilization of hydrogen atoms in the rare-gas lattice. In order to prepare H$_2$O/Xe solid mixtures in the present experiments, we co-deposited water...
vapor (H$_2$O, D$_2$18O, H$_2$18O, and D$_2$18O) and xenon gas from two separate nozzles onto a CsI window. The deposition temperature was 30 K, the deposition time was about 1 h, and the typical sample thickness was 100 μm. The measurements were performed at temperatures down to 7 K in a closed-cycle helium cryostat (DE-202A, APD). The IR absorption spectra in the 4000 to 400 cm$^{-1}$ region were recorded with a Nicolet 60 SX FTIR spectrometer in most cases by using resolution of 1 cm$^{-1}$ and co-adding 500 scans. The as-deposited H$_2$O/Xe samples are quite monomeric, and no complexes with normal matrix impurities like N$_2$ are visible. In the experiments with isotopically substituted water, ordinary water (H$_2$18O) always interferes in the spectra with a typical ratio of ~1/3. In order to decompose water molecules to H+OH, the H$_2$O/Xe solid mixtures were irradiated with an excimer laser (MPB, MSX-250) operating at 193 nm. Typically, we used in photolysis ~2×10$^3$ pulses with a pulse energy density of ~10 mJ/cm$^2$. Under irradiation at 193 nm, water decomposes and the IR absorption band of isolated OH radicals appears in the spectra. Upon annealing of a photolyzed sample at ~40 K, hydrogen atoms become globally mobile, and HXeH (1166 and 1180 cm$^{-1}$) and HXeOH (1577 cm$^{-1}$) are formed (see Fig. 1). For the D-enriched samples, DXeH (1094 and 1121 cm$^{-1}$), DXeD (846 and 857 cm$^{-1}$), HXeOD (1572 cm$^{-1}$), DXeOH (1149 cm$^{-1}$), and DXeOD (1141 cm$^{-1}$) are known to rise in annealing. It should be mentioned that these Xe-containing molecules exhibit some matrix site structure as seen in Fig. 1, and our data below were obtained for the strongest site bands.

In the experiments, we carefully analyzed the changes of the H–Xe stretching frequency of various HXeOH isotopomers concentrating on the effect of oxygen isotopic substitution. The IR absorption bands were fitted with Gaussian functions and appropriate backgrounds. Most importantly, the H–Xe stretching frequency of HXeOH and HXeOD is found to be lower for the lighter oxygen isotope (see Table I and Fig. 2 for HXeOD), which is considered as anomalous behavior. For other isotopomers, oxygen isotopic substitution in DXeOH produces no measurable shift of the D–Xe stretching mode, and the effect on the D–Xe stretching mode of DXeOD is quite large (~3 cm$^{-1}$) and has the normal direction down in energy for the heavier mass.

The reliability of the experimentally detected anomalous shifts should be discussed because the bandwidths in question (~2 cm$^{-1}$) are rather larger than the observed anomalous shifts. However, the experimental results are completely repeatable. In order to demonstrate this repeatability, we list the actual fitting realizations for the H–Xe stretching band centers of HXeOD. For the 16O isotope, they are 1572.18, 1572.15, 1572.13, and 1572.11 cm$^{-1}$ whereas for the 18O-enriched samples, they are 1572.40, 1572.43, 1572.41, and 1572.39 cm$^{-1}$. The typical error given by the least-square fitting procedure is 0.02 cm$^{-1}$. In order to estimate the possible influence of local matrix morphology on the vibrational frequency, we analyzed the absorption frequency of the oxygen-free HXeD molecule in the same experiments, and the typical results are 1577 cm$^{-1}$ whereas for the 18O-enriched samples, they are 1577 cm$^{-1}$, HXeOD, HXeOD 1547, HXeOD 1577, HXeOD 1577, and DXeOD 1141 cm$^{-1}$, i.e., similarly very repeatable. Thus, we feel confident in the main conclusion of the present work that the H–Xe stretching mode of HXeOD is reliably higher in energy than that of HXeOD. In average, this difference is measured to be +0.27 cm$^{-1}$. However, the true band shift might be even somewhat larger than the measured one due to

