Optical cycling in polyatomic molecules with complex hyperfine structure

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We have developed and demonstrated a scheme to achieve rotationally closed photon cycling in polyatomic molecules with complex hyperfine structure and sensitivity to hadronic symmetry violation, specifically, ¹⁷¹YbOH and ¹⁷³YbOH. We calculate branching ratios for spontaneous decay into the different hyperfine levels and identify repumping schemes. We demonstrate our scheme by cycling photons in a molecular beam and verify that we have achieved rotationally closed cycling by measuring optical pumping into unaddressed vibrational states. Our work makes progress along the path toward utilizing photon cycling for state preparation, readout, and laser cooling in precision measurements of polyatomic molecules with complex hyperfine structure.

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I. INTRODUCTION

Recent advances in the cooling and trapping of increasingly complex molecules [1,2] are enabling a multitude of applications that leverage molecular complexity for applications in quantum science and precision measurement [3–7]. One avenue for increased complexity is the use of molecules with a heavy nucleus and nonzero nuclear spin, *I*. Precision measurements with these molecules can probe new physics related to the nucleus, such as charge-parity (CP)-violating nuclear moments [6], nuclear-spin-dependent parity violation (NSD-PV) [8,9], and measurements of nuclear structure [10,11].

One example is the odd isotopologues of the linear polyatomic molecule YbOH, which are promising candidates for probing symmetry-violating physics in the hadronic sector: ¹⁷¹YbOH for parity violation [12] and ¹⁷³YbOH for the CP-violating nuclear magnetic quadrupole moment (NMQM) [13,14]. Interactions of core-penetrating valence electrons with the heavy Yb nucleus enhance sensitivity to symmetry violation [15,16], and the quadrupole shape deformation of the Yb nucleus provides further collective enhancement of CP-violating moments [13]. Additionally, the vibrational bending mode of YbOH exhibits closely spaced, opposite parity levels. These parity doublets allow for control of molecular polarization in modest electric fields [17] and suppression of systematic errors in precision measurements [14,18,19], and can be tuned even closer to degeneracy by modest magnetic fields, enhancing sensitivity to NSD-PV [12]. Bending modes are one example of parity doublet structures that are generic to polyatomic molecules and which exist independently of the electronic structure, thereby enabling their combination with efficient optical cycling and laser cooling [7,14,20–22].

The additional complexity of polyatomic molecules presents both opportunities and challenges. In particular, the

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increased number of vibrationally excited states in the electronic ground state provide many pathways for spontaneous decay after optical excitation. Molecules that decay to "dark states" not addressed by repumping lasers are lost from the optical cycle [1,23]. Addressing rotational and vibrational branching to dark states has enabled laser cooling and trapping of both diatomic and polyatomic molecules [1,24]. However, eliminating dark states in species with large and complicated hyperfine structure, which is a byproduct of many molecules with spin on a heavy nucleus, poses additional challenges [25]. For example, in the same rotational manifold, ¹⁷⁴YbOH has 12 Zeeman sublevels and ¹⁷³YbOH has 6 sublevels. In particular, hyperfine splittings will have a multiplicative effect on the number of existing rotational and vibrational dark states, with frequency splittings large enough such that bridging with acousto-optical or electro-optical modulators becomes a challenge.

In this manuscript, we report the design and experimental realization of a scheme for achieving rotationally closed cycling of ¹⁷¹YbOH ($I_{Yb} = 1/2$) and ¹⁷³YbOH ($I_{Yb} = 5/2$). With only two modulation frequencies on a single laser, we achieve photon cycling with closure of rotational, spin-rotational, and hyperfine structure. Furthermore, such a scheme should be broadly applicable to other molecules with similar structure, especially with the implementation of additional techniques, such as computer-generated holography [26].

II. BRANCHING RATIO CALCULATION

In this manuscript, we address the branching of spontaneous decays $\tilde{A}^2 \Pi_{1/2}(0, 0, 0) \rightsquigarrow \tilde{X}^2 \Sigma^+(0, 0, 0)$. Here, $\tilde{A}^2 \Pi_{1/2}(0, 0, 0)$ is the vibrational ground state of the excited electronic state $\tilde{A}^2 \Pi_{1/2}$, and $\tilde{X}^2 \Sigma^+(0, 0, 0)$ is the vibrational ground state of the ground electronic state $\tilde{X}^2 \Sigma$. We will often refer to the states as simply "A" and "X", respectively, and if the vibrational quantum numbers ($\nu_{\text{Yb-O stretch}}$, ν_{bend} , $\nu_{\text{O-H stretch}}$) are omitted, it is assumed that

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we mean the ground vibrational state (0,0,0). The vibrational branching ratio of $A \rightsquigarrow X$ decay has been previously measured [22] to be 89.44(61)%. Here, we are primarily concerned with the rotational, spin-rotational, and hyperfine branching, which are significantly more complicated in lasercoolable species with nonzero nuclear spins on the metal and the ligand [25].

