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ABSTRACT
The IR spectra of liquid hydrogen isotopologues (Q2 = H2, D2, T2, HD, HT, DT) are dominated by the interaction induced absorption. Therefore, the complexity tremendously increases with the number of different isotopologues in the sample. As we aim for a system independent calibration of IR absorption spectroscopy against all six isotopologues and three ortho–para ratios, we need a minimal and complete set of descriptors to predict the spectra and to decrease the needed calibration effort. For this, we grouped the absorption lines into three groups: absorption on monomers, phonons, and molecular dimers. In particular, molecular dimers contribute to the absolute number of absorption lines in the spectra of mixed isotopologues. To develop and test the set of descriptors, we make use of three spectra: a pure H2 sample, a pure D2 sample, and a mixed H2–D2 sample. We show a detailed analysis of these three spectra in the first and second vibrational branch in the range from 2000 cm−1 to 9000 cm−1. The set of descriptors found within this work can be used to identify and predict all lines in this range for liquid H2–D2 mixtures.

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I. INTRODUCTION
The development of analytic systems and methods for tritium and other hydrogen isotopologues is a main research field at the Tritium Laboratory Karlsruhe (TLK). One of the methods under development is IR absorption spectroscopy for the concentration measurement of liquid hydrogen isotopologues at temperatures between 15 K and 25 K. This method can be used for in-line and real-time process monitoring of cryogenic distillation columns or the measurement of the ortho/para ratio of the homonuclear hydrogen isotopologues (H2, D2, and T2).

We are aiming for a full calibration of the IR absorption spectra of liquid hydrogen isotopologue mixtures against all six concentrations (Q2 = H2, D2, T2, HD, HT, DT) and the three ortho/para ratios. The first calibration for H2, D2, and HD mixtures with fixed ortho/para ratios leads to a total accuracy of better than 5%. Previous studies of absorption spectra of liquid H2 and D2 for different ortho/para ratios and temperatures show a strong influence of these parameters on the first vibrational branch. This limits the trueness of our calibration. Additionally, for isotopologue mixtures, molecular dimers do complicate the IR absorption spectra tremendously. For the interpretation of the experimental data, we aim for a minimal and complete set of descriptors for the IR spectra of liquid hydrogen isotopologue mixtures. This will be crucial for the upcoming measurement campaigns with the tritium compatible setup that is under construction at the TLK, where we plan to extend this calibration to all six hydrogen isotopologues and variable ortho/para ratios. To extract the minimal and complete set of descriptors, we studied the IR absorption spectra of pure H2 and D2 liquid samples, and a H2–D2 mixture.

II. THEORETICAL OVERVIEW
As diatomic molecules, the hydrogen isotopologues undergo rotational and vibrational excitations by the absorption of an infrared photon.

The hydrogen isotopologues follow the vibrational selection rule of Δν ≥ 0. Δν = 0 transitions create the pure rotational transition...
spectrum; $\Delta \nu = 1$ constitutes the common ro-vibrational “fundamental absorption” spectrum; higher vibrational excitation $\Delta \nu \geq 2$ represent the “overtone bands” with transition probabilities at least one to two orders of magnitude lower than that for $\Delta \nu = 1$. Only the (small) anharmonicity in the molecular potential makes these transition probabilities non-vanishing.

The rotational selection rules of the hydrogen isotopologues for IR absorption differ from the general selection rules for diatomic molecules due to the ortho/para modifications of the homonuclear hydrogen molecules. Infrared absorption follows the rotational selection rule of $\Delta J = 0$, $\pm 1$ for diatomic molecules. For homonuclear hydrogen isotopologues ($H_2$, $D_2$, $T_2$), the $\Delta J = \pm 1$ transition is forbidden due to the forbidden ortho ↔ para transition. For the heteromolecular hydrogen isotopologues (HD, HT, and DT), these transitions are suppressed by the vanishing inertia tensor along the main axis $\theta$.\textsuperscript{9,10}

This implies that only “pure vibrational” transitions ($\Delta \nu \geq 1$; $\Delta J = 0$) could be observed, without rotational companion lines with $\Delta J \neq 0$. However, experimental observations contradict this by exhibiting clear rotational S-branch lines;\textsuperscript{11–13} for an explanation of the related selection rule $\Delta J = \pm 2$, see further below.

Theory discusses different mechanisms to solve this contradiction; van Kranendonk gave the first overview on this in 1957, which was followed by Ref. 8, 19, and 20. These mechanisms are mainly based on the van der Waals interactions between hydrogen molecules and enable the formation of oligomers. Parameters of these bonds, such as binding potentials, are investigated to the present day.\textsuperscript{9,12,21–24} Already the simplest oligomer, the dimer, offers an additional rotational and vibrational quantum number\textsuperscript{21,23} to explain the rotational splitting in the IR absorption spectra and the observed selection rules of $\Delta \nu \geq 0$ and $\Delta J = 0$, ±2.

