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Ultra-wide band near-infrared (NIR) optical thermometry (12–673 K) performance enhanced by Stark sublevel splitting in Er^{3+} ions near the first biological window in the $\text{PbZr}_{0.53}\text{Ti}_{0.47}\text{O}_3:\text{Er}^{3+}/\text{Yb}^{3+}$ phosphor†

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The fluorescence intensity ratio (FIR) approach, which relies on thermally coupled levels (TCLs), is significantly important for optical thermometry at room temperature and above, but was found to be impractical for low temperature sensing due to limited population density (thermal) or lack of spectrum at extremely low temperatures. Herein, we report a wide temperature range (12–673 K) sensing capability of the $\text{PbZr}_{0.53}\text{Ti}_{0.47}\text{O}_3:\text{Er}^{3+}/\text{Yb}^{3+}$ (C1:PZT) phosphor utilising the bandwidth of Stark sublevel split near-infrared (NIR) emission bands as one sensing parameter and FIR as another. Motivated by our previous studies on upconversion (UC) and the promising thermometry performance of the C1:PZT phosphor for real time nanothermometer monitoring (using visible TCLs), this work extends to the same thermometry application using UC-NIR emission as TCLs. The temperature-dependent UC spectra were measured across the ranges of 12–313 K and 313–673 K under 980 nm excitation, and their sensing capabilities were thoroughly evaluated. An enveloped single emission band, comprising multiple peaks in the NIR region, was observed and subsequently deconvoluted using Gaussian fitting. These individual peaks were analyzed in relation to Stark sublevel splitting, which was particularly evident at 12 K, and the variations in their full width at half maximum (FWHM) were compared across temperatures up to 313 K. Based on the temperature-dependent bandwidth, two prominent peaks $\sim 11655\text{ cm}^{-1}$ (858 nm) and 11454 cm^{-1} (873 nm), were identified as TCL levels, and a sensitivity (S_r) of $0.68 \pm 0.01\% \text{ K}^{-1}$ at 673 K was observed, making it a suitable thermometer for reading low temperatures using NIR bands.

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1. Introduction

The advancement of optical thermometry in the biophotonics window (including NIR-I & NIR-II) is practically limited in the region between 700 nm and below 1300 nm and has drawn considerable attention from researchers due to exceptional spatial resolution and deep tissue penetration capability.^{1,2} Potential phosphor materials such as organic dyes,³ quantum dots, semiconductors,³ and rare-earth (RE^{3+}) ion-doped nanocrystals^{4,5} have been extensively explored for various applications, including optical nanothermometry in the Near-Infrared I (NIR-I) region. Among these, NIR-I emitting RE ion-doped phosphors stand out

due to their excellent photostability, resistance to photobleaching, tunable emission profiles, and extended fluorescence decay times. These properties make them highly desirable for applications requiring durable and reliable luminescent probes. However, a key challenge with NIR-I emitting RE-doped phosphors is the temperature and doping concentration-dependent quenching of fluorescence intensity. This quenching is often linked to cross-relaxation (CR), inefficient energy transfer (ET) mechanisms, and other energy dissipation pathways inherent to RE^{3+} doping. In contrast, several studies have reported instances of suppressed NIR-II emissions^{6,7} in comparison to NIR-I emissions,⁸ with conclusions indicating that minimizing energy dissipation pathways can significantly enhance emission efficiency. Further studies have shown that coating of $\text{NaGdF}_4:\text{Nd}^{3+}$ @ NaGdF_4 nanocrystals⁹ with a passivating layer of NaGdF_4 of appropriate thickness can lead to notable improvements in NIR-II emission. Similar findings suggest that the application of various coatings on RE-doped core-shell structures can enhance overall NIR-II emissions by reducing surface quenching and improving energy transfer efficiency.^{10,11} These conclusive

