Piezospectroscopic measurement of high-frequency vibrations in a pulse-tube cryostat

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Vibrations in cryocoolers are a recurrent concern to the end user. They appear in different parts of the acoustic spectrum depending on the refrigerator type, Gifford McMahon or pulse-tube, and with a variable coupling strength to the physical system under interest. Here, we use the piezospectroscopic effect in rare-earth doped crystals at low temperature as a high resolution, contact-less probe for the vibrations. With this optical spectroscopic technique, we obtain and analyze the vibration spectrum up to 700kHz of a 2kW pulse-tube cooler. We attempt an absolute calibration based on known experimental parameters to make our method partially quantitative and to provide a possible comparison with other well-established techniques.

I. INTRODUCTION

Compared to liquid helium cryostats, cryocoolers open many perspectives in the scientific community. First, closed-cycled systems allow to access cryogenic temperatures even when liquid helium is not available on-site. They can also be operated continuously with very limited maintenance, which is mandatory in many integrated applications or when remote control is required. The price to pay is a higher level of vibration for the cryocoolers compared to their wet equivalent. The measurement and the reduction of the vibrations in a cryocooler are active subjects of research, finding applications in very different fields including metrology for gravitational wave detection, the definition of high stability oscillators, or as a routine instrument for fundamental research.

These prospects have motivated many studies to accurately measure the vibration spectrum and to model it up to few tens of kHz. The noise is essentially dominated by the low-frequency components driven by the compression cycle. Higher frequencies are also present because of high-order harmonics of the tubes’ mechanical distortion modes and smaller parts of the apparatus with higher resonant frequencies. The high frequency region of the spectrum may have an impact on micro or nano-mechanical resonators investigated at low temperatures to explore the quantum mechanical nature of massive objects, or cryogenic mirrors for gravitational wave observation, since both could have resonances falling into the high frequency range.

More fundamentally, proper modeling requires to measure a large spectrum in order to design accordingly a passive stabilization scheme. Active stabilization offers an alternative and elegant solution. This technique also requires the largest acquisition bandwidth to obtain a fast and reliable servo-control.

Accelerometers represent a commercial solution for measuring vibrations that are already compatible with cryogenic temperatures. They are routinely used to produce technical reports and allow a rapid comparison between different mechanical assembly of cryocoolers. When higher measurement bandwidths are targeted, optical techniques are particularly interesting. Fast response is guaranteed by current optoelectronics components, but they also offer a fine spatial resolution given by the beam size and an intrinsic nondestructive, contact-less character. Optical measurements are of two types: interferometric, when the surface under study reflects light as the end mirror of a Michelson interferometer, for example or intensity sensor, usually based on fiber-coupled excitation and detection port. Both give access to the kHz frequency range, up to 20kHz, at least an order of magnitude better than most accelerometers. The detection bandwidth of optical techniques could also be pushed to higher frequencies, away from the audible spectrum, which have been mostly unexplored despite the absence of clear technical limitations.

We propose a new non-interferometric optical approach based on the piezospectroscopic effect in rare-earth doped oxide crystals at low temperature. The technique has the same advantages as other optical measurements: high bandwidth and no electrical contact on the sample, this latter representing a possible source of thermal and mechanical perturbations. The laser beam probes the stress induced by vibrations in the crystal instead of the change in distance between two reference points (as in, for example, interferometers). Piezospectroscopy has been used as a diagnostic tool for sapphire-based materials in which the inclusion of chromium can be used directly as a photoluminescent stress probe. This in-situ method is particularly well-adapted to extreme fabrication conditions, as in high temperature reactors. It was also proposed in diamond NV-centers as a tool to detect the damage trail left by a weakly interacting massive particles (WIMPs). Piezospectroscopy is even more relevant at low temperatures because atomic optical transitions become narrower, giving an enhanced sensitivity. This transition narrowing can be enhanced in rare-earth doped solids with the spectral-hole burning (SHB) mechanism, whose spectral resolution is in the MHz range well below the already existing limitations.
narrow inhomogeneous linewidth at cryogenic temperature\textsuperscript{23} (chap.4).