The branching ratios are calculated using effective Hamiltonians [27]. The energies and eigenvectors within the X and A states are obtained by diagonalizing Hamiltonian matrices in the Hund's case (a) basis, and then transition dipole moments (TDMs) are calculated between the eigenvectors of the two states. For simplicity, the hydrogen nuclear spin from -OH was not included in the basis set, as the hyperfine splitting is not optically resolved [28]. However, we must remember that it is present when calculating level degeneracies.

The effective Hamiltonians used for the two states are [29]

$$H_X^{\text{eff}} = BN^2 + \gamma N \cdot S + b_F I \cdot S + c \left(I_z S_z - \frac{1}{3} I \cdot S \right) + e^2 Q q_0 \frac{3I_z^2 - I^2}{4I(2I - 1)}, \qquad (1)$$

$$H_A^{\text{eff}} = AL_z S_z + BN^2 + \frac{1}{2}(p + 2q)(J_- S_- e^{+2i\phi} + J_+ S_+ e^{-2i\phi}) + h_{1/2} I_z L_z - \frac{1}{2}d(S_+ I_+ e^{-2i\phi} + S_- I_- e^{+2i\phi}) - 3I^2 - I^2$$

$$+e^{2}Qq_{0}\frac{3I_{z}^{2}-I^{2}}{4I(2I-1)}.$$
(2)
re, S is the Yb-centered electron spin, I is the Yb nu-

Here, *S* is the Yb-centered electron spin, *I* is the Yb nuclear spin, and *N* is the total nonspin angular momentum. All angular momentum subscripts (z, \pm) denote molecule frame components. Spin-orbit *A*, rotation *B*, spin-rotation γ , and Λ -doubling (p + 2q) are present in all isotopologues. Both ¹⁷¹Yb(I = 1/2) and ¹⁷³Yb(I = 5/2) have nuclear spins, which give rise to additional hyperfine parameters, namely, orbital hyperfine *a*, Fermi contact b_F , spin-dipolar *c*, and parity-dependent dipolar *d*. In the excited state, the diagonal hyperfine shifts are determined by an effective parameter $h_{1/2}$, which can be written as $h_{1/2} = a - (\frac{b_F}{2} + \frac{c}{3})$ [29]. Further, ¹⁷³Yb also has an electric quadrupole moment, which gives rise to the term e^2Qq_0 . The exact parameter values used here are taken from Ref. [29] and listed in Table I.

After diagonalizing the Hamiltonians, the eigenvectors are labeled according to the Hund's cases that best represent their structure: Hund's case $(b_{\beta S})$ for X and Hund's case $(a_{\beta J})$ for A [29]. The labels are simplified to include only the quantum numbers relevant to this manuscript: $|NGF\rangle$ for the X state and $|J(P)F\rangle$ for the A state. Here, for the X state, G = I + S results from the hyperfine interaction between I and S, while for the A state, J results from both molecule rotation and spin-orbit coupling, and (P) is the parity label for the parity doublets resulting from Λ -doubling. The total angular momentum F is given by F = G + N in the X state and F = J + I in the A state. We will use double primes to denote ground-state quantum numbers (e.g., N'') and single primes to denote excited-state quantum numbers (e.g., J').

TA	ABLE I.	Relevant	parameters,	from	[29],	in	wave	numbers
(cm^{-})	¹) for the	$\tilde{X}^2 \Sigma^+(0,$	$(0, 0)$ and \tilde{A}^2	$\Pi_{1/2}(0$, 0, 0)	sta	ates of	¹⁷¹ YbOH
and 17	⁷³ YbOH.							

