Laser spectroscopy of aromatic molecules with optical cycling centers: strontium (I) phenoxides

Guanming Lao,1 Guo-Zhu Zhu,1,∗ Claire E. Dickerson,2 Benjamin L. Augenbraun,3,4 Anastassia N. Alexandrova,2,5 Justin R. Caram,2,5 Eric R. Hudson,1,5,6 and Wesley C. Campbell1,5,6

1 Department of Physics and Astronomy, University of California, Los Angeles, California 90095, USA
2 Department of Chemistry and Biochemistry, University of California, Los Angeles, California 90095, USA
3 Department of Physics, Harvard University, Cambridge, MA 02138, USA
4 Harvard-MIT Center for Ultracold Atoms, Cambridge, MA 02138, USA
5 Center for Quantum Science and Engineering, University of California, Los Angeles, California 90095, USA
6 Challenge Institute for Quantum Computation, University of California, Los Angeles, California 90095, USA

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We report the production and spectroscopic characterization of strontium (I) phenoxide (SrOC6H5, or SrOPh) and variants featuring electron-withdrawing groups designed to suppress vibrational excitation during spontaneous emission from the electronic excited state. Optical cycling closure of these species, which is the decoupling of vibrational state changes from spontaneous optical decay, is found by dispersed laser-induced fluorescence spectroscopy to be high, in accordance with theoretical predictions. A high-resolution, rotationally-resolved laser excitation spectrum is recorded for SrOPh, allowing the estimation of spectroscopic constants and identification of candidate optical cycling transitions for future work. The results confirm the promise of strontium phenoxides for laser cooling and quantum state detection at the single-molecule level.

INTRODUCTION

Optical cycling transitions in atoms allow laser cooling of the center-of-mass motion, laser state preparation, and laser-induced fluorescence (LIF) state detection — open-channel operations at the heart of many promising applications of quantum technology, including quantum computation [1, 2], atomic clocks [3, 4], and quantum simulation [5, 6]. Optical cycling and cooling schemes have also been demonstrated in diatomic [7, 8] and even some small polyatomic molecules [9, 10], including SrF [11], YO [12], CaF [13, 14], YbF [15], BaF [16, 17], MgF [18], AlF [19], SrOH [20], CaOH [21], YbOH [22] and CaOCH3 [23]. Because they possess rich internal structures and complex interactions, molecules provide new opportunities in studies of dark matter detection [24, 25], measurement of electron’s electric-dipole moment [26–28], parity violation tests [29, 30], and changes to fundamental constants [31, 32].

The somewhat unexpected atom-like transitions supporting optical cycling and cooling in these small molecules have inspired searches for similar transitions in complex polyatomic molecules with an M-O-R structure [10], where M is an alkaline-earth metal atom ionically bonded to oxygen (O) forming an optical cycling center (OCC) and R is a molecular ligand [33–35]. In these molecules, the remaining metal-centered radical electron forms the highest-occupied and the lowest-occupied molecular orbitals, HOMO and LUMO. For molecules with R having strong electron withdrawing capability, the HOMO and LUMO are localized on M, which typically indicates that the OCC is highly decoupled from the vibrational degrees of freedom. As a result, the diagonal vibrational branching ratio (VBR, which is to say the probability that spontaneous decay occurs on the 0-0 transition) is high, indicating that the spontaneous emission happens without a vibrational state change. This allows such molecules to repeatedly scatter photons before being pumped to the vibrational dark states, furnishing mechanical control and state detection of single molecules via laser illumination.

Since optical cycling in this motif is predicted to be enhanced by the electron-withdrawing strength of the ligand, the diagonal VBR of M-O-R molecules could be tuned by functionalizing the ligand to promote this effect [36]. For example, according to a recent measurement of the VBRs [37], laser cooling of CaOPh-3,4,5-F3 appears feasible from the perspective that each molecule could scatter ≈1000 photons with six to eight lasers. Compared to CaOPh, the three substitutions of H → F in the 3,4 and 5 positions on the ring enhances the electron-withdrawing strength of the ligand, rendering the Ca atom more ionic and thus suppressing spontaneous decays to excited vibrational states of the electronic ground state.

As molecules of M-O-R type, the strontiummum variants, SrOPh-X, were also predicted to have high and tunable diagonal VBRs [36]. Compared to CaOPh-X, although the diagonal VBRs were predicted to be lower, the predicted difference is of the same order as the variation in measured VBRs of various calcium species [37], suggesting that some of the strontium species may show better cycle closure if the variation is due to M-specific features. Further, Sr-containing molecules allow exploration of the role of stronger spin-orbit coupling [38] and nuclear spin structures [39]. For the strontium variants, the excitation and repumping wavelengths can be directly produced by diode lasers.
RESULTS AND DISCUSSION

In the calcium- and strontium-phenoxydes, transitions to the two lowest electronic states (A and B, Figure 1a) have been proposed for laser cooling, since almost all photon scatters go back to the vibrationless ground state X [36, 37, 42]. Figure 1b shows the measured transition energies of all molecules show a linear correlation with the acid dissociation constants, pK_a, of the precursor phenol. This linear trend has recently also been observed for CaOPh-X molecules [37, 42]. Lower pK_a implies a weaker electron-withdrawing capability of the R-O ≡ ligand, which pulls the single electron away from the Sr atom, making it more ionic and increasing the HOMO-LUMO gap [36]. Also shown are excitation energies calculated by time-dependent density functional theory [40] which give a similar trend but systematically undershoot the excitation energies likely due to the lack of spin-orbit coupling (SOC) in calculations [44] and the wider difference in SrOPh-X is due to a stronger SOC effect in Sr.