Here, we use the piezospectroscopic effect in rare-earth doped oxides as a diagnostic tool, but it was initially identified as a strong limitation to high resolution measurements based on SHB performed using cryocoolers. In recent years, interest has arisen in using SHB for optical signal processing applications, such as RF spectral analysis\textsuperscript{28} and acousto-optic filtering\textsuperscript{29}. The piezospectroscopic effect of rare-earth luminescent centers has been studied in the references\textsuperscript{30–34} and in Table I we give some representative values. Despite very different combinations of transitions and crystals, the piezospectroscopic coefficients (in Hz/\textmu Pa) have very similar values and are dispersed by less than an order of magnitude. This similarity is related to the structural resemblance of the crystals. The piezospectroscopic effect is an indirect Stark shift induced by a modification of the crystalline electric field surrounding the rare-earth dopant. The applied pressure translates into a compression (strain) which brings the ionic charges closer. The dopant transitions are then shifted due to their sensitivity to electric fields (the Stark effect). The linear Stark shift in rare earth 4f-4f transitions varies only by about two orders of magnitude across different ions and host materials\textsuperscript{35} (Table 2). The variation in Young’s modulus and inter-atomic distances is even smaller, resulting in comparable Stark coefficients across very different materials.

A wide range of dopants, transitions and host matrices has been studied in the references\textsuperscript{36,37} and in Table I we give some representative values. Despite very different combinations of transitions and crystals, the piezospectroscopic coefficients (in Hz/\textmu Pa) have very similar values and are dispersed by less than an order of magnitude. This similarity is related to the structural resemblance of the crystals. The piezospectroscopic effect is an indirect Stark shift induced by a modification of the crystalline electric field surrounding the rare-earth dopant. The applied pressure translates into a compression (strain) which brings the ionic charges closer. The dopant transitions are then shifted due to their sensitivity to electric fields (the Stark effect). The linear Stark shift in rare earth 4f-4f transitions varies only by about two orders of magnitude across different ions and host materials\textsuperscript{35} (Table 2). The variation in Young’s modulus and inter-atomic distances is even smaller, resulting in comparable Stark coefficients across very different materials.

Meanwhile, lanthanide inter-shell 4f-5d transitions have much stronger piezospectroscopic coefficients, as for example 4200 Hz/\textmu Pa for Pr\textsuperscript{3+} in YAG\textsuperscript{39}. The reason is that the 4f orbitals of the lanthanides are inner shells benefiting from a lower sensitivity to environmental changes, including the piezospectroscopic effect. In terms of measurement technique, the lower sensitivity of the intra-shell 4f-4f transitions is nevertheless largely compensated by the much narrower linewidth when probed by narrowband lasers. The resolution is, in that case, not limited by the inhomogeneous broadening (few GHz), but the homogeneous linewidth (below MHz) when the SHB technique is used, as we will see in section 11(B).

| Refs | Ion\textsuperscript{3+} | Transition | Crystal | Shift (Hz/\textmu Pa) |
|------|----------------|------------|---------|-------------------|
| 134  | (Table III) Pr | \(^{3}H_{4} \rightarrow \text{avg.}\) | LaOCl | -242.92 |
| 135  | (Table I) Nd  | \(^{4}F_{3/2}(1) \rightarrow ^{4}I_{9/2}(2)\) | YLiF\textsubscript{4} | -37 |
| 136  | (Table I) Pr  | \(^{3}H_{4}(1) \rightarrow ^{1}D_{2}(1)\) | YLiF\textsubscript{4} | -172 |
| 137  | (Table I) Nd  | \(^{4}F_{3/2}(1) \rightarrow ^{4}I_{9/2}(1)\) | YVO\textsubscript{4} | -193 |
| 138  | (fig.1) Nd    | \(^{4}F_{3/2} \rightarrow ^{4}I_{9/2}\) | LiNbO\textsubscript{3} | \(-270\) |
| 139  | (fig.1a) Pr   | \(^{3}H_{4} \rightarrow ^{1}D_{2}\) | YAG | \(-100\) |
| 140  | (fig.3) Eu    | \(^{1}F_{0} \rightarrow ^{1}D_{0}\) (site 1) | Y\textsubscript{2}SiO\textsubscript{5} | -211 |
| 141  | (fig.3) Eu    | \(^{1}F_{0} \rightarrow ^{1}D_{0}\) (site 2) | Y\textsubscript{2}SiO\textsubscript{5} | -52 |
| 142  | (fig.7a) Yb   | \(^{2}F_{7/2}(1) \rightarrow ^{2}F_{5/2}(1)\) | GGG | 76 |

TABLE I: Review of piezospectroscopic measurements in rare-earth doped oxides and fluorides (see Table I).
In summary, a typical value of 100 Hz/Pa can be kept in mind to evaluate the order of magnitude of pressure-induced shifts. As we will see now, the \( ^3H_6 \rightarrow ^3H_4 \) transition of Tm\(^{3+}:\text{YAG} \) at our wavelength of interest (793nm) falls in this range as well.