State	Parameter	¹⁷¹ YbOH	¹⁷³ YbOH
$\overline{\tilde{X}^2\Sigma^+(0,0,0)}$	В	0.245	0.245
	γ	-0.00270	-0.00270
	\dot{b}_F	0.228	-0.0628
	с	0.0078	-0.00273
	$e^2 Q q_0$	N/A	-0.111
$\tilde{A}^2 \Pi_{1/2}(0,0,0)$	Α	1350	1350
,	В	0.253	0.253
	p+2q	-0.439	-0.438
	$h_{1/2}$	0.0148	-0.00422
	d	0.0320	-0.00873
	$e^2 Q q_0$	N/A	-0.0642

TDMs are calculated between the eigenvectors by representing all states in the Hund's case (a) basis and performing computations only in this basis. Case (b) quantum numbers labelled N'', G'' are assigned to the ground-state eigenvectors by identifying patterns in the eigenvalues. To obtain branching ratios and account for the degeneracy of M_F sublevels, the TDMs are summed over light polarization p and ground-state M''_F levels, and averaged over excited-state M'_F levels. We obtain

$$\frac{1}{2F'+1} \sum_{p,M''_F,M'_F} \langle N''G''F'' | T_p^1(d) | J'(P)F' \rangle.$$
(3)

In this way, the branching ratio derived by squaring the TDM is naturally normalized, which means that the branching ratios originating from the same excited state add up to 1. Note that we use the same reduced matrix element convention as Ref. [27].

The calculated branching ratios are shown in Fig. 1. Note that these branching ratio numbers add up to 1 only within the same vibration level, and they will be referred to as rotational branching ratios. The total branching ratio of a transition can be derived by multiplying the rotational branching ratio by the vibrational branching ratio, which is known for ¹⁷⁴YbOH [22] and expected to be the same for ^{171,173}YbOH.

III. METHOD FOR ACHIEVING ROTATIONALLY CLOSED CYCLING

We achieve rotationally closed optical cycling via $N'' = 1 \rightarrow N' = 0$ type transitions [30], specifically, $N'' = 1 \rightarrow J' = 1/2(+)$ for YbOH (since N' is not well defined in A, J' = 1/2 corresponds to the lowest rotational level in A). Based on our calculations, rotationally closed cycling is straightforward to achieve for ¹⁷¹YbOH because there is a single transition, $F'' = 1 \rightarrow F' = 0$, with a calculated rotational branching ratio of >0.999. The excited hyperfine splitting of ~400 MHz is sufficiently large, compared to the observed linewidths of ~50 MHz, such that off-resonant excitation to F' = 1 can be avoided, which is not the case in ¹⁷³YbOH. The experimental linewidth is larger than the radiative width of ~10 MHz, mostly due to power broadening, and is beneficial for addressing all the sidebands for ¹⁷³YbOH.



FIG. 1. Calculated rotational branching ratios for (a) 171 YbOH and (b) 173 YbOH. Numbers 1–6 in circles label the transitions targeted for achieving rotationally closed cycling.

In order to fully address rotational branching for ¹⁷³YbOH, we must address all six ground hyperfine levels, as the excited F' = 2 and F' = 3 states are separated by ~150 MHz, which can result in off-resonant excitation when slightly power broadened. We address the ground levels by generating sidebands on a single laser using two fiber electro-optical modulators (EOMs) used in series.¹ Fortunately, the transitions are spaced such that it is possible to use only two EOMs, each with a single sinusoidal drive, to address all of transitions within the slightly power-broadened linewidth, as shown in Fig. 2.

¹EOSPACE PM-0S5-10-PFA-PFA-1154-UL-SOP125mW.

The two EOMs are used sequentially with input from an 1154 nm seed laser, and the output is then amplified and doubled to produce the needed visible light at 577 nm. Extra, unwanted sidebands are produced from both the EOMs and the doubling crystal, and the latter also modifies the sidebands due to the nonlinear nature of second harmonic generation. These sidebands can accidentally drive undesirable transitions that cause leakage by optically pumping N'' = 1 population into rotational dark states within the same vibrational manifold.

Here, we were able to avoid leakage-inducing transitions by using modulation frequencies set such that the three intense, unused sidebands, as well as smaller, higher-order sidebands, are sufficiently far away from unwanted transitions, as shown in Fig. 2. Specifically, one EOM is driven at 5.63 GHz and the other 495 MHz, with modulation depth of around $0.5V_{\pi}$, chosen such that the end output matches the patterns shown in Fig. 2. Note, however, that the complex and congested spectrum [29] of these molecules means that off-resonant excitation of unwanted lines cannot be ignored, and our method takes them into account when calculating the number of scattered photons. These problems can be alleviated using other techniques, such as computer-generated holography [26].