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improvements in NIR-II emitting phosphors differ partially from those of NIR-I emitting phosphors. Furthermore, extensive studies^{8–12} have been conducted on UC luminescence emissions in the NIR-I (700–1000 nm) region compared to the NIR-II region, owing to its specific applications. Recently, Wang *et al.* reported excellent NIR-NIR temperature sensing *via* TCL variation of Tm^{3+} in a layered polarizing host.¹³ They utilized Tm^{3+} doped anisotropic $\text{Bi}_7\text{F}_{11}\text{O}_5$ nanosheets (layered polarizing host) to achieve wavelength selective and intense NIR-I (800 nm) UC luminescence. Typically, RE-doped TCL band emitters produce broad NIR-I emissions, which can be advantageous in some applications due to the broader spectral coverage. However, sharp NIR-I emissions, such as the 808 nm band, are particularly valuable for laser applications; for instance, the 808 nm emission from TCL can serve as an input wavelength for Nd:YAG lasers. In our previous study,¹⁴ we investigated UC emission and optical thermometry of $\text{Er}^{3+}/\text{Yb}^{3+}:\text{PZT}$ across a temperature range of 313–673 K, revealing interesting NIR emissions with TCL variation. The performance of this flexible thin-film thermometer, composed of $\text{Er}^{3+}/\text{Yb}^{3+}:\text{PZT}$ on a PDMS substrate, was validated through visible green and reddish emission bands, highlighting its potential for accurate temperature measurement in electronic devices. These findings, particularly the temperature-dependent NIR emissions, underscore the potential for further exploration of its sensing properties. Dong *et al.*¹⁵ presented related work, demonstrating NIR-NIR thermometry using Tm^{3+} -based thermometers that exploit Stark transitions of the $^3\text{H}_4$ sublevels. According to research by Lili Xing *et al.*,¹⁶ the host's crystal structure is correlated with spectral splitting, *i.e.* the degree of splitting is proportional to sensitivity (S_T) within 323–773 K. In the majority of publications on NIR thermometry insights on either Er^{3+} or Tm^{3+} ion doped phosphors, especially at room temperature, and in the low temperature range, the sensitivity is explained using Stark sublevels. The present work on the C1:PZT ($\text{Er}^{3+}/\text{Yb}^{3+}:\text{PZT}$) phosphor represents a novel approach, aiming to correlate Stark level splitting with NIR thermometry, while also enabling low-temperature measurements with Er^{3+} ions within a single phosphor sample.

As a continuation of our previous study, this work will examine the NIR-to-NIR optical thermometry performance of $\text{Er}^{3+}/\text{Yb}^{3+}:\text{PZT}$ phosphor (C1:PZT)¹⁴ samples. The choice of doping PZT with $\text{Er}^{3+}/\text{Yb}^{3+}$ ions, optimization and concentration quenching has been elaborated in our previous studies.^{14,17–19} In particular, in the morphotropic phase boundary region *i.e.* the $\text{PbZr}_{0.53}\text{Ti}_{0.47}\text{O}_3$ composition, the coexistence of both tetragonal (PT) and rhombohedral (PZ) crystal phases endows PZT with exceptional hosting capabilities and enhanced piezoelectric properties. These properties peak around the MPB due to abrupt changes in crystal symmetry, as discussed in our earlier works.^{14,18,19} This study employs two strategic approaches to derive its conclusions: (1) conducting absolute sensitivity analyses of (C1:PZT) samples¹⁴ across a broad temperature range (313–673 K) with NIR-I emission; (2) the overall performance of optical thermometry based on UC emission (NIR-I) in relation to TCL change is discussed on the basis of low perturbation crystal field introduction and Stark splitting. To the extent of our understanding, this report emphasizes the importance of explaining

closely spaced NIR bands, which were identified using a parametric model. The deconvoluted bands were then used for wide-range temperature sensing, with FWHM/FIR as the sensing parameter. The multiplet splitting of Er^{3+} ion energy levels into $(2J+1)/2$ Stark components due to small perturbations is analyzed based on FIR and sensitivity calculations. Because the variation in these NIR bands with temperature could not be defined within a single calibration, three distinct temperature regions were individually calibrated using the FIR to calibrate NIR emissions and address these limitations. This study follows a systematic approach, beginning with the identification and deconvolution of upconverted NIR emission bands, utilizing a parametric model and Gaussian distribution to account for Stark splitting within the Er^{3+} ion. Subsequently, absolute and relative sensitivities are calculated based on the FIR of Stark-split emission bands, specifically the $^4\text{S}_{3/2(\text{II})} \rightarrow ^4\text{I}_{13/2}$ transition at 875 nm ($i=4$) and the $^4\text{S}_{3/2(\text{II})} \rightarrow ^4\text{I}_{13/2}$ transition at 897 nm ($i=9$), across a wide temperature range. With this method, a sensitivity calibration curve may be predicted with NIR emitting upconverted bands. Moreover, these results also comply with absolute and relative sensitivity with excellent stability throughout the calibrating region.