### III. EXPERIMENTAL STRATEGY

For the present work, we studied three different samples in the liquid phase: pure $H_2$, pure $D_2$, and a mixture of both (see Table I).

At low temperatures of about 20 K, almost only the ground state of the molecules is expected to be occupied. Due to the forbidden ortho ↔ para transition of the homonuclear isotopologues, the sum of the odd J states and the sum of even J states are almost constant in time. Therefore, the first excited rotational state $J = 1$ is occupied in the case of a sufficiently fast cool down. The $J_1$ to $J_0$ ratio is 3:1 for $H_2$ (6:3 for $D_2$) directly after the liquefaction.\textsuperscript{25} This ratio changes by natural equilibration, which can be neglected due to its long conversion times (>days\textsuperscript{26,27}). For this work, the precise ortho/para ratio is not needed for the discussion. However, it is a major systematic uncertainty for the calibration of an IR absorption based concentration-measurement and requires a separate treatment.\textsuperscript{26,27} Higher energetic J states ($J > 1$) are negligibly occupied at the low temperatures of the liquid phase, therefore no $\Delta J = \pm 2$, ±3 transitions are expected for both homonuclear and heteronuclear hydrogen isotopologues.

The present study does not use the common method of comparison for calculation with the measured line position, which is limited to the broad lines in the liquid phase (see Sec. V A). Instead, we use a twofold approach. First, we check for missing lines predicted by the current set of descriptors to verify or falsify these. Second, we investigate lines in a $H_2$–$D_2$ mixture that are not present in pure $H_2$ or pure $D_2$ spectra (see Table I for the composition of all investigated samples). HD as the origin of these additional lines is ruled out by a reference measurement by Raman spectroscopy in the gas phase before the liquefaction and after the re-vaporization of the liquid sample having completed the IR-measurements. Therefore, these additional lines must be associated with heteromolecular hydrogen dimers. A rough analysis of the line positions of these lines also makes it possible to verify or reject specific selection rules.

### IV. EXPERIMENTAL DETAILS

As described by Größle et al.,\textsuperscript{2} the hydrogen isotopologues are liquefied in an infrared sample cell with a volume of 100 ml and an absorption length of 4.5 cm. The sample cell operates at a pressure up to 3.5 bars and is equipped with sapphire view-ports (diameter 3.2 cm). The sample cell is cooled by gaseous helium provided by a helium refrigeration system with a cooling power of 250 W at 16 K helium temperature. It is surrounded by an insulation vacuum container with a pressure $<10^{-4}$ mbar, which is also equipped with sapphire view-ports.

The sample temperatures are between 19 K and 21 K. Infrared absorption spectra are obtained with a Bruker Tensor 27 FT-IR spectrometer with a resolution of 0.9 cm\textsuperscript{-1} combined with an external liquid nitrogen cooled HgCdTe detector.

#### A. Measurement description

The typical measurement consists of several steps. First, a pure $H_2$, $D_2$, or a mixed $H_2$–$D_2$ gas sample is prepared. In the case of a mixed sample, its composition is measured by Raman spectroscopy as described in Refs. 28 and 29. The isotopologue concentrations of the samples of the present analysis are listed in Table I. Second, the sample cell is cooled below the boiling point of the gas mixture and the condensation process starts. Third, the sample cell is constantly cooled and filled with liquid by condensation, which takes about 8 h. Fourth, IR absorption spectra are taken for up to 30 min. Fifth, the liquid is evaporated and the sample cell is evacuated to take IR reference spectra with the empty sample cell.

For the mixed $H_2$–$D_2$ sample, another Raman measurement is taken in the gasphase. By this, the formation of HD during the measurement process can be monitored and in this case excluded.