B. Piezospectroscopy of Tm\(^{3+}:\text{YAG}\)

The SHB technique has been extensively used to measure the homogeneous linewidth of many rare-earth transitions and has been described in many textbooks, see (chap.4) for example. Put simply, the technique involves using a narrow band laser to select out a spectrally narrow portion of a broader atomic line by pumping atoms into a non-resonant shelving state. The width of this artificially narrow line (the spectral hole) is determined by the linewidth of the laser, well below MHz for narrowband lasers (such as extended cavity diode lasers). For very narrow lasers, the resolution is eventually limited to a few kHz, the homogeneous broadening of rare-earth transitions such as those in Tm\(^{3+}:\text{YAG} \). The lifetime of the spectral hole can be extremely long, hours or days, by applying a field and shelving population in the Zeeman hyperfine structure. The SHB technique has been refined to the extreme in the context of optical metrology and used to accurately measure the detrimental shifts induced by the environmental changes, including pressure shifts.

In Fig. 1 we show the effect of a small applied pressure on the shape and position of a spectral hole on the \( ^3H_6 \rightarrow ^3H_4 \) transition of Tm\(^{3+}:\text{YAG} \). The initial hole (without pressure) is 50 kHz broad, significantly larger than the homogeneous linewidth (\( \sim 3 \)kHz) because of power broadening. This resolution is sufficient to observe the dramatic effect of even a few kPa.

![FIG. 1: Example of SHB absorption spectrum in Tm\(^{3+}:\text{YAG}\) (blue line). The hole has a roughly Lorentzian shape (black dashed line) of 50kHz width (FWHM). When a 3.3kPa pressure is applied along the [110] crystal axis, the line splits in two components (red line) that shift (and broaden) independently (see text for details).](image)

A splitting of the line is actually expected when a pressure is applied along the [110] crystal axis in \( \text{YAG} \). Tm substitutes for Y in the YAG cell at a site of D\(_2\) symmetry. This site has 6 possible orientations relative to the crystallographic axes, numbered from 1 to 6 (Ref. 43, Fig. 1 and references therein). Five of these sites contribute to the spectral hole in Fig. 1 site 1, and sites 3 to 6. Site 2 does not, because the optical transition is polarized, and for site 2 this polarization is parallel to the beam propagation direction, so the site is not excited. For pressure along [110], sites 3 to 6 are equivalent, and shift as a group in one direction, while site 1 shifts in the other, resulting in the split line in Fig. 1. Which peak corresponds to which site was determined by looking at the polarisation dependence of the split peaks.

The goal of our paper is not to describe extensively the piezospectroscopy effect in Tm\(^{3+}:\text{YAG} \), but rather to give a general idea of the link between the pressure vibrations and the optical transition. The measurement in Fig. 1 is illustrative and gives an order of magnitude of the pressure shifts. It was obtained in a dedicated variable temperature insert (VTI) cryostat (2-3K) that was used to apply a calibrated pressure and perform the piezospectroscopic analysis of our Tm\(^{3+}:\text{YAG} \) sample. The shift in Fig. 1 was measured as a function of the applied pressure. To vary the pressure, we used a piezoelectric actuator (Thorlabs AE0203D04F) that was inserted in the cryostat and constrained in contact with the crystal. To calibrate the piezo response, which is unknown at low temperature, we applied a static pressure by loading, with a calibrated weight, an inner rod that presses on the crystal through the VTI sample holder. A few volts across the piezo corresponds to 10 kPa (a few tens of grams on the crystal surface).