2. Experimental

2.1 Synthesis

The optimized phosphor, designated as C1:PZT, was prepared using a low temperature assisted combustion route described in our previous work.¹⁴ The UC optimization studies were performed, and the concentration of $\text{Er}_x^{3+}/\text{Yb}_y^{3+}$ ratio *i.e.* ($x = 0.3$ mol% and $y = 3.0$ mol%) was found to be suitable for performing NIR-NIR temperature sensing studies. It should be noted that the concentration of Yb^{3+} ions was kept at 3.0 mol% due to the optimized concentration ratio (10:1) for efficient energy transfer, as already demonstrated in our previous studies.^{14,17,18,20,21} The final C1:PZT powder sample was first made into a dispersed solution in toluene and then coated using a spin-coating unit from POLOS (SPS POLOS-SPIN150-Standard), Germany, on an electronic printed circuit board (PCB) from Novastar, Alibaba Corp., China for measuring the temperature of the electronic counterpart. Detailed NIR temperature sensing studies for real time optical thermometry (thin film thermometer) were conducted, examining both absolute and relative sensitivity. For this purpose, the authors have prepared a solution by dispersing C1:PZT powder in approximately 20 mL of ethanol. The phosphor solution (SOL) was coated onto a p-type Si-wafer (100) using a spin-coating technique. After dispensing the solution onto the horizontal substrate, it was spun at 2500 rpm for 30 seconds to create a thin gel film, allowing excess liquid to evaporate quickly. The resulting film was then heat-treated at 900 °C for 1 hour in a muffle furnace. Finally, the prepared thin-film Si substrate underwent UC temperature sensing performance evaluation.

2.2 Characterization

The structural and optical characterization of the optimized C1:PZT sample, including X-ray powder diffraction (XRPD) with



Rietveld analysis, UV-vis spectroscopy, and UC measurements, has been detailed in our previous work.¹⁴ The room temperature UC emission spectra indicated that the NIR emission bands vary in a power-dependent manner, suggesting a correlation with thermally coupled level (TCL) bands. Consequently, NIR optical thermometry was performed using a different setup, as detailed below: temperature fluctuations were measured with a heating-cooling vacuum/gas stage (MHCS600, Microptik Instruments), which offers a temperature adjustment range of 100 to 870 K with 0.1 K stability. Additionally, a resistive temperature detection sensor and a K-type hybrid thermocouple sensor were employed for accurate temperature readings. Approximately 100 to 300 accumulations of UC emission spectra were recorded at each stage, once the probe reached thermodynamic equilibrium, in order to minimize the signal-to-noise ratio (SNR) ($\text{Signal/Noise} = 2^N$). The UC emission data were collected using a spectrograph (FLS F980, Edinburgh instruments), connected to a Hamamatsu R931 PMT and a InGaAs 1.7 DU490A high-resolution detection system. For rapid optical thermometry measurement, a fiber-coupled spectrometer from the Avantes (AvaSpec-650) low resolution detection system was used to measure the emission spectra.

3. Results and discussions

3.1 X-ray powder diffraction analysis

The powder XRD profile of the optimized C1:PZT ($\text{Er}^{3+}/\text{Yb}^{3+}$:PZT) sample confirms the formation of rhombohedral crystal symmetry with the ($R3m$) space group from profiles of PbZrTiO_3 (JCPDS #732022), as mentioned in our previous work.¹⁴ The Rietveld refinement was also conducted for the formed phases and crystal structural parameters mentioned in detail in our earlier report. For convenience, the powder XRD patterns for the $\text{PbZr}_{0.53}\text{Ti}_{0.47}\text{O}_3:\text{Er}^{3+}/\text{Yb}^{3+}$ (C1:PZT) samples are provided in the ESI,[†] as Fig. S1.

3.2 Discussion

The UC emission spectrum of the C1:PZT phosphor at 313 K with 980 nm excitation ($\sim 35 \text{ mW cm}^{-2}$) is shown in Fig. 1 and 2, respectively. It is evident that the phosphor displays a wide array of NIR emission bands consisting of different narrow intense peaks. Splitting of the $^4\text{S}_{3/2(\text{II})}$ level of Er^{3+} into seven Stark sublevels in the PZT crystal field has been established by the UC mechanism depicted in the inset of Fig. 1 using a low temperature (12 K) narrow spectrum.