To measure the VBRs from the two electronic states, we performed DLIF spectroscopy of all molecules. Electronic excitation is provided by a pulsed dye laser (PDL) tuned to the 0-0 line and the spectrometer grating was scanned in time (over repeated excitation) to select the wavelength of LIF photons sent to a photomultiplier tube (PMT) [40]. Figure 2 shows the measured DLIF spectra of SrOPh while those of other species are presented in Figure S1. Figure 2a shows the spectrum of \( A^2 B_2 \rightarrow X^2 A_1 \) of SrOPh (Figure 1a) at an excitation of 668.90 nm. The strongest peak at the origin, labeled as \( A^2 0^2 \), is due to the diagonal decay from \( A \) to \( X \) (\( \nu'' = 0 \)). The strong peak at \(-440 \text{ cm}^{-1}\) is from excited atomic Sr created during laser ablation [45]. The peak at \(-238 \text{ cm}^{-1}\) is assigned to the strongest off-diagonal stretching mode \( \nu_3 \) (theo. 241 cm\(^{-1}\)) and the weak peak at \(-54 \text{ cm}^{-1}\) is assigned to the low-frequency bending mode \( \nu_1 \) (theo. 54 cm\(^{-1}\)). The other two weak...
peaks at $-100 \text{ cm}^{-1}$ and $-297 \text{ cm}^{-1}$, which do not match the calculated frequencies of any fundamental vibrational modes, are assigned to the overtone of the bending mode $^A2_1^0$ and a combinational mode of $^A2_1^0\nu_0^0$, respectively.

Figure 2b shows the spectrum of $\tilde{B} \rightarrow \tilde{X}$ of SrOPh (Figure 1a) at 655.52 nm. Aside from the strongest diagonal peak $B0^0_0$, four peaks are observed. The strong peak with a shift of $-300 \text{ cm}^{-1}$ is due to a diagonal decay $A0^0_0$ from the $\tilde{A}$ state. The origin of the appearance of $A0^0_0$ when exciting the $\tilde{B} \rightarrow \tilde{X}$ is unknown, but could be due to the collisional relaxation from $\tilde{B}$ to $\tilde{A}$ followed by fluorescence decay to the ground state $\tilde{X}$ [37, 42, 46]. The identification of this feature as originating from the $\tilde{A}$ state is further confirmed by the observation of the decay to the stretching mode $\nu_3$ at $-534 \text{ cm}^{-1}$ from $\tilde{A}$. The other two weak peaks, $-298 \text{ cm}^{-1}$ and $-55 \text{ cm}^{-1}$, are due to the vibrational decay to the stretching mode $\nu_3$ and bending mode $\nu_1$, respectively. The full width at half maximum of all peaks is $\approx 22 \text{ cm}^{-1}$ mainly due to the spectrometer resolution of approximately $20 \text{ cm}^{-1}$. Another measurement using a narrow-band continuous-wave (cw) laser to excite the $\tilde{B} \rightarrow \tilde{X}$ of SrOPh and an electron-multiplying charge-coupled device (EMCCD) camera to capture the fluorescence photons dispersed by the spectrometer. This technique obtained a better spectral resolution ($\approx 5 \text{ cm}^{-1}$), allowing the resolution of the combinational vibrational mode of $B2^2_0\nu_0^0$ (Figure S2), which is overlapped with the diagonal decay $A0^0_0$ from the $\tilde{A}$ state and not observed in Figure 2b. The experimental and theoretical vibrational frequencies of all resolved fundamental modes are summarized in Table S1.

The relative heights of the peaks $^A0^0_0$ and $B0^0_0$ in Figure 2 and Figures S1-S2 imply that both transitions are very diagonal with few vibration-changing decays. To extract the VBRs, all peaks are fitted with Gaussian functions, as shown by the red traces in Figure 2, and the peak areas are extracted from the fits to obtain VBRs. A strict definition of VBR requires measurements of all vibrational decays. Due to finite measurement sensitivity ($\approx 10^{-2}$) and detection range ($< 600 \text{ cm}^{-1}$), while we predict that our measurement is sensitive to the dominant leakage channels, the possibility of undetected decays contributes a systematic uncertainty on the measured VBRs.

For the vibrational decays that were identified for each molecule, and the ratios of line intensities to the total intensities of all observed peaks are presented in Figure 3a. In both electronic transitions, the relative ratios of observed peaks show good agreement with the calculated VBRs. The vibrational decays to the strongest off-diagonal Sr-O stretching mode ($\nu_3$, $\nu_4$, $\nu_5$ or $\nu_6$) and the low-frequency bending mode ($\nu_1$ or $\nu_2$) have been observed for all molecules. The theoretical VBRs of the low-frequency bending modes are underestimated possibly due to the vibronic coupling and anharmonicity effect not considered in the calculation [36, 37]. SrOPh also shows unpredicted decays to the overtone of mode $\nu_2$ and a combinational mode $\nu_2\nu_3$ where the intensities could be from the vibronic coupling. The intensity ratios of all observed decays are summarized in Table S2. Figure 3b plots the estimated VBRs of the diagonal peak $0^0_0$ of each transition as a function of $pK_a$. The scaled $0^0_0$ VBRs are obtained by adding the estimated contribution of the unobserved peaks predicted by the theory to the normalized intensity ratios of the observed $0^0_0$ intensities [40]. Both SrOPh-3-F and SrOPh-3,4,5-F$_3$ molecules show VBRs $> 95\%$ for the $\tilde{B} \rightarrow \tilde{X}$ transition and $> 90\%$ for the $\tilde{A} \rightarrow \tilde{X}$ transition, while SrOPh has the lowest VBR of $82.2\%$ for $\tilde{A} \rightarrow \tilde{X}$ transition.

The VBRs for SrOPh-3-CF$_3$ shows the largest discrepancy between the calculation and the measurement, potentially due to the larger vibronic mixing between the $\tilde{A}$ and $\tilde{B}$ caused by the low symmetry and large electron inductive effect from the CF$_3$ group [37]. The error bars include both the statistical uncertainties from

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**Figure 2.** (a) and (b) Dispersed spectra of $\tilde{A} \rightarrow \tilde{X}$ and $\tilde{B} \rightarrow \tilde{X}$, respectively, of SrOPh excited by pulsed dye laser and measured by a spectrometer coupled with PMT. The experimental curves (black) are fitted with the Gaussian functions (red). The positions of the blue, vertical lines illustrate the theoretical frequencies while the intensities show the vibrational branching ratios of different vibrational modes of SrOPh. The Sr impurity peak in (a) is from the Sr emission of $5s5p^2\text{P}_1^0 \rightarrow 5s^2\text{S}_0^0$ at 689 nm [45]. The assignments of all resolved vibrational peaks are indicated.
FIG. 3. (a) Intensity ratio of observed decays for $A \to X$ and $B \to X$ transitions. Error bars are statistical errors from Gaussian fits. The vibrational label $\nu_i$ indicates the final vibrational modes of the $X$ state. $\nu_0$ implies the decay that don’t change vibrational state. (b) Scaled $0\nu_0$ VBRs as a function of $p_{K_{a}}$ of all species. The scaling adds the contributions of those unobserved vibrational decays predicted by the theory to the observed intensity ratios of $0\nu_0$ in (a). Error bars include the statistical errors from Gaussian fits and the systematic errors from the unobserved peaks [40].