![FIG. 2: Pressure shifts of the different crystallographic sites when the pressure is applied along [110]. The laser polarisation is along [110] as well, and so addresses only sites 1 and 3 to 6. Site 1 causes the peak with large positive shift 68 Hz/Pa, and sites 3 to 6 the peak with a small negative shift \(-13\) Hz/Pa.](image)

We can see in Fig. 2 that the lines splits with different pressure coefficients for the different crystallographic sites. The splittings as a function of the piezo calibrated voltage are linear and allow us to extract a 68 Hz/Pa coefficient for site 1 and \(-13\) Hz/Pa to sites 3, 4, 5 and 6. More generally, we ob-
served shifts ranging from 13 Hz/Pa to 150 Hz/Pa depending on the sites and on the direction of applied pressure, demonstrating the anisotropic character of the piezospectroscopic response in Tm$^{3+}$:YAG. Nevertheless, a complete analysis of the piezospectroscopic tensor for the different sites is beyond the scope of this paper. In the following, we will keep the value of 68 Hz/Pa as a reference because it is somehow intermediate amongst the various measurements made. We first remark that Tm$^{3+}$:YAG has a similar shift to other rare earth materials (see Table I). We note that the line showed substantial broadening as well as splitting, indicating that not all ions in the crystal experienced the same pressure. This shows that it is difficult, even in a well-controlled measurement, to apply a uniaxial pressure that only shifts and does not broaden the line. We attribute the large broadening (with respect to the shift) to the inhomogeneity in the local strain field. Because of the surface roughness, the force may concentrate on few contact points. The spread from the surface to the interior of the crystal requires a specific mechanical analysis. We indeed discard any inhomogeneity of the local microscopic piezospectroscopic tensor (local distortion of the crystal cell) which could also lead to a broadening even under a uniform strain field. The local distortion of the crystal cell can be estimated by comparing the optical inhomogeneous line broadening (10 GHz~0.3 cm$^{-1}$) and the typical crystal field splitting (a few 100 cm$^{-1}$). So we expect the microscopic piezospectroscopic tensor to be homogeneous at less than the percent level. This estimation indicates by default that the broadening is due to the inhomogeneity of internal strain field and not the piezospectroscopic tensor. In addition, the presence of multiple site orientations means that an applied pressure would shift all the sites differently. These two features combined (intrinsic and multi-site broadenings) mean that the vibrations in a cryocooler are most likely to broaden the spectral line, with the amount of broadening as a measure of the strain in the crystal. Thus, we will use the measured piezospectroscopic coefficient $\kappa = 68$ Hz/Pa as representative of the line broadening in Tm$^{3+}$:YAG.

C. Piezospectroscopy and vibrations

Above we described the physical origin of the piezospectroscopic effect and illustrated the discussion with a static pressure measurement in Tm$^{3+}$:YAG. The extension of this work to the case of vibrations, as dynamical pressure fluctuations, is not direct and requires some modeling. We propose a toy model in which the sample is attached to a vertically vibrating plate (see fig. 3).

The piezospectroscopic effect is due to the compression of the crystal when pressure is applied. This latter is related to the strain by $\sigma = E \Delta L / L$ where $E$ is the Young’s modulus and $\Delta L / L$ the relative compression ($L$ is the sample length along $x$). In our model, the crystal is attached at one end where the vibration are induced as $x(t)$. For a unidirectional propagation of the vibration, the compression is due to the retarded propagation of sound in the material, so quantitatively, $\Delta L = \dot{x}(t) \cdot (t - L / V)$ where $V$ is the sound velocity in the solid. For acoustic waves below 1 MHz, the wavelength is much longer than the crystal so we can write $\Delta L = \dot{x}(t) / V$ or for the strain $\sigma = E \dot{x}(t) / V$ to the first order.

The relation between the strain induced in the sample and the atomic line shape is not trivial. As we discussed in section II B, a uniaxial pressure should in principle split the different sites. As we have seen, obtaining a pure shift without broadening is quite challenging in the static case because of the effective inhomogeneity of the strain field in the crystal along the axis of the laser. In the dynamical case, we expect the strain field to be even less homogeneous so the line is essentially broadened by the vibration, summing the effect of ions in different locations in the crystal and of each crystallographic site. The line broadening $\Gamma_\sigma = \kappa |\sigma|$ can then be related to the crystal velocity:

$$\Gamma_\sigma^{[\text{Hz}]}(t) = \kappa |E V | \dot{x}(t) | = 2.5 \times 10^9 |x[m/s]|(t)$$

where the numerical example corresponds to YAG with $E = 300$ GPa and $V = 8165$ m/s. The value of $\kappa = 68$ Hz/Pa is inferred from the spectral hole shift measurements reported in section II B.