We observed three primary emission bands, with centers at $\sim 11655 \text{ cm}^{-1}$ (858 nm), 11350 cm^{-1} (881 nm), and 11135 cm^{-1} (898 nm) as shown in Fig. 2. With temperature variations, the bands shift to 11494 cm^{-1} (870 nm), 11454 cm^{-1} (873 nm), 11428 cm^{-1} (875 nm), and 11123 cm^{-1} (898 nm) as shown in Fig. 1 with deconvolution using Gaussian fitting. These are the results of transitions from Stark sublevels $^4\text{S}_{3/2(\text{II})}$ to $^4\text{I}_{13/2(i)}$ ($i = 7, 6, 5, 4, 3, 2$, and 1). With an increase in temperature from room temperature up to 673 K, these transitions exhibit a well-defined upconversion (UC) luminescence variation,

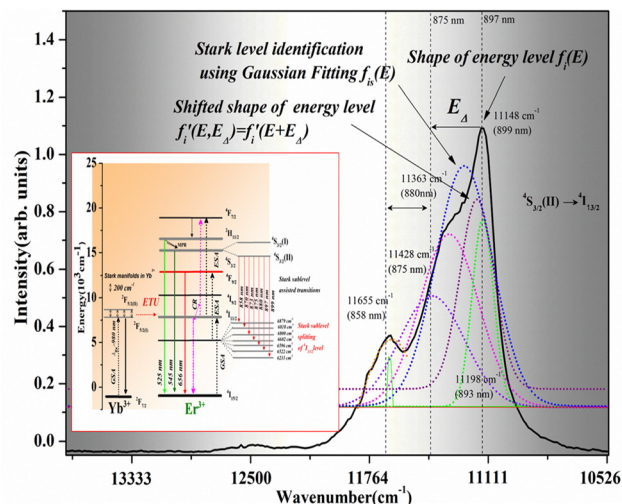


Fig. 1 UC emission spectrum of the C1:PZT phosphor at 313 K, with 980 nm excitation. The inset displays the possible UC mechanism responsible for the NIR emission process, for accommodating Stark sublevel splitting $^4\text{S}_{3/2(\text{II})}$ (j) ($j = 1, 2, 3, 4, 5, 6$ and 7) of the Er^{3+} ion.

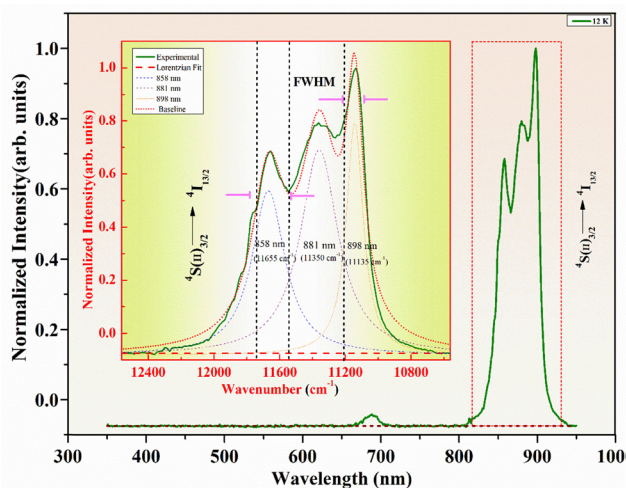


Fig. 2 UC emission spectra of the C1:PZT phosphor at 12 K, with 980 nm excitation. The zoomed red region displays the UC spectrum at 12 K within the range of 810–940 nm ($12345\text{--}10638 \text{ cm}^{-1}$).

resulting in notable FIR sensitivity across the thermally coupled luminescence (TCL) bands.

The C1:PZT phosphor's temperature sensing behaviour was investigated by measuring the UC spectra in the 12–300 K range, which (at 12, 70, 120, 220 and 313 K) are shown in Fig. 3. Throughout the studies, the excitation power was limited to reduce the photo-thermal effect. It is noteworthy that there is a considerable correlation between temperature and the intensity of luminescence, as well as the bandwidths. The increase in bandwidths can be attributed to the increase in electron-phonon interaction²² occurring in the temperature range of 12–313 K.