FIG. 4. High-resolution rotationally-resolved excitation spectrum of the $B \leftarrow X$ transition of SrOPh. The upper trace (blue) shows the experimental spectrum and the lower trace (black) is the simulated spectrum with a Gaussian linewidth of 70 MHz and a rotational temperature $T_{\text{sim}} = 2.5$ K. Insets (a), (b) and (c) are expansions of some local features. (a) displays detailed spectrum near 0-0 transition, while (b) and (c) show the $K_{a}' = 3 \leftarrow K_{a}'' = 2$ and $K_{a}' = 4 \leftarrow K_{a}'' = 3$ rotational bandheads, respectively. (d) shows the inferred position of the candidate rotational cycling transition between the spin-rotation manifold of the $N''' = 1$ state and $N'' = 0$ state.

To further investigate the potential of these species for optical cycling, a high-resolution excitation spectrum (obtained by collecting LIF as a continuous-wave (cw) excitation laser is scanned) of SrOPh for the $B(v' = 0) \leftarrow X(v'' = 0)$ transition is recorded at a step size of 25 – 50 MHz in a cryogenic buffer-gas beam (CBGB) [40, 47] and fitted with PGOPHER [41], as presented in Figure 4. Since SrOPh is an asymmetric-top molecule,
the rotational states are labeled as \( N\,K_aK_c \), where \( N \) is the rotational angular momentum, \( a \) and \( c \) label the inertial axes lying along the Sr-O bond and perpendicular to the molecular plane (Figure 1a), respectively, \( K_a \) and \( K_c \) are the projection of \( N \) onto the two axes in the prolate and oblate limits, respectively. Figure 4a shows the expansion of the two congested bands at 15238.5 cm\(^{-1}\), while Figures 4b,c show two well-resolved rotational bands. A full rotational analysis is difficult due to the high density of rotational lines in the middle of the spectrum (Figure 4a), but the individually resolved lines (Figures 4b,c) make it possible to fit the spectrum to extract some spectroscopic constants.

Using a custom program to fit the spectral contour and pgopher [41] to refine and iterate the line assignments [40], we have assigned nearly 400 rotational transitions and obtained the final fitted spectrum given as the black traces in Figure 4. The fitting is in agreement with the experimental measurement for the middle broad bands and the \( K_a' = 3 \leftarrow K_a'' = 2 \) and \( K_c' = 4 \leftarrow K_c'' = 3 \) bands, as expanded in Figures 4a-c. The best fit molecular constants, including the transition energy, rotational constants, spin-rotational constants and centrifugal distortion corrections, are reported in Table 1. The measured rotational constants are in good agreement with the calculated values. The spin-rotation constant \( \epsilon_{aa} \) in the ground state is too small to be determined from the spectrum, and \( \epsilon_{aa} \) in the \( B \) state is large because of the coupling to the \( A \) state. The larger value of spin-rotational constant than the rotational constants in \( B \) implies a strong SOC effect apart from the direct coupling between the spin and molecular rotation. Based on the second order perturbation theory [48, 49] and the measured constants, the SOC constant in SrOPh is estimated to be \( \approx 272 \) cm\(^{-1}\), which is close to that of SrOH (\( A^3 \Pi \approx 265 \) cm\(^{-1}\)) [50]. The large SOC also dominates the energy separation of \( A - B \), elucidating the discrepancy between the calculation and the measurement in Figure 1b [40, 44].

While involving more parameters has been able to enhance the accuracy of fitting, many parameters in such scenarios tended to fit to values consistent with zero, and we therefore omit those in our analysis. The rotational temperature from the fit is 2.5 K [40]. The colder temperature is due to the free expansion of neon buffer gas from the cryogenic cell (\( \approx 23 \) K) to form a beam with SrOPh entrained [47]. As the SrOPh \( B \leftarrow X \) transition dipole moment lies along the principle axis \( c \) (Figure 1b), the rotationally closed photon cycling transition is the \( c \)-type transition \( N\,K_aK_c = 000 - N\,K_a',K_c' = 100 [35] \), which is estimated to be at 456.8391(7) THz based on the fitting results and shown in Figure 4d.

### Table 1. Molecular constants of SrOPh obtained by fitting the rotationally-resolved excitation spectrum in Figure 4 with pgopher. All quantities are presented in cm\(^{-1}\).

| Constant | \( B^2B_2 \) | \( X^2A_1 \) |
|----------|----------------|----------------|
|          | Exp. | Cal. | Exp. | Cal. |
| \( T_0 \) | 15238.7155(23) | 15238.7155(23) | 1.28(20) | 1.28(20) |
| \( A \) | 0.1923(6) | 0.1915 | 0.1934(11) | 0.1916 |
| \( \frac{1}{2}(B + C) \) | 0.01520(36) | 0.01522 | 0.01508(36) | 0.01513 |
| \( (B - C) \times 10^3 \) | 1.28(20) | 1.21 | 1.13(12) | 1.19 |
| \( \epsilon_{aa} \) | -0.6894(6) | - | - | - |
| \( \epsilon_{bb} \times 10^3 \) | 34(10) | 34(10) | 1.3(1.7) | 1.3(1.7) |
| \( \epsilon_{cc} \times 10^3 \) | 16(7) | 16(7) | -1.3(1.8) | -1.3(1.8) |
| \( D_N \times 10^8 \) | -14(8) | -14(8) | -14(8) | -14(8) |
| \( D_{NK} \times 10^7 \) | -5(11) | -5(11) | -28(23) | -28(23) |
| \( D_K \times 10^4 \) | 1.3(5) | 1.3(5) | 5.2(1.1) | 5.2(1.1) |
| \( H_K \times 10^6 \) | 3.0(1.4) | 3.0(1.4) | 21(4) | 21(4) |

**T_0**: electronic transition energy; \( A, B, C \): molecular rotational constants; \( \epsilon_{aa}, \epsilon_{bb}, \epsilon_{cc} \): spin-rotation coupling constants; \( D_N, D_{NK}, D_K \): centrifugal distortion constants; \( H_K \): sextic centrifugal distortion correction.