The absolute value of the velocity in Eq. 1 makes it mathematically impossible to determine the signed velocity. An algorithmic retrieval of the signed signal is still possible but an algorithmic retrieval is still possible but is a complex signal processing task that we do not consider in a first approach. To proceed further with the calculation we are then compelled to crudely removing the absolute value, which comes down to assuming that negative strain leads to a narrowing of the spectral hole or that there is no negative strain at all. We expect this simplification to distort the sample displacement spectral density, with a possible doubling of the dominating frequencies, and the appearance of high frequency artefacts in the case of sharp features close to zero. The displacement spectral density will therefore be analyzed with caution.

The linewidth broadening induced by the vibration can be...
estimated optically by probing the absorption at the center of the hole as sketched in Fig. 3 (right) and as will be discussed in more details in Sec. V A. A line broadening (for example by a factor of 3 in Fig. 3, right) increases the absorption in the center of the spectral hole accordingly.

We can write the hole linewidth in terms of the measured absorption at the center of the spectral hole, and the original spectral hole width \( \Gamma_{HB} \) at the instant of the time-resolved measurement. \( \Gamma_{HB} \) is not obviously the hole width in absence of vibration but it can be the result of a complex hole burning dynamics. It will be measured experimentally. The vibration makes the hole width increase up to \( \Gamma_{HB} + \Gamma_{\sigma}(t) \), leading to a varying absorption coefficient \( 0 \leq \alpha(t) \leq \alpha_0 \) where \( \alpha_0 \) is the absorption in the absence of the spectral hole. Assuming a spectral hole broadening as previously discussed, and the conservation of the hole area, we write:

\[
[\alpha_0 - \alpha(t)](\Gamma_{HB} + \Gamma_{\sigma}(t)) = \alpha_0 \Gamma_{HB}
\]

leading to:

\[
\Gamma_{\sigma}(t) = \frac{\alpha(t)}{\alpha_0 - \alpha(t)} \Gamma_{HB}
\]

This simply reflects the conservation of the absorbing centers involved in the SHB process, which is valid as long as the hole is measured over times shorter than the hole decay time.

As a conclusion, by using a simple model, we can relate the vibration (Eq. 1) to the time-resolved SHB spectroscopy of \( \alpha(t) \) (Eq. 2) and finally extract quantitatively the characteristic vibration through its velocity \( \dot{x}(t) \).

III. MEASUREMENT SETUP

We used a pulse-tube cryocooler (TransMIT PTD-009) with an Oerlikon COOLPAK 2000A compressor. The rotary valve is rigidly attached to the cold head and has a cycle rate of 2 Hz. A sound level meter (RadioShack 33-099) was positioned in contact with the outer shield of the rotary valve and records the audio signal in the 1Hz–20kHz range.

A copper inner vacuum chamber is attached to the second stage at 2.9 K. It will be used for further isolation with exchange gas injection in the bottom of the chamber, thermally contacted by Apiezon-N grease. This should ensure a rigid mechanical contact at low temperature (below 236 K, the glass transition temperature Apiezon-M). The thermal contact is clearly maintained during the cooling cycle (absence of laser heating) so we can safely consider that the sample is rigidly contacted through the grease layer on cold stage.

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The crystal was fabricated by Scientific Materials and is anti-reflection-coated on both sides to reduce interferometric effects. The laser propagated along the [110] crystallographic axis. The thulium concentration in the sample is 0.25 at.% so that the total absorption is \( \alpha L \approx 2 \) for \( L = 5 \) mm. Further details about SHB in Tm\(^{3+}\):YAG including the population dynamics under optical excitation can be found in [16] (and references therein).

The narrow-band laser source (below 1kHz) was an extended cavity diode laser stabilized to a Fabry-Perot cavity via a Pound-Drever-Hall feedback loop and tuned close to the center of the Tm\(^{3+}\):YAG \( \text{3H}_6(1) \rightarrow \text{3H}_4(1) \) absorption line (12604.42 cm\(^{-1}\)). Diode lasers offer a moderate level of power fluctuations that could be interpreted as frequency shifts in our case. The use of a well-isolated reference locking cavity has the advantage to avoid the coupling of the cryocooler’s vibrations and laser cavity through air or optical table which would induce an acoustic frequency noise of the laser. This would be incorrectly interpreted as a piezooptroscopic shifts. The laser beam was spectrally and temporally shaped with an acusto-optic modulator (AA-Optoelectronics MT110) driven with an arbitrary waveform generator (Tektronix AWG5004). The 60 µW beam was focused into the sample with a 100 µm waist, corresponding to a Rabi angular frequency around 100 rad⋅s\(^{-1}\). We detected the transmitted light through the crystal with an avalanche photodiode (Thorlabs APD110A). The photodiode and the soundmeter signals were acquired via a digital oscilloscope (Agilent DSO5034A) as a 2 s long trace containing 4 million points per channel, leading to a 2 MHz sampling frequency.