IV. MEASURING THE NUMBER OF SCATTERED PHOTONS

We experimentally study optical cycling using a cryogenic buffer gas beam (CBGB) [31] of YbOH. We create cold, slow beams of YbOH using methods similar to those described elsewhere [17,29,32]. We use a ~1.5 K cell with 2 SCCM flow of helium buffer gas, and enhance the production of YbOH using optically driven chemical reactions [32]. We use cw lasers to scatter photons, and the resulting laser-induced fluorescence is monitored with photomultiplier tubes (PMTs).

To verify that we have achieved rotational closure, we optically cycle and measure the number of photons scattered per molecule. If we have addressed all branching within X, the molecules will cycle until they are optically pumped into a dark vibrational state. The majority of molecules are pumped into one of the hyperfine levels of the $\tilde{X}^2\Sigma^+(1, 0, 0)$ state with one quantum of Yb-O stretching motion (v = 1), which



FIG. 2. Scheme for EOMs to address ¹⁷³YbOH branching. Blue (thin) lines are the target transitions that we want to address. From left to right, they correspond to transitions in Fig. 1(b) labeled 1–6. Yellow (thick) lines are generated by the pump laser and EOMs, which reflects the frequency spacings and relative amplitudes of the sideband generated EOMs as verified by using a Fabry-Perot cavity. As shown, the scheme used here can address all the target transitions within their linewidths, while avoiding the unwanted lines that can cause leakage to dark states.



FIG. 3. Schematic for measuring the number of photons cycled by measuring the population transfer into one of the hyperfine levels of the first excited vibration state (1,0,0) or alternatively labeled v = 1. We derive the number of cycled photons by comparing the probe fluorescence with the cycling pump and noncycling pump.

has a branching ratio of 9.11(55)% from A. After cycling upstream in the molecular beam, we measure the population downstream in one of the hyperfine levels of the v'' = 1 state (see Fig. 3).

The main benefit of such a population transfer measurement, compared to a direct fluorescence measurement of optical cycling, is the ability to reject accidental fluorescence from nearby transitions. There are five isotopes of Yb with significant abundance, and both ^{171,173}YbOH have extra complexity arising from their nuclear spins. As a result, the band head region, where the cycling transitions are located, is extremely congested with additional lines, and a direct fluorescence measurement would make it difficult to disentangle the effects of photon cycling versus merely addressing multiple lines in different states and isotopologues.

By using a pump-probe scheme to measure the population transfer, most of these unwanted transitions do not contribute to our results. The exception is when both pump and probe coincide with optical pumping of a different isotopologue or level that is unrelated to cycling. By using both resonant chemical enhancement [32] of a specific Yb isotope and intentional selection of the hyperfine level in v'' = 1 for the probe transition, contaminant contributions to optical pumping are relatively small and can be estimated using detunings and branching ratios.

Our figure of merit is the quantity *D*, defined as the difference in v'' = 1 probe fluorescence caused by the cycling pump laser, normalized against probe fluorescence without the pump. Because the $v'' = 1 \rightarrow v' = 0$ probe transition has a very small branching ratio of 0.0911, we operate with low saturation parameter, and the measurement of fluorescence increase is equivalent to an increase in population. To extract the number of photons cycled from *D*, we use the following relationship:

$$D = \frac{\text{fluorescence}_{\text{pump}} - \text{fluorescence}_{\text{no pump}}}{\text{fluorescence}_{\text{no pump}}}$$
$$\approx \frac{N_{f,1} - N_{i,1}}{N_{i,1}} = \frac{N_P}{N_{i,1}} \approx \frac{N_{i,0}PB_1}{N_{i,1}}, \tag{4}$$

where N_P is the population transferred into v'' = 1 from optical pumping, $N_{i,1}$ is the initial population of the v'' = 1 state being probed, $N_{i,0}$ is the initial population of the v'' = 0 states being pumped, $P \approx 1/(1 - B_0)$ is the number of photons cycled per

pumped molecule, B_0 is the sum of total branching ratios of all the pumped states in v'' = 0, and B_1 is the branching ratio down to the specific hyperfine state probed in the v'' = 1 probe state (see Fig. 3).

The ratio $N_{i,0}/N_{i,1}$ can be calculated from measurements by making two assumptions about the initial populations of the relevant states: First, that the states within a vibrational manifold are well thermalized coming out of the cryogenic buffer gas beam source, and second, that the ratios of population between the vibration ground and first excited vibration mode, $R = N_{\text{total}(v=1)}/N_{\text{total}(v=0)}$, are the same for different isotopologues of YbOH under the same source condition. These assumptions are supported by tests done with ¹⁷⁴YbOH and from behavior in other such molecular sources [31].