#### TABLE I. Compositions of the investigated hydrogen samples. Samples a and b are pure samples; the indicated concentrations are the manufacturer’s data. The $D_2$ calibration gas has an atomic purity of 99.7%, which corresponds to a fraction of 0.6% HD. Sample c has been mixed within the experimental setup. Since the ideal gas law has been used and the liquefaction process influences the composition, it is difficult to deduce accurate values; the actual composition has been measured by Raman spectroscopy.

| Sample | $c_{H_2}$ | $c_{HD}$ | $c_{D_2}$ |
|--------|-----------|----------|-----------|
| A      | $D_2$     | 0        | 0.006     | 0.994     |
| B      | $H_2$     | 1        | 0         | 0         |
| C      | $H_2 + D_2$ | 0.429   | 0.003     | 0.568     |
B. Data analysis

The recorded infrared absorption spectra are evaluated in four steps, as described by Größle et al.\textsuperscript{2} The first step is a Fourier transformation and corrections to the 15–25 recorded sample and reference interferograms applied by the OPUS software from Bruker. Second, the recorded transmittance spectra are normalized to their intensity at 2500 cm\(^{-1}\) and treated with an elliptical rolling circle filter as described by Mikhailjov and Razhivin.\textsuperscript{50} We adapted this filter procedure by taking values for independent filter radii for the wavenumber \(\tilde{\nu}\) and intensity \(I\) as \(r_1 = 2500\ \text{cm}^{-1}\) and \(r_2 = 2.5\), respectively. This filter step removes the influence from background effects, such as the varying detector sensitivity, source fluctuations, and the influence of the surface reflection between the sample and the sapphire window. Third, each of the sample spectra is divided by the associated reference spectra, the resulting transmittance spectra are averaged, and the 68.2\textsuperscript{th} percentile is calculated as an estimation for the square root of the variance. Fourth, the decadic absorbance \(A = -\log_{10}(T)\) is calculated from the transmittance \(T\) and spectral lines are integrated over the intervals listed in Table III.\textsuperscript{51}

C. Calculation of line positions and occupational numbers

For the interpretation of the recorded spectra, the molecular energy levels are calculated based on molecular constants for the gas phase published by Huber and Herzberg and the calculation given in their publication.\textsuperscript{52} The transitions are then derived as the energy difference between the initial and the final states, where the initial state is always in the vibrational ground state (\(v = 0\)) and in the rotational ground state (\(J = 0\)) or the first excited state (\(J = 1\)). The final state is then selected based on the initial state by the selection rules \(\Delta\nu \geq 0\) and \(\Delta J = 0, 2\). This approximation is valid under the assumption that the energy levels and, therefore, the rotational and vibrational quantum numbers of hydrogen molecules in the gaseous phase are still valid in the liquid phase and the corrections are small compared to the expected line width.\textsuperscript{32} The energies of dimer transitions are calculated as the combination of the two monomer transitions.\textsuperscript{33} Additional excitation energies of a possible dimer rotation are neglected.

Transitions are shortly denoted as \(\Delta J\nu_{\text{v}}(J''')\), where \(\Delta J\) is the rotational excitation expressed by letters \(Q, R,\) and \(S\) (\(Q\) corresponds to \(\Delta J = 0, R\) to \(\Delta J = 1,\) and \(S\) to \(\Delta J = 2\)) and \(\nu_{\text{v}}\) is the initial rotational state. \(\Delta\nu\) represents the vibrational excitation, where the initial vibrational state is \(\nu = 0\).

At the cryogenic temperatures encountered here, no measurable population is found in any higher vibrational state. The occupational numbers can be calculated by the energy level of the initial states and the Boltzmann distribution, where one has to take into account the spin degeneracy of the nuclei. This is 3:1 (odd \(J''\), even \(J''')\) for H\(_2\) and 6:9 for D\(_2\). In addition, the slow conversion between ortho and para has to be considered.\textsuperscript{53} This is done by executing the Boltzmann distribution in two iterations. First, with a sample temperature that corresponds to the ortho–para equilibrium. We used 300 K. From this first iteration one will get the ortho–para ratio (sum of the odd or even states), here, almost the nuclear spin degeneracy for the corresponding isotopologue. This ratio is then maintained, and the Boltzmann distribution is executed a second time. This time separately for the odd states and then for the even states, where the normalization factor for the odd and the even states is used from the first iteration. From this, one simulates that the transitions only occur within the even or within the odd states during the cool down from room temperature to around 20 K.

From this, significant population is only found for the rotational levels \(J'' = 0\) and \(J'' = 1\). In addition, the transition lines are indexed (as superscript) with the associated isotopologue, equally for monomers and dimers. For example, \(S_{02}^{12}(1)Q_{22}^{22}(0)\) indicates a mixed isotopologue dimer exhibiting an ortho-S-transition (on H\(_2\)) and an ortho-Q-transition (on D\(_2\)).\textsuperscript{24} Selected strong transitions are summarized in Tables II and III, with a full list of all calculated transitions provided in the supplementary material.