**CONCLUSION**

In summary, we have produced strontium (I) phenoxide (SrOPh) and derivatives featuring electron-seeking withdrawing groups in a cryogenic cell. Two proposed laser cooling transitions (\( A - X \) and \( B - X \)) of each molecule have been identified and the transition energies show linear trends as the ligand pKa, which can be used to look for transitions of new molecules containing Sr. The overall vibrational branching ratios considering contributions of unobserved vibrational decays are estimated to be 82.2% - 95.8%. Among them, SrOPh-3-F and SrOPh-3,4,5-F_3 molecules show diagonal VBRs > 95%, potentially enabling laser cooling with fewer than ten rotational repumping lasers. The rotationally-resolved spectrum for the \( B \leftarrow X \) transition of SrOPh is presented and molecular constants are obtained. The spin-orbit interaction that couples the \( A \) and \( B \) states is estimated to be 275 cm\(^{-1}\), which has a strong effect on the energy splitting of \( A - B \). The rotational closure transition for optical cycling is estimated to be centered near 456.8391 (7) THz. This work paves the way for optical cycling of SrOPh and other large molecules using diode lasers.

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Molecule production. All SrOPh-X molecules were produced by the reaction of Sr atoms generated by laser ablation of a metallic Sr metal chunk using the Minilite pulsed Nd:YAG laser at 1064 nm (pulse energy $\approx 6$ mJ, repetition rate 10 Hz) with different ligand precursors in a cryogenic buffer-gas cell operated at $\approx 23$ K. Five ligands – phenol, $m$-cresol, 3-fluorophenol, 3-(trifluoromethyl)phenol and 3,4,5-trifluorophenol – purchased from Sigma Aldrich were individually heated in a reservoir to supply the respective vapors, which were guided via a heated gas line into the cryogenic cell with a density $\approx 10^{13}$ cm$^{-3}$. The reaction products were then cooled by colliding with a neon buffer gas of density $\approx 10^{15}$–$16$ cm$^{-3}$.

**DLIF measurement in the cryogenic cell.** The resulting SrOPh-X molecules have two low-lying electronic states proposed for laser cooling. To look for those states, a tunable, pulsed dye laser (10 Hz, LiopStar-E dye laser, linewidth 0.04 cm$^{-1}$ at 620 nm) were used to excite molecules in the cryogenic cell and the laser wavelength was scanned. When the laser wavelength hit electronic resonance, the molecule was excited and followed by the emission of molecular fluorescence. The fluorescence was then collected via an imaging system into a model 2035 McPherson monochromator equipped with a 1200 lines/mm grating and detected by a PMT. The dispersed measurement were done by parking the laser wavelength at the electronic resonance and continuously scanning the grating of the spectrometer at an increment of 0.10 nm while monitoring the fluorescence photons. The entrance and exit slit widths were both set at 0.20 mm, resulting in a spectrometer resolution of $\approx 20$ cm$^{-1}$.

A different method was also used to measure the DLIF spectrum for SrOPh $\bar{B} \rightarrow \bar{X}$ transition (Figure S2). In this method, the SrOPh molecules were excited by a cw laser illuminated from a home-build external-cavity diode laser (ECDL). The fluorescence was dispersed by the monochromator at a fixed grating position and detected by an EMCCD camera. The entrance slit of the spectrometer was set to 0.03 mm to achieve a resolution of $\approx 3$ cm$^{-1}$. In this measurement, all fluorescence photons were collected simultaneously by the EMCCD, which avoided the systematic errors due to the ablation or PDL energy drift in scanning the grating in the first method.

**High resolution excitation spectroscopy of SrOPh $\bar{B} \rightarrow \bar{X}$.** To look for the rotational closure transitions, We performed the high-resolution excitation spectroscopy of SrOPh in a molecular beam. SrOPh molecules formed in the cryogenic cell were extracted out via a 9 mm cell aperture and entrained into a neon buffer gas beam. The excitation zone is $\approx 23$ cm downstream the cell aperture. A cw laser from an ECDL was scanned with an increment of $25$–$50$ MHz near the $\bar{B}(v' = 0) - \bar{X}(v'' = 0)$ transition of SrOPh determined by the low-resolution PDL measurement. The fluorescence were collected by a PMT at the perpendicular direction. Due to the cooling effect in the expansion, SrOPh molecules in the beam are colder than those thermalized in the cell at a temperature $\approx 23$ K.

**SUPPLEMENTARY INFORMATION**

**A. Experimental methods**

**Molecular geometries, excitation energies, and Franck-Condon factor (FCFs) calculations**

Molecular geometries, excitation energies, and Franck-Condon factor (FCFs) calculations were performed in Gaussian16 [51]. Density functional theory (DFT) was used for the ground states while time-dependent DFT was used for excitation energies of excited states. A superfine grid was used with the PBE0-D3 functional with dispersion corrections and the def2-TZVPPD basis set [52–55]. An effective core potential (ECP) was used for the Sr atom within the def2-TZVPPD basis set.
Molecular orbitals were generated with an isosurface of 0.03 in the Multiwfn program [56]. The FCFs calculated included Duschinsky rotations, which seems to be sufficient for an overall trend in FCF. However, the anharmonicity and vibronic coupling effects do play a key role in the the FCFs of the low-frequency bending modes [36, 37], which were underestimated here in comparison to the experimental measurements. The calculated FCFs can be converted to the VBRs using the formula [35, 37]:

\[
b_{i\nu',f\nu''} = \frac{A_{i\nu',f\nu''}}{\sum_{f\nu'} A_{i\nu',f\nu''}} = \frac{|\mu_{i\nu',f\nu''}|^2 \times (\nu_{i\nu',f\nu''})^3}{\sum_{f\nu'} |\mu_{i\nu',f\nu''}|^2 \times (\nu_{i\nu',f\nu''})^3} \approx \frac{\text{FCF}_{i\nu',f\nu''} \times \nu_{i\nu',f\nu''}^3}{\sum_{f\nu'} \text{FCF}_{i\nu',f\nu''} \times \nu_{i\nu',f\nu''}^3}
\]

where \(i\) and \(f\) imply the initial and final states, respectively. \(b_{i\nu',f\nu''}\) is the branching ratio, \(A_{i\nu',f\nu''}\) is the Einstein coefficient for spontaneous emission, \(\mu_{i\nu',f\nu''}\) is the transition dipole moment and \(\nu_{i\nu',f\nu''}\) is the transition frequency.