V. CALIBRATION WITH ¹⁷⁴YbOH

We first made measurements with ¹⁷⁴YbOH to validate our method since photon cycling in ¹⁷⁴YbOH has been carefully characterized elsewhere [22], as well as to derive the population ratio *R* between the vibrational modes in our molecular beam. Results are shown in Fig. 4, which gives two final *D* values for two different pumping laser configurations: D_{ro} for rotationally open (ro) pumping, and D_{rc} for rotationally closed (rc) pumping. For ¹⁷⁴YbOH, we only need one sideband to cover the spin-rotation splitting and achieve rotationally closed cycling [21], which gives a scattered photon number of $P \approx 1/(1 - B_0) = 9.1$ per addressed molecule, limited only by the vibrational branching ratio $B_0 = 0.89$. In comparison, when only addressing the J = 1/2 state of the spin-rotation pair, which has a rotational branching of 0.67, the expected photon number is only P = 2.5, where $B_0 = 0.89 \times 0.67 = 0.60$.

With the assumption of thermalization, we can treat all rotational levels and M_F sublevels within a given vibronic manifold as having equal initial population since the level splittings are all far smaller than the thermal energy $k_BT \approx$ 30 GHz, where k_B is Boltzmann's constant and our operating temperature is $T \approx 1.5$ K. Let the initial population of the pumped state be the degeneracy of the state times 1 (including hydrogen nuclear spin). For ¹⁷⁴YbOH, the initial population is $N_{i,0} = 12$ for addressing both components of the spin-rotation (SR) pair, and $N_{i,0} = 4$ for addressing only the J'' = 1/2 state of the pair. Then, using $R = N_{\text{total}(v=1)}/N_{\text{total}(v=0)}$ as an unknown variable, we have the initial population of the probed state as $N_{i,1} = 4R$. Here the probe laser addresses the v'' = 1, N'' = 1/2 state, which has a total branching ratio of $B_1 = 0.06$ coming from J' = 1/2(+) excited state.

As a result, when the pump addresses both components of the SR pair, we would expect the measurement to be $D_{rc} = N_{i,0}PB_1/N_{i,1} = 1.64/R$. Similarly, when the pump addresses only the J'' = 1/2 state, we expect $D_{ro} = 0.15/R$. Experimentally, we measured *D* to be $D_{rc} = 15.0 \pm 1.6$ and $D_{ro} = 1.38 \pm 0.23$, which gives *R* values of 0.11 ± 0.01 and 0.11 ± 0.02 . In other words, the v'' = 1 manifold has approximately 11% of the population of the v'' = 0 manifold. The results are averaged from 40 data sets, where the statistical uncertainties mainly come from shot-to-shot variations of the molecule source. The two different pumping configurations give the same *R* value within our error, validating the



FIG. 4. ¹⁷⁴YbOH fluorescence signals from a probe laser addressing the v'' = 1, N'' = 1, J'' = 1/2 state. Error bars represent 1- σ variation due to shot-to-shot fluctuations over 80 beam pulses. (a) The pump laser has no sidebands and only addresses the J'' = 1/2 state in the ground spin-rotation pair. The increase in fluorescence is a factor of $D_{ro} = 1.38 \pm 0.23$, as described in the main text. (b) The pump laser addresses both J'' = 1/2 and J'' = 3/2, achieving rotationally closed cycling and increasing the fluorescence by $D_{rc} = 15.0 \pm 1.6$.

measurement method. For the measurements on 171,173 YbOH, we will similarly perform both rotationally closed and open pumping, but will use the R = 0.11 as the known parameter to derive the number of scattered photons per molecule.

VI. SPECTROSCOPY OF ^{171,173}YbOH $\tilde{X}^2\Sigma^+(1,0,0)$

In order to measure the population transfer from optical pumping, it is important to establish the correct quantum numbers and energies for the $\tilde{X}^2 \Sigma^+(1, 0, 0)$, or v'' = 1, states within the N'' = 1 manifold for ^{171,173}YbOH. Given the known odd and even isotopologue parameters for the v'' = 0 state [28,29], and the even isotopologue parameters for the v'' = 1 state [33], we first predict the expected spectral region for the $\tilde{X}^2 \Sigma^+(1, 0, 0) \rightarrow \tilde{A}^2 \Pi_{1/2}(0, 0, 0)$ transition. We then measure spectra around the predicted region and look for splittings that match the known excited splittings between hyperfine levels of $\tilde{A}^2 \Pi_{1/2}(0, 0, 0)$, J = 1/2(+), and J = 3/2(+). For