Interestingly, the Boltzmann relationship in FIR based optical thermometry is correlated with the temperature dependent

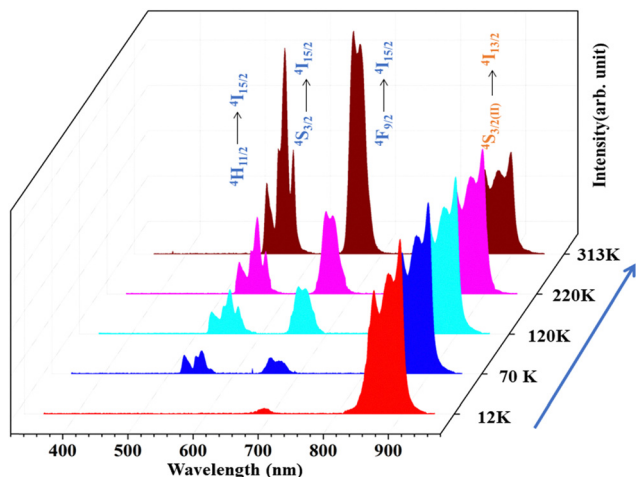


Fig. 3 UC emission spectra of the C1:PZT phosphor with 980 nm excitation (35 mW cm^{-2}) at different temperatures (12 K, 70 K, 120 K, 220 K and 313 K).

FWHM^{22,23} of the emission bands and can be expressed in a simplified equation,²³

$$\Delta\nu = \Delta\nu_0 + A \cdot e^{R_0 T} \quad (1)$$

here, A and R_0 are the fitting constants used where T indicates the absolute temperature and $\Delta\nu_0$ is the bandwidth. It is observed that among different Stark sublevel emission bands, these two bands at 11655 cm^{-1} (858 nm) and 11454 cm^{-1} (873 nm) are likely to be thermally coupled bands, which led us to show interest in their thermometry behaviour. Thus, the FWHM of the emission bands at 11655 cm^{-1} (858 nm) and 11454 cm^{-1} (873 nm) may be represented as a function of temperature,

$$\Delta\nu_{11454} = 17 + 2.9 \exp(1.05 \times 10^{-2} T) \quad (2)$$

$$\Delta\nu_{11655} = 11 + 5.1 \exp(7.15 \times 10^{-3} T) \quad (3)$$

The FWHMs of these two TCL emission bands, 11655 cm^{-1} (858 nm) and 11454 cm^{-1} (873 nm), are shown in Fig. 4(a) and (b).

Simultaneously, the FIR of these bands originated due to Stark splitting from very low temperature to room temperature (12–313 K), as shown in Fig. 4(c).

To detect an accurate temperature utilising the present sample in the NIR region, the UC spectra ($\sim \text{NIR}$ region) of C1:PZT with the strongest emission intensity at different temperatures within the range from 313 to 673 K were recorded and are shown in Fig. 5.

Three temperature regions were divided and the sensing behaviour was observed in the NIR region to be similar to our previous study,¹⁴ where sensing behaviour was studied in the visible region. The main reason for the distinctive three temperature region can be interpreted as UC emission peaks (three major transitions) showing either an incremental or decremental nature with temperature variation which is unique. Therefore, to calculate sensitivity in a wide temperature region, three temperature regions with distinctive incremental or decremental natures are reported. In practice, the UC based FIR technique utilizes the intensity ratio variation to measure sensitivity, which is advantageous and does not rely on the incremental/decremental nature. The FIR variation between $^4\text{S}_{3/2(\text{II})} \rightarrow ^4\text{I}_{13/2(\text{I})}$ (875 nm) and $^4\text{S}_{3/2(\text{II})} \rightarrow ^4\text{I}_{13/2(\text{I})}$ ($\lambda_{\text{em}} = 897 \text{ nm}$) i.e. Stark energy level transitions of the Er^{3+} ion established the basis of dual mode temperature sensing if we consider ($^2\text{H}_{11/2}/^4\text{S}_{3/2} \rightarrow ^4\text{I}_{15/2}$) transitions in the visible region as another TCL pair from the same sample. Moreover, the following transitions, $^4\text{S}_{3/2(\text{II})} \rightarrow ^4\text{I}_{13/2}(\text{Er}^{3+})$ and $^2\text{F}_{7/2}(\text{Yb}^{3+})$ achieve normalization at 899 nm and 979 nm, resulting in an inverse nature of UC emission with temperature in the $^4\text{S}_{3/2(\text{II})} \rightarrow ^4\text{I}_{13/2}(\text{Er}^{3+})$ transition. The energy level difference between TCLs of Er^{3+} i.e. $^4\text{S}_{3/2(\text{II})}$ and $^4\text{S}_{3/2(\text{I})}$ is around 250 cm^{-1} , which is under the suitable range ($200\text{--}2000 \text{ cm}^{-1}$) for excellent sensing using the TCL pair referred to. However, the $^4\text{S}_{3/2(\text{II})} \rightarrow ^4\text{I}_{13/2}$ transitions have greater probability of relative population gathering, which makes the transitions being referred to, 11655 cm^{-1} (858 nm) and 11454 cm^{-1} (873 nm) emissions, that originated due to Stark level splitting, a suitable TCL pair. The Boltzmann relationship governs the relationship between the relative population of