![FIG. 1. IR absorption spectra of Xe-containing molecules prepared from water and Xe. The H$_2$O/Xe solid mixtures were first irradiated at 193 nm and then annealed at 45 K. The spectra were measured at 7 K. The smooth background line is introduced by interference of light in the thin matrix layer.](image)

| TABLE I. Experimental and theoretical shifts of the H(D)–Xe stretching frequencies (in cm$^{-1}$) of various H/D isotopomers of HXeOH upon 16O to 18O substitution. For comparison, the shift of the H–N stretching frequencies of HNC upon 12C to 13C substitution is analyzed. |
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| Species | Experiment$^a$ | CC-VSCF theory | Anharmonic theory without coupling | Harmonic theory |
| HXeOH | 1577.65 | +0.13 | 1720.78 | +0.14 | 1762.85 | −0.03 |
| HXeOD | 1572.14 | +0.27 | 1717.61 | +0.30 | 1760.03 | −0.03 |
| DXeOH | 1149.14 | −0.03 | 1250.01 | +0.04 | 1271.84 | −0.10 |
| DXeOD | 1141.23 | −0.30 | 1247.60 | −1.28 | 1266.90 | −0.06 |
| HNC | 3643.1$^b$ | −0.6$^b$ | 3634.88 | −0.40 | 3690.83 | −1.21 |

$^a$The experimental error obtained from the fitting inaccuracy and data distribution is smaller than (+0.04).

$^b$The H–12C stretching frequency and the shift upon 12C to 13C substitution from Ref. 17 (experiment in solid neon).
overlap between the HXe\textsuperscript{16}OD and HXe\textsuperscript{18}OD bands in the experiments with D\textsubscript{2}\textsuperscript{18}O. In fact, we did not mix D\textsubscript{2}\textsuperscript{18}O and H\textsubscript{2}\textsuperscript{18}O on purpose, and H atoms in the D\textsubscript{2}\textsuperscript{18}O experiments originate simply from residual H\textsubscript{2}\textsuperscript{16}O that is always present in the samples as a natural impurity. It means that the Xe-containing products with the \textsuperscript{16}O isotope always form in some amounts. The HXe\textsuperscript{16}OD/HXe\textsuperscript{18}OD proportion is not precisely known, and no reliable deconvolution is possible for such a small frequency difference and a decent signal-to-noise ratio. Furthermore, the bands are broadened limiting the optimal experimental resolution to 1 cm\textsuperscript{-1}. We can mention here that the measured band centers shift down in energy when in some experiments the initial H\textsubscript{2}\textsuperscript{18}O/H\textsubscript{2}\textsuperscript{16}O ratio decreases, and this supports the proposed image. The measured anomalous difference of +0.13 cm\textsuperscript{-1} between the H–Xe stretching vibrations of HXe\textsuperscript{18}OH and HXe\textsuperscript{16}OH has a high level of confidence as well.

The structure, energy and vibrational properties of the present Xe-containing molecules have been calculated by the CC-VSCF method. The approach in this work is of the same level of theory as used recently to calculate the anharmonic vibrational spectra of a number of HRgY molecules.\textsuperscript{16} As emphasized earlier, HXeOH is quite anharmonic as evidenced, for instance, by the calculated anharmonicity 2 × \nu\textsubscript{2}−2 \nu\textsubscript{3} = −80 cm\textsuperscript{-1} where \nu\textsubscript{2} is the H–Xe stretching mode and 2 \nu\textsubscript{2} denotes its first overtone. In this situation, the agreement between experimental and calculated fundamental vibrational frequencies can be essentially improved by accounting this property (see Table I) even though the result is not of spectroscopic accuracy yet. We also tested the CC-VSCF method for the \textsuperscript{12}C to \textsuperscript{13}C isotopic substitutions in HNC molecules comparing the theoretical results with the experimental data available for this molecule.\textsuperscript{17} The idea of this test is to evaluate the CC-VSCF method for the heavy atom substitution in respect with fine perturbation of the H–N stretching mode. Importantly, HNC is also quite anharmonic as judged by the experimental H/D ratio of 1.311 for the H–N stretching vibration. Indeed, the harmonic approximation applied to HNC overestimates strongly the H–N stretching vibration energy although the isotopic shift is properly described meaning that the reduced-mass effect still dominates in the isotopic shift of this molecule. The anharmonic approach improves the quantitative agreement with the experiment with respect to the fundamental frequencies but it does not feature for HNC qualitatively different results.