FIG. 5. Example of v'' = 1 spectroscopy for ¹⁷¹YbOH. Transitions are identified by matching spectral splittings to known excited-state splittings [29]. (a) The level diagram of the involved ground and excited states, with calculated branching ratios for the target transitions. (b) The measured fluorescence spectrum with the same four transitions marked in (a). Some of the unmarked peaks that are nearby are transitions from F'' = 0, 2 ground hyperfine levels to the same excited states, identifiable via the common splittings.

transitions with a shared ground state and different excited state, the observed line splittings will match the known structure of the excited $\tilde{A}^2 \Pi_{1/2}(0, 0, 0)$ state [29].

The spectral region is congested with overlapping transitions from multiple rotational and hyperfine states, as well as from other isotopologues. Furthermore, the v = 1 states have an order-of-magnitude-smaller population coming out of the cryogenic buffer gas source compared to v = 0, as we learned from the ¹⁷⁴YbOH measurements in the previous section. In order to differentiate the multiple transitions, we used chemical enhancement [32] and population pumping from v = 0.

As an example, Fig. 5 shows four transitions from the X, v'' = 1, N'' = 1, G'' = 1, F'' = 1 ground state to four hyperfine levels in the excited A, v' = 0, J' = 1/2, and J' = 3/2 states. The spectrum in Fig. 5(b) is taken downstream of a power-broadened pumping laser with EOM-generated sidebands that roughly addresses all the v'' = 0, N'' = 1 ground states connected to the J' = 1/2(+), F' = 1 excited state. The EOM sideband scheme is not optimized here because we need the initial spectroscopy to confirm our probe state selection. After the prerequisite spectroscopy, we are able to optimize the sideband scheme to achieve rotationally closed photon cycling.

A full spectroscopic analysis and parameter fit for all the v = 1 states is beyond the scope of this work. However, we were able to identify all of the relevant hyperfine levels within



FIG. 6. ¹⁷¹YbOH fluorescence signals from the probe laser addressing v'' = 1 |N'' = 1, G'' = 1, F'' = 1). Error bars represent 1σ spreads due to shot-to-shot fluctuations over 80 beam pulses. (a) Pump laser addressing the $F'' = 1 \rightarrow F' = 1$ rotationally open transition. The increase in fluorescence is $D_{ro} = 0.48 \pm 0.17$. (b) Pump laser addressing the $F'' = 1 \rightarrow F' = 0$ rotationally closed cycling transition. The fluorescence increase is $D_{rc} = 7.4 \pm 1.3$.

the v'' = 1, N'' = 1 manifold for ^{171,173}YbOH (see Appendix). We are confident of our state identification based on the matching splittings in both ground and excited states, and based on the observation of expected population transfers.

VII. PHOTON CYCLING IN 171,173 YbOH

After identifying all of the v'' = 1, N'' = 1states for both 171,173 YbOH, we decided to use the $\tilde{X} \,{}^2\Sigma^+(1,0,0) | N'' = 1, G'' = 1, F'' = 1 \rangle \rightarrow$ $\tilde{A}^2\Pi_{1/2}(0,0,0) | J' = 1/2(+), F' = 0 \rangle$ transition for the 171 YbOH probe since it is the rotationally closed analog of the corresponding $(0,0,0) \rightarrow (0,0,0)$ transition, and relatively well separated from other hyperfine levels. For 173 YbOH, we decided to use the $\tilde{X} \,{}^2\Sigma^+(1,0,0) | 1,3,2 \rangle \rightarrow$ $\tilde{A}^2\Pi_{1/2}(0,0,0) | 1/2(+),2 \rangle$ transition, as it minimized accidental pump or probe signal contaminants from unwanted states and since it has very similar branching from both excited F' hyperfine levels, simplifying data analysis and modeling. With the probe transitions identified, we were



FIG. 7. ¹⁷³YbOH fluorescence signals from probe laser addressing $v'' = 1 | N'' = 1, G'' = 3, F'' = 2 \rangle$. Error bars represent 1 σ spreads due to shot-to-shot fluctuations over 80 beam pulses. (a) Pump laser with no sidebands, addressing only the $|1, 2, 3\rangle \rightarrow F' = 3$ transition. The change in integrated fluorescence is $D_{ro} = 0.40 \pm 0.14$. (b) The pump laser addresses all the v'' = 0, N'' = 1 hyperfine levels using the optimized EOM scheme, and thus achieves rotationally closed cycling. The increase in fluorescence is $D_{rc} = 13.0 \pm 1.9$.

able to optimize the cycling schemes and lower the pump laser power required to less than 25 mW per sideband in a 5-mm-diameter beam.