### Table II. Overview of the dominant transitions in the fundamental vibrational band of liquid D\(_2\) (sample a), H\(_2\) (sample b), and mixtures (sample c). The dominant transitions are given with their calculated position \(\nu_0\) and calculated occupational number \(N(I)\). In the first vibrational branch, the transitions cannot be spectrally separated, so calculating the intensities and the center of gravity is strongly dependent on the selection of integration ranges.

| Transition          | \(\nu_0\) (cm\(^{-1}\)) | \(N(I)\) |
|---------------------|---------------------------|----------|
| Sample (a)          |                           |          |
| \(Q_{12}^{22}(1)\)  | 2989.7                    | 0.33     |
| \(Q_{12}^{22}(0)\)  | 2991.9                    | 0.67     |
| \(S_{02}^{12}(0)Q_{22}^{22}(1)\) | 3168.7 | 0.44 |
| \(S_{02}^{12}(0)Q_{22}^{22}(0)\) | 3170.9 | 0.89 |
| \(S_{02}^{12}(1)Q_{22}^{22}(1)\) | 3287.1 | 0.22 |
| \(S_{02}^{12}(1)Q_{22}^{22}(0)\) | 3289.3 | 0.44 |
| Sample (b)          |                           |          |
| \(Q_{12}^{42}(1)\)  | 4152.4                    | 0.75     |
| \(Q_{12}^{42}(0)\)  | 4158.6                    | 0.25     |
| \(S_{02}^{12}(0)Q_{22}^{22}(1)\) | 4506.7 | 0.38 |
| \(S_{02}^{12}(0)Q_{22}^{22}(0)\) | 4512.8 | 0.13 |
| \(S_{02}^{12}(1)Q_{22}^{22}(1)\) | 4739.1 | 1.12 |
| \(S_{02}^{12}(1)Q_{22}^{22}(0)\) | 4745.2 | 0.38 |
| Sample (c)          |                           |          |
| \(S_{02}^{12}(0)Q_{22}^{22}(1)\) | 3343.9 | 0.04 |
| \(S_{02}^{12}(0)Q_{22}^{22}(0)\) | 3346.1 | 0.08 |
| \(S_{02}^{12}(1)Q_{22}^{22}(1)\) | 3576.3 | 0.12 |
| \(S_{02}^{12}(1)Q_{22}^{22}(0)\) | 3578.5 | 0.25 |
| \(S_{02}^{12}(0)Q_{22}^{22}(1)\) | 4331.4 | 0.25 |
| \(S_{02}^{12}(0)Q_{22}^{22}(0)\) | 4337.6 | 0.08 |
| \(S_{02}^{12}(1)Q_{22}^{22}(1)\) | 4449.9 | 0.12 |
| \(S_{02}^{12}(1)Q_{22}^{22}(0)\) | 4564.0 | 0.04 |
A. Spectral resolution and line broadening

The spectral resolution of our setup is 0.9 cm\(^{-1}\) according to the manufacturer. In the gas phase,\(^{17}\) this is enough to resolve almost all ro-vibrational spectral features for H\(_2\), D\(_2\), and mixtures. In the liquid phase, there is a severe line broadening due to which many spectral features cannot be resolved. A comparison of water vapor absorption lines and those of liquid H\(_2\) can be found in the supplementary material. This demonstrates that the resolution of our setup is not the limiting factor to resolve all the spectral features, but it is limited due to the line-broadening in the liquid phase.

In the end, our analysis shows that it is not possible to separately analyze Q\(_3\)\(_0\)\(_0\) and Q\(_3\)\(_0\)\(_1\) since the difference between these lines is equal to or much smaller than the typical linewidth. For D\(_2\), Q\(_2\)\(_1\) and Q\(_3\)\(_0\) are separated by 2.2 cm\(^{-1}\), and at \(\Delta v = 2\), they are separated by 4.3 cm\(^{-1}\). For H\(_2\), the line-spacing is 6.2 cm\(^{-1}\) in the first vibrational branch and 12.2 cm\(^{-1}\) in the second. This is due to the low rotational anharmonicity contribution and could be resolved in the gas phase. For the liquid phase, these branches cannot be separated easily. However, the separation for H\(_2\) is big enough to deconvolute the two contributions, especially when investigating varying ortho–para ratios, which is beyond the scope of this paper. Therefore, we indicate these lines by denoting Q\(_3\)\(_1\)\(_0\) and Q\(_3\)\(_1\)\(_1\) as Q\(_3\)\(_1\)\(_0\)\(_0\) and Q\(_3\)\(_1\)\(_1\)\(_0\). At the same time, S\(_3\)\(_0\)\(_0\) and S\(_3\)\(_0\)\(_1\) are well-separated due to the rotational energy being proportional to J(J + 1) in the first order.