C. DLIF spectra of other molecules

Figure S1 presents the DLIF spectra of other four molecules, which were recorded by monitoring the fluorescence signal from the respective excited states when scanning the grating wavelength at a step size of 0.1 nm. All peaks were fitted with Gaussian functions, as shown by the red traces overlapped with the experimental black traces. Comparing to the theoretical vibrational frequencies and the respective VBRs (blue lines), those peaks can be readily assigned. Take SrOPh-3-CH\(_3\) for an example, two peaks show up in the \(\tilde{A} \rightarrow \tilde{X}\) decay in Figure S1a. The origin peak labeled as \(A_{000}\) represents the decay of \(A(v'=0) \rightarrow \tilde{X}(v''=0)\), while a weak peak at a frequency shift of \(-226\) cm\(^{-1}\) matches well with the theoretical vibrational frequency of the Sr-O stretching mode \(\nu_5\) (230 cm\(^{-1}\)), which is also the most off-diagonal vibrational mode. In Figure S1b of \(B \rightarrow \tilde{X}\) decay, except for the diagonal peak \((B_{000})\) at the origin and the stretching-mode peak \((B_{01}^0)\) at \(-230\) cm\(^{-1}\), two additional peaks are observed. The strong peak at \(-300\) cm\(^{-1}\) is assigned to \(A_{000}\), which is always observed and due to the collisional relaxation from \(B\) to \(\tilde{A}\) followed by the radiative decay. A very weak peak at \(-43\) cm\(^{-1}\) is the decay to the low-frequency bending mode \(\nu_2\) (50 cm\(^{-1}\)). The similar spectra features of decays to the most-off diagonal Sr-O stretching mode and the low-frequency bending mode and the appearance of the \(A_{000}\) peak when exciting to \(\tilde{B}\) have been observed for all other molecules. A more complex decay scenario is observed for SrOPh-3-CF\(_3\) mainly due to the low-symmetry structure introduced by the electron-withdrawing group of CF\(_3\). More vibrational decays have been observed. In Figure S1c, three fundamental vibrational modes have been resolved from the decay of \(A\) and assigned to \(\nu_2\), \(\nu_3\) and \(\nu_0\). The decay from the \(\tilde{B}\) state is more complicated due to the relaxation decay to the \(\tilde{A}\) state. Except for the same vibrational decays from \(\tilde{B}\), three more peaks are observed. The strongest peak at \(-300\) cm\(^{-1}\) is due to the collisional relaxation and fluorescence pathways of \(\tilde{B} \rightarrow \tilde{A} \rightarrow \tilde{X}\). The high intensity is caused by the large CF\(_3\) group which increases the collisional relaxation rate from the \(\tilde{B}\) state. Two other weak peaks are assigned to \(A_{52}^0\) and \(A_{63}^0\) due to the relaxation from \(\tilde{B}\) to the vibrationally excited states of \(\tilde{A}\) followed by the fluorescence decays. The vibrational frequencies and VBRs of all observed vibrational modes are given in Tables S1 and S2.

D. Error analysis of VBRs

All observed peaks in DLIF spectra in Figures 2 and S1 are fitted with the Gaussian function using the parameters of peak location, height and width. For the \(i\)th peak in each spectrum, the peak area \((M_i)\) is extracted and the error of the area \((\delta M_i)\) is estimated from the covariance matrix of the fitting parameters. The intensity ratios of each peak, as shown in Figure 3a, is obtained from the ratio of \(M_i\) and the total area of all the observed peaks, \(\sum_{i=0}^{p} M_i\). The statistical error of each intensity ratio is then calculated from the relative uncertainties \(\delta M_i/M_i\). Besides the statistical fitting errors, several sources of systematic error in the DLIF measurement are discussed and listed in Table S3.

The first systematic error come from the unobserved peaks which contribute to the VBRs. A true VBR depends on contributions of all possible decay pathways. Due to a low measurement sensitivity and a small detection window, only a few vibrational decays have been observed for each transition. Compared to a complete description of vibrational decays obtained from calculated FCFs, all unobserved vibrational decays are therefore a source of the systematic uncertainty, which is estimated by [37]:

\[
S'_0 = \frac{S_0}{\sum_{i=0}^{p} S_i + \sum_{i=p+1}^{N} \frac{C_i}{T_0}}.
\]
diagonal VBR, while the scaled VBR $S'_0$ is used as the plot data points in Figure 3. As summarized in Table S3, the uncertainties of the unobserved peaks as differences of $S_0$ and $S'_0$ are in the range of 1.1 – 3.2%.

Another source of systematic uncertainty is the signal drifting in the measurement due to the change of experimental conditions. Except for the DLIF measurement of SrOPh $B \rightarrow \bar{X}$ by the cw laser and EMCCD (Figure S2), all other DLIF spectra were taken by scanning the grating of the spectrometer to disperse photons onto a PMT. This means that the fluorescence photons at different wavelengths were not detected simultaneously. A typical scan of 15 nm wavelength range would take 75 minutes with an increment of 0.1 nm and 300 averages for each wavelength. During the scan, the signal was slowly drifting mainly due to the dust accumulation on the imaging lens and the pulse intensity drifting from both the ablation and the excitation lasers. We kept track of the signals before and after the whole scan and found that the signal change can vary by up to 20%, which can lead to an error of 1.0% in VBR estimation as such signal change affects the off-diagonal vibrational transition signal. In addition, the dispersed photons at different wavelength were detected simultaneously in the EMCCD measurement, which eliminates the error due to signal drifting. The error can be estimated by the difference of the diagonal VBR of SrOPh $B \rightarrow \bar{X}$ from the two different methods, which is 1.2%.