Following the same procedure used for ¹⁷⁴YbOH laid out in the previous section, we measured the scattered photon number per molecule for ¹⁷¹YbOH and ¹⁷³YbOH to confirm that our cycling schemes work as expected. Figure 6 shows the results for ¹⁷¹YbOH. The rotationally closed cycling transition is from F'' = 1 to F' = 0, which gives an initial population, including unresolved –OH hyperfine degeneracy, of $N_{i,0} = 6$. The pump transition for the rotationally open comparison is from the same ground state, and hence the same initial population, but different excited state, F' = 1, which gives a calculated branching ratio of $B_0 = 0.22$. The probe transition addresses the equivalent ground state in v'' = 1, so $N_{i,1} = 6R$, and the total branching ratio from A to v'' = 1 is $B_1 = 0.09$ from F' = 0 and $B_1 = 0.02$ from F' = 1.

Figure 6(a) shows results from pumping on the rotationally open transition, which gives an increase in fluorescence

TABLE II. A comparison of expected and measured numbers of photons scattered per molecule, denoted as *P*. Here, RO refers to rotationally open pumping and RC refers to rotationally closed pumping. Measurements of *P* in ¹⁷⁴YbOH are used, along with the theoretical prediction, to derive the vibration population ratio *R*. Both RO and RC for ¹⁷⁴YbOH yielded the same resulting *R*. The high *P* measured in RO ¹⁷¹YbOH is caused by the nearby ground hyperfine levels' contribution.

Molecule	RO or RC	Expected P	Measured P
¹⁷⁴ YbOH	RO	2.5	
10011	RC	9.1	
171 VHOH	RO	1.3	2.4 ± 0.8
10011	RC	9.1	8.9 ± 1.5
173 VhOH	RO	1.5	1.4 ± 0.5
1000	RC	9.1	9.1 ± 1.3

of $D_{ro} = 0.48 \pm 0.17$. Using the value $R = 0.11 \pm 0.01$ from 174 YbOH, we obtain a photon number of 2.4 ± 0.8 , which is larger than the 1.3 expected from a branching ratio of 0.22. The reason for the discrepancy is the contribution from nearby v'' = 1 ground-state hyperfine levels, which are 80 and 150 MHz away. The nearby levels contribute both to the initial population $N_{i,0}$ and to the branching ratio B_0 . By calculating and including the effect of photon scattering from these nearby states, we actually expect a fluorescence increase of $D_{ro} = 0.45$, which fits the data well and thus confirms that the assumption holds that the vibrational population ratio Ris isotope independent. For the rotationally closed transition, we measured a fluorescence increase $D_{rc} = 7.4 \pm 1.3$, which gives a scattered photon number of 8.9 ± 1.5 , which matches the expectation of 9.1 from the vibrational branching ratio of 0.89.

Figure 7 shows results for ¹⁷³YbOH. The reference pump transition addressing F'' = 3 has $N_{i,0} = 14$, and a branching ratio of $B_1 = 0.02$ to the probe state in v'' = 1. The cycling pump addresses all six ground hyperfine levels, giving $N_{i,0} = 72$, and the same branching ratio to the probe state. The probe laser addresses F'' = 2, giving $N_{i,1} = 10R$. With the same R = 0.11value as before, the measured normalized difference in fluorescence, $D_{ro} = 0.40 \pm 0.14$, means that the reference pump scattered $P = 1.4 \pm 0.5$ photons per molecule, matching the expectation of 1.5 for the branching ratio of $B_0 = 0.33$ on the open transition. For rotationally closed cycling, the result of $D_{rc} = 13.0 \pm 1.9$ would indicate a photon number of 9.1 ± 1.3 , matching the expected 9.1 from purely vibrational branching. Note that there is an expected rotational leakage of 0.5% due to hyperfine-induced transitions [29], which is below our resolution to observe; however, this could be addressed by adding additional sidebands to address the relevant N'' = 3 levels.