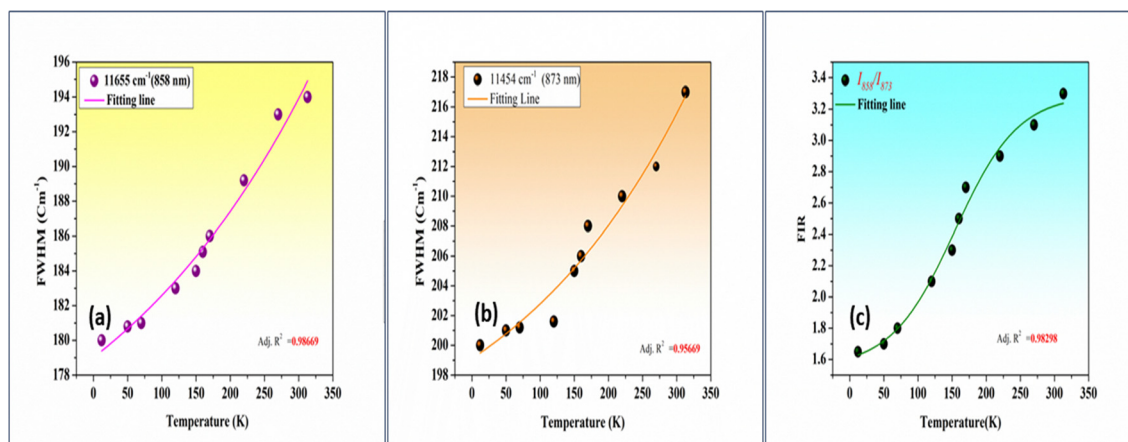


Fig. 4 The full width at half maximum (FWHM) of (a) 11655 cm^{-1} (858 nm) and (b) 11454 cm^{-1} (873 nm) bands as a function of temperature and (c) FIR of 11655 cm^{-1} (858 nm) and 11454 cm^{-1} (873 nm) emission bands as a function of temperature (cryogenic to room temperature).