In an experiment, the hole amplitude at line center can be directly measured by monitoring the absorption of a laser burning a continuous spectral hole. A continuous measurement like this allows us to measure up to very high frequencies, limited only by the timing resolution of the acquisition system. However, it is still necessary to measure \( \Gamma_{HB} \), the spectral hole width, by chirping the laser over the hole to measure the hole shape. In the continuous scheme, this reading chirp must be done periodically, at an interval of 31.2 ms in our case, comparable to the lifetime of the spectral hole, to provide an accurate reference.

The periodic pulse sequence depicted in Fig. 4 was repeated with a 31.2 ms cycle time, ie around 16 times per rotary valve cycle. The 30 ms burning pulse filled 96% of the cycle time and can therefore be regarded as almost continuous. The hole decay time was typically 10 ms (limited by the lifetime of the Tm metastable state \( \text{3F}_4 \)) so each burning-reading step can be considered as independent. The 100 µs reading pulse was linearly chirped over the burning frequency so as to provide...
We observe that the hole broadening oscillates at frequencies above 20 kHz and its value often exceeds 100 kHz in the first 0.3 s of each rotary valve cycle. The spectral hole perturbations are slightly delayed with respect to the rotary valve cycle. The acoustic noise and the piezospectroscopic perturbation have a common origin and are both triggered at the same time in the compression cycle. However, there is no direct relation between the two spectra, indicating that the vibration in the sample is not directly correlated to the vibration frequencies of the rotary valve itself.

B. Spectro-temporal analysis

We calculate the single-sided amplitude spectral density of the hole broadening for each sequence cycle:

\[
A_{r\sigma}(f) = \sqrt{\frac{2}{T}} |\tilde{\Gamma}_\sigma(f)|, \tag{4}
\]

where \(\tilde{\Gamma}_\sigma(f) = \int_{-T/2}^{T/2} \Gamma_\sigma(t)e^{2\pi ift} dt\) is the Fourier transform of the hole broadening \(\Gamma_\sigma(t)\) over the burning pulse duration \(T = 30\) ms. \(A_{r\sigma}(f)\) is expressed in Hz/√Hz. We calculate the hole broadening spectral density for each sequence cycle, and plot the result in the form of a spectrogram in Fig. 7(a), averaged over 150 acquisitions. It should be noted that the integration time \(T = 30\) ms is characteristic of the population dynamics (longest state lifetime). In other words, the populations are roughly stationary during this period. This ensures that the transmission linearly follows the frequency shift produced by the vibrations. A longer integration time could induce a population change that would give a non-linear response.
FIG. 7: Amplitude spectral density spectrograms of (a) the spectral hole broadening, (b) the corresponding displacement, and (c) the audio sound level on the rotary valve.

In Eq. 1 we gave the link between the hole broadening and the sample velocity. As explained in section II C we drop the absolute value in the equation. Displacement and velocity are linked by a derivation operation in the time-domain, which translates into \( \tilde{x}(f) = 2i\pi f \tilde{x}(f) \) in the spectral domain. We thus obtain the displacement spectral density:

\[
A_x(f) = \frac{V}{\kappa E 2\pi f} A_G(f)
\]

that we display in Fig. 7(b). Finally, Fig. 7(c) shows the spectrogram of the audio signal spectral density \( A_{V_a}(f) \) measured on the rotary valve.

The audio spectrogram clearly shows the rotary valve cycles at 2 Hz, with the high-pitched chirp (around 10 kHz) that can be heard twice per valve cycle, in agreement with previously reported cryocooler audio spectrograms.\(^5\) From the hole broadening and displacement spectrograms, we can define two uneven alternating steps in the rotary valve cycle that were already visible in Fig. 6: a noisier step, starting almost together with the loud rotary valve chirp and lasting 0.3 s, followed by a quieter step lasting 0.2 s. These two steps are not visible in the acoustic spectrogram, confirming that the vibration of the rotary valve is not transmitted to the cold finger in a straightforward manner. By comparing the characteristic frequencies in the spectrograms of the spectral hole broadening and the audio signal, we could not observe any correlation neither, even by anticipating the frequency-doubling (second order harmonics) due the crude removal of the absolute value in Eq. 1.