As discussed in the error analysis of CaOPh-X [37], the wavelength response of the spectrometer and the imperfection of the mirrors and lenses in the imaging system could cause a systematic error up to $\approx 1\%$. The last error source comes from the diagonal excitation of vibrationally excited modes in the ground state. Due to a cell temperature of $\approx 23$ K, the thermalized molecules can have thermal populations of $\approx 5\%$ of the low-frequency bending mode and $\approx 10^{-7}$ of the stretching mode. Those vibrationally excited states in the $\bar{X}$ state could be near-resonantly excited by the pulsed dye laser to the same vibrational levels of the upper states. The following decays from those diagonal excitations can cause an error up to 0.5%.

By adding the four systematic errors in quadrature, a total systematic uncertainty is estimated to be 1.9% – 3.5%. Considering the statistical uncertainties from the Gaussian fits, the final upper and lower bounds of uncertainties for the diagonal $0^0\bar{S}$ VBRs are obtained and plotted in Figure 3b.

### E. Fitting of high-resolution excitation spectrum of SrOPh

The rotationally-resolved excitation spectrum of SrOPh has been fitted to estimate the molecular constants. The energy levels of SrOPh $B$ and $X$ states are computed from an effective Hamiltonian which includes rotation, electron spin-rotation coupling and centrifugal distortion correction terms:

$$H_{\text{eff}} = H_{\text{Rot}} + H_{\text{SR}} + H_{\text{cd}}.$$  \hfill (3)

The rotational Hamiltonian is

$$H_{\text{Rot}} = AN_a^2 + BN_b^2 + CN_c^2,$$ \hfill (4)

the spin-rotation coupling term is

$$H_{\text{SR}} = \epsilon_{aa} N_a S_a + \epsilon_{bb} N_b S_b + \epsilon_{cc} N_c S_c,$$ \hfill (5)

and the centrifugal distortion correction used in this work is

$$H_{\text{cd}} = -D_N N^2(N+1)^2 - D_N K N(N+1)K_a^2 - D_K K_a^4 + H_K K_a^6,$$ \hfill (6)

here the sextic centrifugal distortion correction $H_K$ is included to enhance the fitting quality of the bandheads of the observed $\Delta K_a = \pm 1$ bands.

Although a few rotational bands recognized in the SrOPh excitation spectrum can determine a few rotational constants in good accuracy, the high density of rotational line near 0-0 transition ($\sim 10^7$/GHz in the computing limit of $J_{\text{max}} = 30$) is the main challenge for fitting the rest of the parameters. Therefore, we used a homemade program first to search for parameters that can roughly fit the spectrum in contour and limit the searching range of the parameters for PGOPHER’s subsequent fitting [41]. To avoid the program being trapped by local minima, two algorithms were used in turns: the mini-batch gradient descent (MBGD) [57] and the genetic algorithm (GA) [58]. MBGD works similarly as the well-known gradient descent method, while its gradient is computed from a batch of randomly chosen data instead of the whole data set during each iteration to jump out of local minima with semi-stochastic steps. MBGD is the main algorithm that searches for a potential solution iteratively, and GA checks whether the MBGD result is optimal within a larger parameter space. The target functions of the two algorithms are different, with the purpose of making their local minima to be also different.

A contour fitting result is accepted as the initial input of PGOPHER if it is agreed by both MBGD and GA. In PGOPHER, the fitting of molecular constants is done in following procedure: first, the clearly observed bandheads are matched to the simulated rotational bands, such as the lines labeled in Figure 4b and 4c. With such assignment PGOPHER can calculate parameters $T_0$, $A$, $B$, $\epsilon_{aa}$, $D_K$ and $H_K$ in a better accuracy than the contour fitting. Next, the rest of the parameters are obtained by some details of the spectrum near the 0-0 transition, such as the spacing of lines in different rotational bands, shape of peaks, and the order of line strength. For concreteness, some strong peaks in the middle of the spectrum (Figure S4a) are assigned to the transitions with $K_a'' = 0, 1, 2$ and different P, Q and R branches (P, Q, and R labels refer transitions with $\Delta J = -1, 0$ and 1, respectively): the
strongest few peaks in Q branch are assigned to some observed peaks, see Figure S4b; and for the P and R branch transitions, the line assignment could be made according to some local features, see Figure S4c.

Fine adjustment of fitting is achieved by adjusting the parameters, re-assigning some lines or bands tentatively, and running the fit based on the updated assignment. This procedure should be repeated multiple times before the parameters become converged.

The rotational temperature in pGopher simulation is set based on the normalized strength of the rotational bandhead of $K_a' = 6 \rightarrow K_a'' = 5$, the farthest band we can recognize in experiment. It is found to be close to the experimental results when $T_{\text{sim}}$ is around 2 – 3 K. In the contour fitting procedure, the rotational temperature is estimated to be 4 – 7 K, depending on the fitting condition such as linewidth and the upper limit of the rotational quantum number $J_{\text{max}}$.

The estimated molecular constants reported in Table I are from the best fit result whose simulated spectrum pattern matches most of observed peaks near the 0-0 transition, with the assignment error bars calculated from the standard errors of the estimated values given by multiple fitting attempts. These attempts follow the same rotational bandhead assignment in the first step and have 

The estimated molecular constants reported in Table I are from the best fit result whose simulated spectrum pattern matches most of observed peaks near the 0-0 transition, with the assignment error bars calculated from the standard errors of the estimated values given by multiple fitting attempts. These attempts follow the same rotational bandhead assignment in the first step and have a similar P, Q, R line distributions depicted in Figure S4, while the numbers and the positions of assigned lines near the 0-0 transition are varied to reflect the parameter fluctuations caused by different assignments.