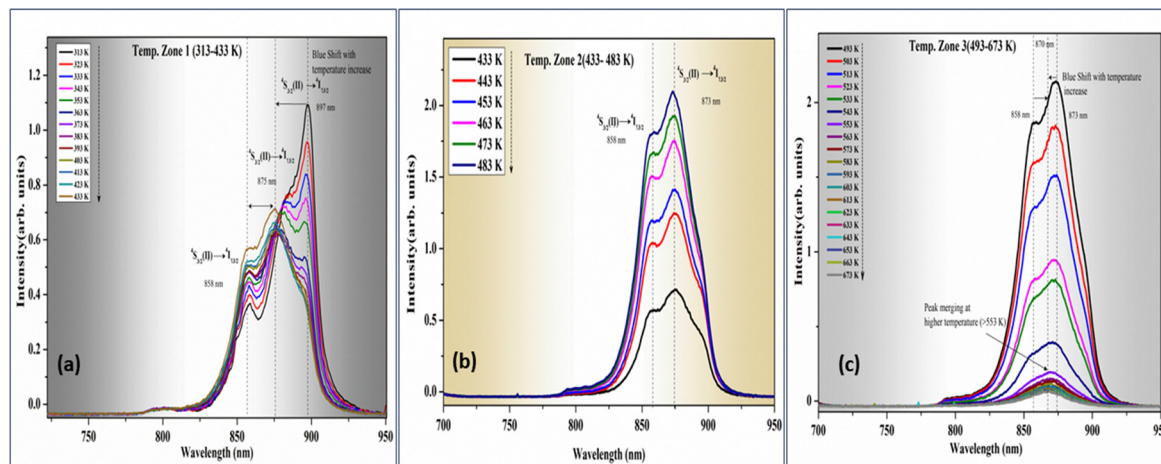


Fig. 5 Stark level splitting assisted UC emission (NIR region) of the C1:PZT phosphor sample in wide temperature regions (a) temp. zone 1 (313–433 K), (b) temp. zone 2 (443–483 K), and (c) temp. zone 3 (493–673 K), under 980 nm excitation.

TCLs and FIR, and can be described as,

$$\text{FIR} = \frac{I_j}{I_i} = \frac{g_j \sigma_{2j} \omega_{2i}}{g_i \sigma_{1i} \omega_{2i}} = C \exp\left(-\frac{\Delta E}{k_B T}\right) \quad (4)$$

where $C = \frac{g_j \sigma_{2j} \omega_{2i}}{g_i \sigma_{1i} \omega_{2i}}$ is the proportionality constant and ΔE is the energy difference between the TCLs; k_B ($0.6950 \text{ cm}^{-1} \text{ K}^{-1}$) is the Boltzmann constant, T is the absolute temperature, and g , σ , and ω are the degeneracy, emission cross-section, and angular frequency of the fluorescence transitions from the upper (j) to lower level (i), respectively. The aforementioned formula may be expressed as a linear equation as follows:

$$\ln(\text{FIR}) = -\frac{\Delta E}{k_B T} \left(\frac{1}{T}\right) + \ln C \quad (5)$$

$$\ln(\text{FIR}) = -B(1/T) + \ln C \quad (6)$$

where $B = \Delta E/k$. Fig. 6 illustrates the FIR variation with incremental temperature of the sample.

When $\ln(\text{FIR})$ is fitted linearly to $1/T$, the slope is $B = \Delta E/k$, and the intercept is $\ln(C)$. The slope and intercept, which represent the proportionality constant (C) and ΔE value for each of the three temperature zones shown in Fig. 6(a), (b), and (c), respectively, were derived by fitting the experimental data. Here, $C = \frac{g_j \sigma_{2j} \omega_{2i}}{g_i \sigma_{1i} \omega_{2i}}$ is the proportionality constant that depends on g , σ , and ω , which are the degeneracy, emission cross-section, and angular frequency of the two levels involved in the occurrence of transitions.

The fitting of temperature zones (1), (2), and (3) yielded values of C and ΔE . Accordingly, they are 1.1699 , $658.60 \pm 2.31 \text{ cm}^{-1}$, 0.679 ± 0.15 , $385.23 \pm 1.29 \text{ cm}^{-1}$, and 7.78 ± 0.29 and $-1.132 \pm 10 \text{ cm}^{-1}$.

It is now possible to determine the material's sensitivity by deriving from eqn (4). As we have previously reported, the relationship may generally be used to analyze two types of sensitivity: absolute and relative. However, when using it in a