In Fig. 8, we plot the displacement spectral density as a multiplot with the same color code as in Fig 5. At 1 kHz, \( A_x(f) \) is equal to \( 2 \cdot 10^{-11} \text{ m}/\sqrt{\text{Hz}} \) in the noisy phase, and \( 2 \cdot 10^{-13} \text{ m}/\sqrt{\text{Hz}} \) in the quiet phase. This is two to three orders of magnitude lower than the values measured in other pulse-tube cryocoolers with no vibration isolation (between \( 10^{-8} \text{ m}/\sqrt{\text{Hz}} \) with optical detection\(^3\) and \( 5 \cdot 10^{-10} \text{ m}/\sqrt{\text{Hz}} \) with accelerometric detection\(^1\)).

C. Vibration spectra

This discrepancy could be explained by an inaccurate estimation of the displacement, or a lower vibration level of our low-power pulse-tube cooler compared to previously measured instruments. We discuss these in turn. There is an inaccuracy in the quantitative estimation of the vibration amplitude, because of the simplistic model used to relate the crystal absorption to the local strain, the somewhat empirical determination of the piezospectroscopic coefficient \( \kappa \) (by taking a typical averaged value of the measured shifts in II B), and the assumption of only positive strain.

It should be also noted that the originality of our method makes the comparison with other optical vibration measurements difficult. This is a general issue when comparing different cryocoolers characterized with different techniques. As an example, our pulse-tube has a low input power (2kW) as compared to other vibrations studies\(^3,15\) (typically 7kW). This makes direct comparisons generally challenging.

More fundamentally, the piezospectroscopic effect is sensitive to the local stress applied on the crystal attached to the second stage. Other optical methods intrinsically measure the relative position of the cold plate with respect to a reference point on the pulse-tube or the optical table base plane. There is
a priori no good or bad method. The appropriateness depends on the measurement of interest driven by the application. Different techniques should be seen as complementary to give a more complete characterization of the vibrations. In that sense, our measurements could be more quantitative if connected at lower frequencies to another type of measurement (ie with 3-axis accelerometers).

Due to its intrinsic sensitivity to velocity, our method’s sensitivity to displacement increases with the frequency. It operates between 100 Hz and 1 MHz. The lower value is imposed by the spectral hole lifetime (10 ms for Tm:YAG (ch.7)), whereas the upper value is a purely technical limit since it corresponds to the Nyquist frequency, ie 1 MHz.

V. CONCLUSION

In this paper, we presented a novel optical method based on the piezospectroscopic effect in a rare-earth ion-doped crystal for the detection of high frequency (> 100 Hz) vibration in a cold environment. Although not fully quantitative, this method is multidirectional and contact-less. Its sensitivity increases with frequency and has no fundamental upper frequency limit.

For our demonstration, we have used a cryocooler with optical windows. Nonetheless, further integration of the setup (primarily the crystal and the detector) is totally possible to form a compact single-component sensor with a fibered input feed-through (probe laser) and electric readout port (photodetector output). This integration step would allow to characterize a much wider range of cryocoolers without optical access. If the contact-less character of the method should be preserved, the fiber port can be attached to the cryocooler first stage and the crystal to the second stage (two-component sensor).

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Appendix A: Discarding the Doppler effect

In this paper, we have identified the piezospectroscopic effect as the principal source of coupling between vibrations and the optical absorption. As we discussed, rare-earth transitions are quite sensitive to strain because of their exceptionally narrow linewidth. This atomic-like feature for a solid impurity may give the impression that the Doppler shift induced by the vibrations could be a source of perturbation. We take the opportunity to show that the Doppler shift or broadening is much weaker than the piezospectroscopic effect.

The Doppler effect that would induce a similar broadening that the one discussed in II C. The latter is also proportional to the crystal velocity. It is given by

$$\Gamma_D^{(Ha)}(t) = \frac{1}{\lambda} |\vec{x}(t)| = 1.3 \times 10^6 |\vec{v}[m/s](t)|$$

where $\lambda = 793$ nm is the transition wavelength for our crystal. A direct comparison with Eq.(1) is then possible. The Doppler effect is three orders of magnitude weaker than the piezospectroscopic shift and can be fully neglected in first approach.

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