B. First vibrational branch

1. First vibrational branch of H\(_2\) and D\(_2\)

The first vibrational branch of D\(_2\) (2900–4000 cm\(^{-1}\)), shown in Fig. 1, can be sufficiently described by a combination of monomer,
FIG. 1. IR absorption spectra of liquid hydrogen isotopologues (first vibrational branch). (a) D$_2$, (b) H$_2$ (with an offset of 0.05), and (c) H$_2$ + D$_2$ mixture (see Table I). Calculated line positions are indicated: solid lines represent dominant transitions, dotted lines represent suppressed quadrupole monomer S transitions, and dashed–dotted lines represent suppressed dimer double-S transitions.

dimer, and phonon transitions. For the absorbance spectra, see Fig. 1 and Table II for a list of calculated dominant transitions.

The pure vibrational monomer transitions $Q_1(0, 1)$ give the lowest energetic lines close to 3000 cm$^{-1}$. A 50–100 cm$^{-1}$ higher wavenumber exhibits the phonon transition [labeled $Q_{1,R}(0, 1)$]. This broad line is followed by the combined rotational and vibrational transitions labeled as $S_0(0)Q_1(0, 1)$ (at 3200 cm$^{-1}$) and $S_0(1)Q_0(0, 1)$ (at 3300 cm$^{-1}$). These absorption lines demonstrate again that the ortho ($J = 0$) and para ($J = 1$) transitions split into well-separated lines only if a rotation is located on the particular molecule. In a pure vibrational excitation, these lines are not separately visible. In addition, the monomer $S_1(0)$ and the corresponding dimer transition $S_0(0)Q_1(0, 1)$ [and as well for $S_1(1)$] are not separately visible in the first vibrational branch. This is caused by the high decadic absorbance of more than 2.5, which exceeds the dynamic range of our setup and the liquid phase that broadens the lines (see also Sec. V A).

In addition, the spectrum shows four dimer double-S transitions whose integral absorbance is one to two orders of magnitude smaller than for the aforementioned transitions. First, the line at
FIG. 2. IR absorption spectra of liquid hydrogen isotopologues (second vibrational branch). (a) D₂, (b) H₂ (with an offset of 0.05), and (c) H₂ + D₂ mixture. Annotations as in Fig. 1.

about 3350 cm⁻¹ is a double-S excitation of a D₂ ortho (J = 0) dimer S₁(0)S₀(0). Second and third, at approximately 3450 cm⁻¹, there are the two excitations of an ortho/para dimer [S₁(1)S₀(0)] and [S₁(0)S₀(1)]. The splitting of these lines is caused by the location of the vibrational excitation on either the ortho or on the para molecule. Fourth, the transition at ~3550 cm⁻¹ corresponds to the para–para dimer S₁(1)S₁(1). The integral absorbance of these four double-S transitions is much lower than that of the single-S transitions. However, giving precise values is difficult since they are located on the high energy flank of the S₀(1)Q₁(0, 1) line, which leads to a strong curvature of the background for the double-S transitions. However, when aiming for an ortho–para calibration, one should look deeper into these four transitions since here the transitions corresponding to para–para, ortho–para, and ortho–ortho dimers are well-separated from each other.

The structure of the H₂ spectrum is very similar. However, the double S transition on the ortho/para dimer shows a better separation due to the lower rotational moment of inertia compared to the D₂ spectrum. Therefore, they should be even more useful for ortho–para studies.

2. First vibrational branch of the H₂–D₂ mixture

Figure 1(c) shows the spectrum of the mixed sample. All lines of the pure H₂–D₂ spectra are also visible, so only the additional lines
that are predicted by the selection for heteromolecular dimers are indicated in the spectrum.

In the $D_2$ branch, two relatively strong new lines, indicated by solid lines, are visible. These $S_0(0)Q_{12}^D(0, 1)$ and $S_0(1)Q_{12}^D(0, 1)$ lines can be associated with a rotational excitation on the para or the ortho $H_2$ molecule and a vibrational excitation on the $D_2$ molecule. These lines are shifted to higher wavenumbers compared to the corresponding pure $D_2$ dimer transitions caused by the lower rotational moment of inertia of the $H_2$ molecule.

In the $H_2$ branch, similar lines can be observed, which originate from the same mechanism. However, in this branch, they are shifted to lower wavenumbers with respect to the pure $H_2$ dimers. In addition, for this mixed sample, four additional lines are visible in each branch. Similar to the pure spectra, these lines are caused by double-$S$ transitions of heteromolecular dimers.

C. Second vibrational branch

The overall structure of the second vibrational branch (Fig. 2) is quite similar to the first vibrational branch. Each branch of the pure samples begins with the pure vibrational monomer excitation $Q_2(0, 1)$, followed by combined ro-vibrational transitions (see Table III).