Table II summarizes the result of our photon cycling measurements. Our measured results for the number of photons scattered per molecule, P, are in good agreement with the expectation from our calculated branching ratios. In particular, once rotational branching is addressed, the value of Pis limited by vibrational branching, which is independent of isotopologue. This is also reflected in our data, with each isotopologue having the same value of P for rotationally closed pumping.

VIII. CONCLUSION

We have demonstrated a scheme that achieves rotationally closed photon cycling of ¹⁷¹YbOH and ¹⁷³YbOH, both of which have a complicated hyperfine structure. The method is technically straightforward and can be easily extended to higher-order vibrational repumps to enable larger numbers of scattered photons. Since molecule-based studies of the nuclear weak force and the search for CP-violating nuclear magnetic quadrupole moments require species with significant electron density at a heavy, spinful nucleus, the ability to address the resulting complicated hyperfine structure will be important for future experimental research. Photon cycling will enable significant increases in the efficiency of state preparation and detection [34,35] for ongoing experiments using ¹⁷³YbOH [19,29], and will be a critical ingredient of future improvements involving laser cooling to perform next-generation beam [36] or trap [14] measurements of this or other lasercoolable molecular species with complex hyperfine structure due to large nuclear spins.

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APPENDIX A: TABLES OF MEASURED LINES

We present transitions measured during spectroscopy in Tables III and IV.

TABLE III. Measured transitions between X''(100)N'' = 1 and A(000) in ¹⁷¹YbOH. Uncertainties are estimated to be 0.0005 cm⁻¹ due to wave-meter drift and uncertainty.

G''	F''	J'(P)	F'	Wave number (cm ⁻¹)
1	1	0.5+	1	16793.8375
1	0	0.5 +	1	16793.8400
1	2	0.5 +	1	16793.8433
1	1	0.5 +	0	16793.8505
1	0	0.5 +	0	16793.8530
1	2	0.5 +	0	16793.8563
1	1	1.5 +	1	16793.9340
1	0	1.5 +	1	16793.9365
1	2	1.5 +	1	16793.9398
1	1	1.5 +	2	16793.9522
1	0	1.5 +	2	16793.9547
1	2	1.5 +	2	16793.9580
0	1	0.5 +	1	16794.0698
0	1	0.5 +	0	16794.0828
0	1	1.5 +	1	16794.1663
0	1	1.5+	2	16794.1845

TABLE IV. Measured transitions between X''(100)N'' = 1 and A(000) in ¹⁷³YbOH. Uncertainties are estimated to be 0.0005 cm⁻¹ due to wave-meter drift and uncertainty.

G''	F''	J'(P)	F'	Wave number (cm ⁻¹)
2	1	0.5+	2	16794.0030
2	3	0.5 +	2	16794.0100
2	3	0.5 +	3	16794.0151
2	2	0.5 +	2	16794.0312
2	2	0.5 +	3	16794.0363
2	3	1.5 +	4	16794.1098
2	1	1.5 +	2	16794.1183
2	2	1.5 +	3	16794.1353
2	2	1.5 +	1	16794.1623
3	2	0.5 +	2	16794.1891
3	2	0.5 +	3	16794.1942
3	4	0.5 +	3	16794.2068
3	3	0.5 +	2	16794.2230
3	3	0.5 +	3	16794.2281
3	4	1.5 +	4	16794.3010
3	2	1.5 +	2	16794.3043
3	4	1.5 +	3	16794.3061
3	2	1.5 +	1	16794.3204
3	3	1.5 +	3	16794.3274
3	3	1.5+	2	16794.3383

APPENDIX B: TABLES OF CALCULATED X(100) ENERGY LEVELS

Based on the measured transitions, and energy levels from previous spectroscopy studies [29], we calculated energy levels relevant for this paper, as shown in Tables V and VI.

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TABLE V. Energies of levels in ¹⁷¹YbOH X(100) determined by this work. Uncertainties are estimated to be 0.0005 cm⁻¹ due to wave-meter drift and uncertainty, and excited-state energy uncertainty.

N	G	F	Wave number (cm ⁻¹)
1	0	1	529.2047
1	1	2	529.4370
1	1	0	529.4345
1	1	1	529.4312

TABLE VI. Energies of levels in ¹⁷³YbOH X(100) determined by this work. Uncertainties are estimated to be 0.001 cm⁻¹ due to wave-meter drift and uncertainty, and excited-state energy uncertainty.

N	G	F	Wave number (cm ⁻¹)
1	2	1	530.146
1	2	3	530.139
1	2	2	530.118
1	3	2	529.960
1	3	4	529.947
1	3	3	529.926

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