F. Spin-orbit coupling effect in SrOPh

According to the second order perturbation theory [48, 49], the relation between the SOC constant $A_{\text{so}}$ and the effective spin-rotation constant of the $\tilde{B}$ state could be estimated by the following equation:

$$\epsilon_{\tilde{B}a} \approx -\frac{4AA_{\text{so}}}{E_B - E_{\tilde{A}}}.$$  \hspace{1cm} (7)

With the measured constants, the SOC constant in SrOPh estimated by Eq.(7) is $A_{\text{so}} \approx 272\text{cm}^{-1}$, which is similar to the $A_{\text{so}}$ of the SrOH $A^2\Pi$ state ($\approx 265\text{cm}^{-1}$) [50].

The SO interaction can also explain why the energy gaps between $\tilde{A}$ and $\tilde{B}$ states ($\approx 300\text{cm}^{-1}$) of all the species shown in Figure 1b are much larger than their prediction. In the calculation, the electronic energies are computed without SOC effect, and the excited state wavefunctions, conventionally labeled as $|A^2B_2\rangle$ and $|B^2B_1\rangle$, are assumed to follow the $C_{2v}$ symmetry. However, the SO interaction strongly mixes the two states and breaks the $C_{2v}$ symmetry in these wavefunctions. We can use a simple two level system model to demonstrate such mixing, in which the total electronic Hamiltonian is contributed by the electronic Hamiltonian ($H_{\text{el}} = E_A|A^2B_2\rangle\langle A^2B_2| + E_B|B^2B_1\rangle\langle B^2B_1|$) and the SO interaction Hamiltonian:

$$H_{\text{tot}} = H_{\text{el}} + H_{\text{so}}.$$  \hspace{1cm} (8)

here the SO coupling Hamiltonian is $H_{\text{so}} = A_{\text{so}}L_aS_a$, $L_a$ and $S_a$ are the projection operators of the orbital and spin angular momentum onto the principle axis $a$, respectively. As written in the basis of the electronic states $\{|A^2B_2\rangle, |B^2B_1\rangle\}$, $H_{\text{tot}}$ reads [44]:

$$H_{\text{tot}}(\Sigma, \Omega) = \left( \begin{array}{cc} E_A & A_{\text{so}} \Sigma \Omega \\ A_{\text{so}} \Sigma \Omega & E_B \end{array} \right) = \frac{E_B + E_A}{2} + \frac{1}{2} \left( \begin{array}{cc} -\Delta E_0 & A_{\text{so}} \\ A_{\text{so}} & \Delta E_0 \end{array} \right),$$  \hspace{1cm} (9)

where $\Delta E_0 = E_B - E_A$ is the energy separation between the $|A^2B_2\rangle$ and $|B^2B_1\rangle$ states without the SO interaction, and it could be regarded as the splitting caused by the molecular asymmetry. $\Sigma$ is the projection quantum number of electronic spin onto the principle axis $a$, and we take $\Sigma = \frac{1}{2}$ since its sign does not affect the energy levels. According to Eq.(9), the energy difference between the two eigenstates of $H_{\text{tot}}$, $\tilde{A}$ and $\tilde{B}$, could be calculated by the following equation:

$$\Delta E = E_{\tilde{B}} - E_{\tilde{A}} = \sqrt{(\Delta E_0)^2 + A_{\text{so}}^2}. $$  \hspace{1cm} (10)

For SrOPh, the total energy gap is $\Delta E = 305\text{cm}^{-1}$, which then gives $\Delta E_0 = 138\text{ cm}^{-1} \ll A_{\text{so}}$. This implies that the orbital angular momentum is partially preserved by the strong SO interaction in the SrOPh $A(\tilde{B})$ state. Therefore, the SrOPh $A(\tilde{B})$ state is similar to a $^2\Pi|\Omega=\frac{1}{2}\rangle (^2\Pi|\Omega=-\frac{1}{2}\rangle$ state in the symmetric top approximation. Noticed that $\Delta E$ is also much larger than $\Delta E_0$ in the other species studied in this work, the SO interaction is expected to dominate the separation between the $\tilde{A}$ and $\tilde{B}$ states of all these species.
TABLE S1. Comparison of the observed and calculated frequencies for resolved fundamental vibrational modes of all species studied in this work. Values are given in units of cm$^{-1}$.

| Vib. modes | SrOPh | | SrOPh-3-CF$_3$ | | Vib. modes | SrOPh-3-CF$_3$ |
|------------|-------|-------|----------------|-------|---|-------|----------------|
| $\nu_2$    | 54(2) | 54    | 42(5)          | 39    | $\nu_2$ | 42(5) | 39 |
| $2\nu_2$   | 102(2)| 108   | 178(5)         | 180   | $\nu_5$ | 219(2) | 222 |
| $\nu_3$    | 238(2)| 241   | 219(2)         | 222   | $\nu_6$ | 226(2) | 230 |
| $\nu_2\nu_3$ | 297(2) | 294  | 263(2)         | 270   |

| Vib. modes | SrOPh-3-F | | SrOPh-3-CH$_3$ | | Vib. modes | SrOPh-3-CH$_3$ |
|------------|-----------|-------|----------------|-------|---|-------|----------------|
| $\nu_2$    | 56(5)     | 56    | 43(5)          | 50    | $\nu_2$ | 43(5) | 50 |
| $\nu_3$    | 226(2)    | 222   | 226(2)         | 230   |

| Vib. modes | SrOPh-3,4,5-F | | | | Vib. modes | SrOPh-3,4,5-F |
|------------|---------------|-------|----------------|-------|---|-------|----------------|
| $\nu_2$    | 47(6)         | 45    |                |       |
| $\nu_4$    | 203(2)        | 204   |                |       |

TABLE S2. The intensity ratios of all observed vibrational decays of all molecules. The errors indicate the statistical uncertainties from the Gaussian fits. The theoretical VBRs are also added for comparison.