The phonon excitations are hardly visible, which is a consequence of our background reduction. In the case of the $D_2$ branch, the residue of the phonon excitation can be seen on the high energy flank of the pure vibrational excitation at $\sim 5900$ cm$^{-1}$. However, there are two main differences to the first vibrational branch. First, the lines are better separated than in the first vibrational branch. Second, the vibrational excitation of $\Delta \nu = \pm 2$ can be split to two molecules in a dimer. This directly leads to a pure vibrational dimer line $Q_1Q_2$. For the homomolecular dimer, this line is the combination of the three excitations $Q_1(0)Q_2(0)$, $Q_1(0)Q_2(1)$, and $Q_1(1)Q_2(1)$. However, the splitting of the $Q_2(0, 1)$ and $Q_2(0, 1)$ $Q_1(0, 1)$ lines directly reveals the an-harmonic vibrational potential. For the heteromolecular dimer, it is a combination of four excitations since $Q_1^{D_2}(0)Q_2^{H_2}(1)$ and $Q_1^{D_2}(1)Q_2^{H_2}(0)$ are distinguishable.

The dimer single $S$-transitions are split into four groups. Two with the rotational excitation on one molecule and the vibration on the other $S_0(0)Q_2(0, 1)$ and $S_0(1)Q_2(0, 1)$, and two where the vibrational excitation is split to both molecules $S_1(0)Q_2(0, 1)$ and $S_1(1)Q_2(0, 1)$. Similar to the pure vibrational excitations, the anharmonic vibrational potential shifts the position of the $S_0Q_2$ lines to higher wavenumbers compared to those of $S_0Q_2$. By this, the homomolecular $S_0(1)Q_2(0, 1)$ and $S_0(0)Q_2(1, 0)$ transitions overlap and cannot be separated. Therefore, only three lines are observed for these four groups.

1. Dimer double $S$ transitions in the second vibrational branch

Double-$S$ transitions are also suppressed in the second vibrational branch. In the region between 6400 cm$^{-1}$ and 6600 cm$^{-1}$ of the $D_2$ sample (see Fig. 3), double-$S$ transitions are predicted and can be identified in the spectrum around 6550 cm$^{-1}$. They occur with a maximum peak height of $0.5 \times 10^{-5}$ to $3.0 \times 10^{-5}$ and an integral absorbance of 0.05 cm$^{-1}$. This corresponds to suppression by one order of magnitude compared to the $Q_2$ monomer transition (integral absorbance of 0.43 cm$^{-1}$ see Table III) and by a factor of 200 compared to dimer transitions. A precise quantification of this suppression is again difficult due to the overlapping lines. The same is true for heteromolecular dimers like in Fig. 4.

FIG. 3. IR spectrum of liquid $D_2$ of double $S$-transitions in the second vibrational branch. Calculated line positions are indicated by the annotated dashed–dotted lines, for details see text.

FIG. 4. IR spectrum of the heteromolecular dimer double-$S$-transition in the second vibrational branch. The calculated line position is indicated by the annotated dashed–dotted line, for further details, see text.
2. Monomer S-transitions

According to the fundamental selection rules of $\Delta J = 0$, $+1$, monomer S-transitions should be forbidden for dipole transitions, but not for quadrupole transitions. In the first vibrational branch, it is impossible to resolve the $S_1$ monomer and the $S_0Q_1$ dimer transitions due to the spectral broadening in the liquid phase. Just to remind, in the gas phase this can be resolved like shown by McKellar et al.$^{2,35,36}$ For the second vibrational branch, the observed broadening is significantly smaller, and, therefore, it is possible to separate $S_2$ and $S_0Q_2$ transitions.

The $S_2(1)$ transition is expected between 8580 cm$^{-1}$ and 8600 cm$^{-1}$ (see Fig. 5). Our recorded spectrum shows no evidence for this transition. In addition, for the $S_2(0)$, expected between 8380 cm$^{-1}$ and 8400 cm$^{-1}$, no evidence is found (see Fig. 6). Therefore, these kinds of transitions seem to be limited to quadrupole transitions, and we can neglect them for our basic set of selection rules.

3. Second vibrational branch of the H$_2$–D$_2$ mixture

The transitions between the H$_2$ and D$_2$ second vibrational branches give a model-free proof for the heteromolecular dimers. The Raman based measurement of the sample isotopologue concentration, before and after the IR measurement, proves that this sample had an HD concentration of less than one percent (see Table I). Therefore, HD as an origin of the lines in this region can be excluded.