Our CC-VSCF results fully support the spectral observations on HXeOH (see Table I). According to this theory, the H–Xe stretching fundamental vibrations of HXeOH and HXeOD exhibit anomalous up-shifts upon substitution with the heavier oxygen isotope, and the computed shifts agree well with the experiment. In accordance with the experiments, practically no shift for the H–Xe stretching mode of DXeOH and a relatively large downshift for this mode of DXeOD was computationally obtained upon the \textsuperscript{16}O to \textsuperscript{18}O substitution. One can see that the harmonic approximation predicts very small but definite downshifts upon substitution with the heavier oxygen isotope in all cases, which qualitatively disagrees with the experiment. This is a case when the harmonic approximation gives qualitatively wrong results. One can speculate, in principle, that the decrease of zero point energy produced by the heavier oxygen isotope makes the molecule more harmonic, which shifts the band up in energy in the case of negligible reduced-mass effect, i.e., it is a purely anharmonic effect. This trend for anharmonicity was found theoretically: the calculated anharmonicities of the H–Xe stretching mode are 80.64 and 80.36 cm\textsuperscript{-1} for HXe\textsuperscript{16}OD and HXe\textsuperscript{18}OD, respectively. However, this change in anharmonicity is insufficient to produce the anomalous shift. Indeed, the anharmonic calculations without accounting for mode couplings are equally wrong (see Table I).

We interpret that the anharmonic mode coupling has a decisive contribution for the anomalous isotopic shifts observed in the present work. The isotopic substitution changes the coupling between normal modes, which changes the vibrational frequencies via the modified coupling coefficients. The contribution of mode coupling to the Xe–H stretching fundamental is computationally quite large (∼40 cm\textsuperscript{-1} for HXeOH), and its modification upon oxygen isotopic substitution is the reason for the obtained anomalous spectral effect. The analysis for HXeOH shows that the coupling potentials of the Xe–H stretching mode (\nu\textsubscript{2}) with the asymmetric bending mode (\nu\textsubscript{3}) and the Xe–OH stretching mode (\nu\textsubscript{6}) change essentially upon the \textsuperscript{16}O to \textsuperscript{18}O isotopic substitution. In particular, the coupling of \nu\textsubscript{2} with \nu\textsubscript{3} is decreased and the coupling of \nu\textsubscript{2} with \nu\textsubscript{6} is increased. It should be also noted that the in-plane bending modes are strongly coupled with the OH stretching mode and they undergo large changes upon oxygen isotopic substitution as well. Of course, the resulting effect of mode coupling can be also of the “normal” direction as we see for DXeOD, and the mode.
coupling essentially establishes it as well. It should be mentioned that we have explained the anomalous shift in a different way compared with Ref. 12 because their mechanism of triple mode interaction induced by the perturbed $D_{sh}$ symmetry is not included into the CC-VSCF model.\(^2\)\(^3\)

The relatively large mode coupling in HXeOH might be connected with the mixed character of its bonding where both ionic and neutral potentials contribute. This Xe-containing molecule dissociates in an unusual way: The detachment of H destroys the chemical character of the Xe–O bond as well, which suggests strong coupling between the H–Xe and Xe–O stretching coordinates. It should be emphasized that this mode-coupling contribution always competes with the reduced-mass effect, and the anomalous result can occur only when the latter effect is small. It is exactly the case of the Xe-containing species due to the large Xe mass, and it can be particularly seen in a very small shift upon oxygen isotopic substitution obtained in the harmonic approximation.

In conclusion, we found experimentally an anomalous shift in the H–Xe stretching vibrational frequency of matrix-isolated HXeOD and HXeOH molecules upon oxygen isotopic substitution when the frequency shifts up in energy for the heavier isotope. This behavior cannot be understood in the framework of harmonic vibrational theory, and the anharmonic coupling between normal modes must be employed to obtain the adequate image. Our anharmonic CC-VSCF calculations describe correctly the experimental observations. This result emphasizes that the effect of anharmonic mode coupling is generally important in vibrational spectroscopy, and it influences vibrational frequencies of various molecules with coupled normal modes.

The Academy of Finland supported this work. R. B. Gerber is thanked for valuable discussions.