| Modes SrOPh-CH$_3$ | Exp.(A) | Theo.(A) | Exp.(B) | Theo.(B) |
|-------------------|---------|----------|---------|----------|
| 0                 | 0.946(6)| 0.926    | 0.918(9)| 0.943    |
| $\nu_2$           | 7 x 10$^{-4}$ | 0.011(2)  | 2 x 10$^{-4}$ | 0.011(2)  |
| $\nu_5$           | 0.054(6)| 0.059    | 0.072(9)| 0.044    |

| Modes SrOPh | Exp.(A) | Theo.(A) | Exp.(B) | Theo.(B) |
|-------------|---------|----------|---------|----------|
| 0           | 0.845(7)| 0.928    | 0.885(5)| 0.945    |
| $\nu_2$     | 0.029(3)| < 10$^{-4}$ | 0.054(3) | < 10$^{-4}$ |
| $2\nu_2$    | 0.025(3)| < 10$^{-4}$ | 0.060(3)| 0.043    |
| $\nu_3$     | 0.079(5)| 0.059    | 0.060(3)| 0.043    |
| $\nu_2\nu_3$| 0.021(3)| < 10$^{-4}$ | 0.035(3)| 0.037    |

| Modes SrOPh-3-F | Exp.(A) | Theo.(A) | Exp.(B) | Theo.(B) |
|-----------------|---------|----------|---------|----------|
| 0               | 0.956(13)| 0.936   | 0.965(3)| 0.954    |
| $\nu_1$         | 0.007(3)| 9 x 10$^{-4}$ | 0.035(3)| 0.037    |
| $\nu_3$         | 0.037(16)| 0.051   | 0.035(3)| 0.037    |

| Modes SrOPh-CF$_3$ | Exp.(A) | Theo.(A) | Exp.(B) | Theo.(B) |
|-------------------|---------|----------|---------|----------|
| 0                 | 0.893(5)| 0.939    | 0.867(11)| 0.950    |
| $\nu_2$           | 0.014(2)| 0.003    | 0.021(5) | < 10$^{-4}$ |
| $\nu_5$           | 0.018(2)| 0.007    | 0.026(6)| 0.007    |
| $\nu_6$           | 0.074(3)| 0.037    | 0.086(8)| 0.035    |

| Modes SrOPh-3,4,5-F | Exp.(A) | Theo.(A) | Exp.(B) | Theo.(B) |
|---------------------|---------|----------|---------|----------|
| 0                   | 0.946(7)| 0.945    | 0.964(4)| 0.963    |
| $\nu_1$             | 0.005(2)| < 10$^{-4}$ | 0.008(2)| < 10$^{-4}$ |
| $\nu_4$             | 0.049(7)| 0.044    | 0.028(4)| 0.030    |

TABLE S3. Systematic error budget for the vibrational branching ratio measurements.

| VBRs measurement error source | Percentage |
|------------------------------|------------|
| Contributions from unobserved peaks | 1.1% – 3.2% |
| Signal fluctuation           | 1.0%       |
| Instrument wavelength response| 1.0%       |
| Diagonal excitation           | 0.5%       |
| Total error                  | 1.9% – 3.5% |
TABLE S4. Measured intensity ratios and scaled VBRs of the diagonal 0-0 decay of all molecules. The scaling process considering contributions of unobserved vibrational decays is detailed in the section of error analysis of VBRs.

| Molecules       | Measured $VBR_{S0}$ | Scaled $S'_0$ | Scaled excluding main peak $S''_0$ | Upper error bar | Lower error bar |
|-----------------|---------------------|---------------|------------------------------------|-----------------|-----------------|
| SrOPh-3-CH$_3$  | 0.946(6)            | 0.933         | 0.933                              | 0.015           | 0.006           |
| SrOPh           | 0.845(7)            | 0.822         | 0.804                              | 0.024           | 0.020           |
| SrOPh-3-F       | 0.956(13)           | 0.943         | 0.941                              | 0.019           | 0.013           |
| SrOPh-3-CF$_3$  | 0.893(5)            | 0.868         | 0.857                              | 0.026           | 0.012           |
| SrOPh-3,4,5-F   | 0.946(7)            | 0.930         | 0.924                              | 0.017           | 0.009           |

| Molecules       | Measured $VBR_{S0}$ | Scaled $S'_0$ | Scaled excluding main peak $S''_0$ | Upper error bar | Lower error bar |
|-----------------|---------------------|---------------|------------------------------------|-----------------|-----------------|
| SrOPh-3-CH$_3$  | 0.918(9)            | 0.896         | 0.880                              | 0.023           | 0.019           |
| SrOPh           | 0.885(5)            | 0.869         | 0.859                              | 0.017           | 0.012           |
| SrOPh-3-F       | 0.965(3)            | 0.958         | 0.958                              | 0.008           | 0.003           |
| SrOPh-3-CF$_3$  | 0.867(11)           | 0.853         | 0.840                              | 0.018           | 0.017           |
| SrOPh-3,4,5-F   | 0.964(4)            | 0.953         | 0.950                              | 0.011           | 0.005           |
FIG. S1. Dispersed fluorescence spectra of all species. The experimental curves (black) are fitted with the Gaussian functions (red). The blue sticks illustrate the vibrational branching ratios of different vibrational modes. The assignments of resolved vibrational peaks are also given.
FIG. S2. Dispersed spectrum of SrOPh $\tilde{B} \rightarrow \tilde{X}$ excited by cw laser and measured by a spectrometer coupled with an EMCCD camera. The inset shows the expansion of a broad peak at $-300$ cm$^{-1}$, which is due to the overlapping of two peaks. The assignments of the resolved vibrational peaks are also given.
FIG. S3. Fluorescence decay traces of all the transitions studied in this work. The experimental data points (red circles) are obtained from the sum of PMT signal for the $0 \rightarrow 0$ decay in the DLIF measurements and the respective error bar represents the standard errors. For each trace, the data points are normalized to the maximum signal counts. The radiative lifetimes $\tau$ and errors for all species are estimated from exponential fits (green curves) by bootstrapping the data.
FIG. S4. A typical line assignment near 0-0 transition of SrOPh. The experimental (a) and simulated data (b,c) are normalized to the maximum signal strength in experiment and simulation, respectively. The observed peaks labeled with green, purple and red ticks in the (a) measured spectrum are assigned to the simulation peaks of (b) Q, (c) R and P branch transitions, respectively. The Gaussian linewidth for the simulation is set as 70 MHz to roughly fit the contour, with the rotational temperature set as $T_{\text{sim}} = 2.5$ K. Each simulation peak in this range usually contains multiple rotational lines, to avoid overfitting, only the strongest 2 - 4 lines in each peak are assigned to the corresponding observed peak.