The Raman based measurement of the sample isotopologue concentration, before and after the IR measurement, proves that this sample had an HD concentration of less than one percent (see Table I). Therefore, HD as an origin of the lines in this region can be excluded. The most obvious lines in the mixture are between 7100 cm$^{-1}$ and 7800 cm$^{-1}$. These can be assigned to the heteromolecular $Q_1Q_1$ and $S_1Q_1$ transitions. The $S_1Q_1$ transition splits into four different components, depending on the location of the rotational excitation that can occur on a ortho-H$_2$ (at 7700 cm$^{-1}$), para-H$_2$ (at 7500 cm$^{-1}$), para-D$_2$ (at 7420 cm$^{-1}$), or ortho-D$_2$ (at 7300 cm$^{-1}$) molecule.

Within the second vibrational branch of H$_2$ and D$_2$, additional single and double-S transitions with a vibrational excitation of $\Delta v = +2$ are visible.

VI. DISCUSSION

The primary objective of this study is to obtain a minimal but complete set of descriptors for the IR absorption spectra of liquid hydrogen isotopologues. This is mandatory when aiming for an optimized calibration procedure to improve the accuracy from 5% to 1% and to extend our calibration for H$_2$, D$_2$, and HD mixtures to all six hydrogen isotopologues.

We find that the spectra of homomolecular mixtures are well-described using the following set of descriptors:

- monomer vibrational excitations $\Delta v = +1, +2$ with $Q_{J,2}(0, 1)$,
- phonon excitations, 50 cm$^{-1}$ to 100 cm$^{-1}$ higher than the monomer excitations,
- dimer excitations as combinations of $\Delta j_{1,2} = 0, +2$ and $\Delta v_{1,2} = 0, +1, +2$, where each molecule in the dimer must undergo at least one transition.

This set satisfactorily predicts all observed spectral features. However, regarding the integral absorbance, three important effects need to be taken into account.

First, the second vibrational branches are suppressed by one to two orders of magnitude compared to the first vibrational branches. These transitions are only allowed due to the anharmonicity of the vibrational potential and are, therefore, suppressed. For the other transitions, a closer investigation is needed since there is no simple reason that the $Q_1Q_1$ lines compared to $Q_1$ are suppressed by the same order of magnitude as $Q_2$ to $Q_1$.
Second, all dimer double-S-transitions are strongly suppressed compared to their corresponding single-S-transitions, but not forbidden. These transitions were identified in the spectra of the first and second vibrational branch and have to be taken into account for a calibration accuracy of about 1% or better. The observed suppression could be explained by the fact that, in a double-S-transition, the rotational excitation and the configuration of all three spins \((J_A, J_B,\text{ and the dimer rotational quantum number } l)\) need to fulfill the photon selection rule of \(\Delta J = -1, 0, +1\) in combination. However, precise information about all bound states of the dimers is needed for a full calculation of the phase space of these transitions, which could be gathered by calculations similar to those by Hinde.\(^{[17]}\)

Third, the missing monomer S-transitions are in agreement with the suppression of quadrupole transitions and, therefore, do not need to be considered for a calibration with an accuracy of 1%. However, from the literature, it is known that these lines can be found in high resolution spectra.\(^{[18]}\) In order to achieve high precision calibration with an accuracy much better than 1%, these lines need to be investigated in detail as it can be seen for gas phase spectra (see Ref.\(^{[19]}\) for HD, Ref.\(^{[20]}\) for pure rotational excitation, and Ref.\(^{[12]}\) for dimers). It may, therefore, be concluded that the missing dimer transition lines, for which only one molecule exhibits vibration and the dimer itself is rotationally excited \((S_A, Q_B)\), seem to be prohibited or suppressed.

Additionally, various effects complicate the calibration of an IR absorption based measurement system for the isotopologue concentration.

First, the samples are mixtures of all six isotopologues, which complicates the spectra due to additional heteromolecular-dimer lines and overlaps caused by the broad absorption lines in the liquid phase.

Second, the absorbance depends not only on the six isotopologue concentrations, but also on the three ortho/para ratios.\(^{[19]}\)

Third, the underlying physics of this set of selection rules causes mutual influence between the isotopologues and the spin isomer states ortho and para.

This results in a dependence even for the absorbance of the lines of one isotopologue from the concentrations and ortho/para ratios of the other isotopologues. This is due to the fact that the transition matrix elements are induced by molecular interactions with the neighboring molecules;\(^{[18,40–42]}\) therefore, the transitional dipole moment can be described as first order approximation as the product of the polarizability of the absorbing molecule and the field strength of the colliding molecule.\(^{[41]}\) The polarizability \(\rho_J\) of the molecules weakly depends \((<1%)\) on the rotational state,\(^{[41]}\) but the multipole moments \(F_J\) of the \(J = 0\) and \(J = 1\) H\(_2\) molecules differ by two orders of magnitude. Thus, the transition matrix element needs to be a function of both concentrations and surface field strength\(^{[43]}\) following \(M_{Q_1(0)}(p_{00},c_{00} \cdot F_{Q_2},c_{11} \cdot F_{11})\). This should be also true for mixtures of H\(_2\) and D\(_2\) and directly leads to the dependence of the integral absorbance of the D\(_2\) Q\(_1\) and Q\(_2\) transition on the H\(_2\) ortho–para ratio and vice versa.

This mechanism can be further investigated by a careful investigation of the dependence of the line absorbance on the ortho–para ratios. The information gathered by highly accurate calibration of the integral absorbance against the isotopologue and ortho–para composition can be combined with the information gathered from line shapes. As shown by Mengel et al.,\(^{[17]}\) the differences in line shapes between the liquid and solid phase allow deep insights into the underlying physics.\(^{[45,46]}\)

The mutual influence between the different isotopologues can also be used to acquire macroscopic information of the sample, such as the homogeneity of the ortho/para and isotopologue distribution in the sample. The ortho/para homogeneity, for example, can be derived from the double S transitions in the first vibrational branch (see Fig. 1, between 3300 cm\(^{-1}\) and 3600 cm\(^{-1}\)). For ortho–para lines, the absorbance is at its maximum in a homogeneous sample, while for ortho–ortho and para–para lines, the absorbance is at its minimum. With decreasing homogeneity (corresponding to a physical separation of the ortho and para molecules in the sample), the possibility to find ortho–para dimers will decrease and the absorbance of the ortho–ortho and para–para lines will increase. As a consequence, one may be able to determine the homogeneity from the relative absorbance of these lines. The same method can be used to determine the homogeneity of the isotopologue composition, where pure H\(_2\) or pure D\(_2\) transitions can be compared to the transitions of H\(_2\)–D\(_2\) dimers. However, only a careful investigation and calibration at the 1% level can give more detailed information about the liquid state and the underlying physics. An application of this can be of interest when studying phase transitions from liquid to solid where the solubility of one of the molecules can be different in the one or the other phase.

VII. CONCLUSION

The prediction of detailed line absorbance needs the calculation of transitional matrix elements, which are not available at the present time. However, there are some calculations for solid para-hydrogen absorption coefficients\(^{[47]}\) and a six-dimensional potential surface for ground state H\(_2\) and D\(_2\) dimers. A full set of absorption coefficients for the liquid phase would be appreciated and combined with the data here, enabling detailed studies of the interaction processes in the liquid phase of hydrogen isotopologues.

However, with the basic set of selection rules of \(\Delta v \geq 0\) and \(\Delta J = 0, \pm 2\), we are able to sufficiently describe the first and second vibrational branches of liquid hydrogen IR absorption spectra of pure H\(_2\), pure D\(_2\), and H\(_2\)–D\(_2\) mixtures.

We are confident that this set also describes mixtures with the other four isotopologues, including the tritiated species.\(^{[8,38–36]}\) This will be verified with the new tritium compatible setup TAPIR 2 currently under construction.\(^{[3]}\) The selection rules will also allow us to develop an optimized strategy for a calibration with an improved accuracy of 1% or better.

The main systematic effect in this procedure is expected to be the ortho/para ratios of the three homonuclear isotopologues. Therefore, one should think of the ortho and para molecules as fully distinguishable species in the liquid phase. In conclusion, the parameter space for a full calibration will contain nine parameters, one for each of the heteronuclear molecules and two for each of the homonuclear molecules.

Further studies of the liquid should address in more detail the dependence of the absorbance on the ortho–para and isotopologue composition, instead of the line positions, which are not an ideal parameter in the liquid phase. Future work should also try to deconvolute the lines according to the identified underlying transitions that contribute to the observed line features. Specifically, the change
during the ortho–para conversion promises a clear and testable deconvolution of several lines, such as the double-S-transitions in the first vibrational branch, the collision induced Q2 monomer lines, and the pure vibrational and single S dimer lines in the second vibrational branch. Combined with an accurate calibration, this will enable investigations of the liquid phase itself to gain a deeper understanding of liquid hydrogen.

**DATA AVAILABILITY**

Data files of the spectra shown in this publication are available from the corresponding author upon reasonable request.

**SUPPLEMENTARY MATERIAL**

See the supplementary material for the complete list of calculated transitions in the investigated region.

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