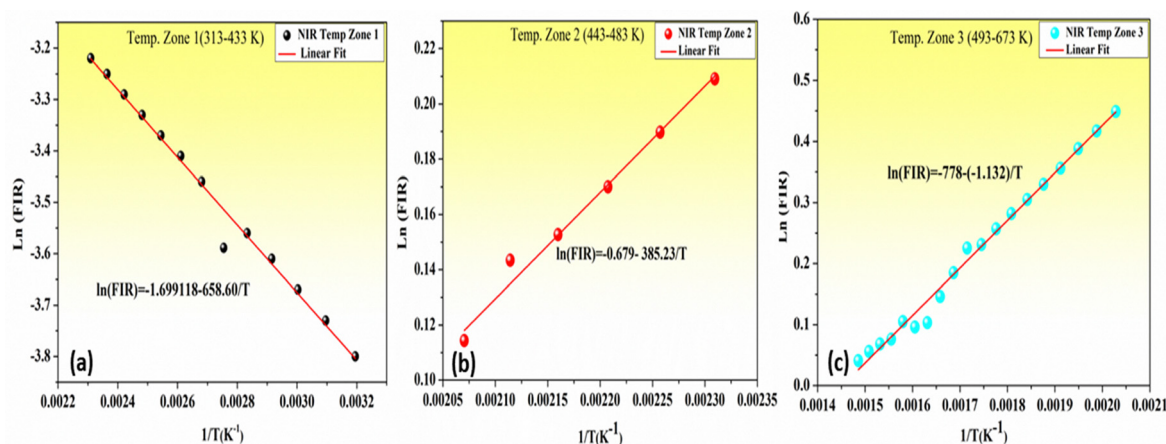


Fig. 6 Experimental and fitted FIR vs. temperature plots for the NIR emission in (a) temp. zone 1 (313–433 K), (b) temp. zone 2 (443–483 K), and (c) temp. zone 3 (493–673 K).



real-world thermometer for sensing purposes, the absolute sensitivity (S_a) is more crucial,

$$S_a = \frac{d(\text{FIR})}{dT} = \frac{\Delta E}{k_B T^2} C \exp\left(-\frac{\Delta E}{k_B T}\right) \quad (7)$$

However, the relative thermal sensitivity (S_r), is used to compare the performance of various thermometers and can be written in the formula as,

$$S_r = \frac{1}{\text{FIR}} \left| \frac{d(\text{FIR})}{dT} \right| \times 100\% = \frac{\Delta E}{k_B T^2} \times 100\% \quad (8)$$

where ΔE is in cm^{-1} , k_B is in cm^{-1} per K and T is in K. According to eqn (8), S_r is proportionally correlated with the energy difference between the TCL energy levels. However, energy levels that are energetically too far apart should be avoided, as chances of thermalization loss in this situation will be lower. Additionally, the upper thermalizing level depopulates and the emission intensity increases with the energy difference between the levels. It may not be feasible to measure very low intensity signals, which would not be economically viable as per the detection system. The Eu^{3+} ion as a thermal probe²⁴ has shown its potential as an optical nanothermometer with identical S_r values that can operate at various temperature ranges including 180 K (${}^7\text{F}_0 \rightarrow {}^7\text{F}_1$) i.e. in cryogenic, 300 K (${}^7\text{F}_0 \rightarrow {}^7\text{F}_2$) i.e. in physiological, and even in further temperature ranges using (${}^7\text{F}_0 \rightarrow {}^7\text{F}_3$) transitions because of its suitable energy difference among ${}^7\text{F}_j$ energy levels.

In our previous study¹⁴ we demonstrated that Er^{3+} ions can be used efficiently using the green (${}^2\text{H}_{11/2}/{}^4\text{S}_{3/2} \rightarrow {}^4\text{I}_{15/2}$) TCL emission bands for thermometry within an extensive temperature range (313–673 K). In the current study, we further explore the capability of Er^{3+} ions to operate at different temperature ranges, such as the cryogenic near 12 K (${}^4\text{S}_{3/2}(\text{II}) \rightarrow {}^4\text{I}_{13/2}$), the physiological near 313 K and beyond for temperatures above the physiological up to 700 K utilizing NIR upconverted emission due to Stark sublevel splitting (${}^4\text{S}_{3/2}(\text{II}) \rightarrow {}^4\text{I}_{13/2}$). Combining insights from these two studies, this level of tunability is highly advantageous for optical temperature probes, as it enables consistent relative sensitivity across a wide range of temperature regions. Practical demonstration in practical device fabrication as in a thin film thermometer has already been reported.¹⁴ The NIR emission bands of the Er^{3+} ion offer reliable strategic insight of the relative sensitivity parameter. We have evaluated both absolute (S_a) and relative (S_r) sensitivities of the TCL pair at $11\,655\text{ cm}^{-1}$ (858 nm) and $11\,454\text{ cm}^{-1}$ (873 nm) across the three temperature regions as shown in Fig. 7(a–f). The absolute sensitivity ($S_{a\text{max}}$) values observed were notably high across the defined temperature zones, with a maximum of 0.0144 K^{-1} at 313 K in temperature zone 1 (313–433 K), 0.00884 K^{-1} at 433 K in temperature zone 2 (433–484 K), and 0.045 K^{-1} at 493 K in temperature zone 3 (493–673 K). These values surpass those of other NIR-emitting TCL bands in similar luminescent centers. Additionally, the maximum relative sensitivity ($S_{r\text{max}}$) recorded was $0.68 \pm 0.06\%$ K^{-1} at 313 K in temperature zone 1,

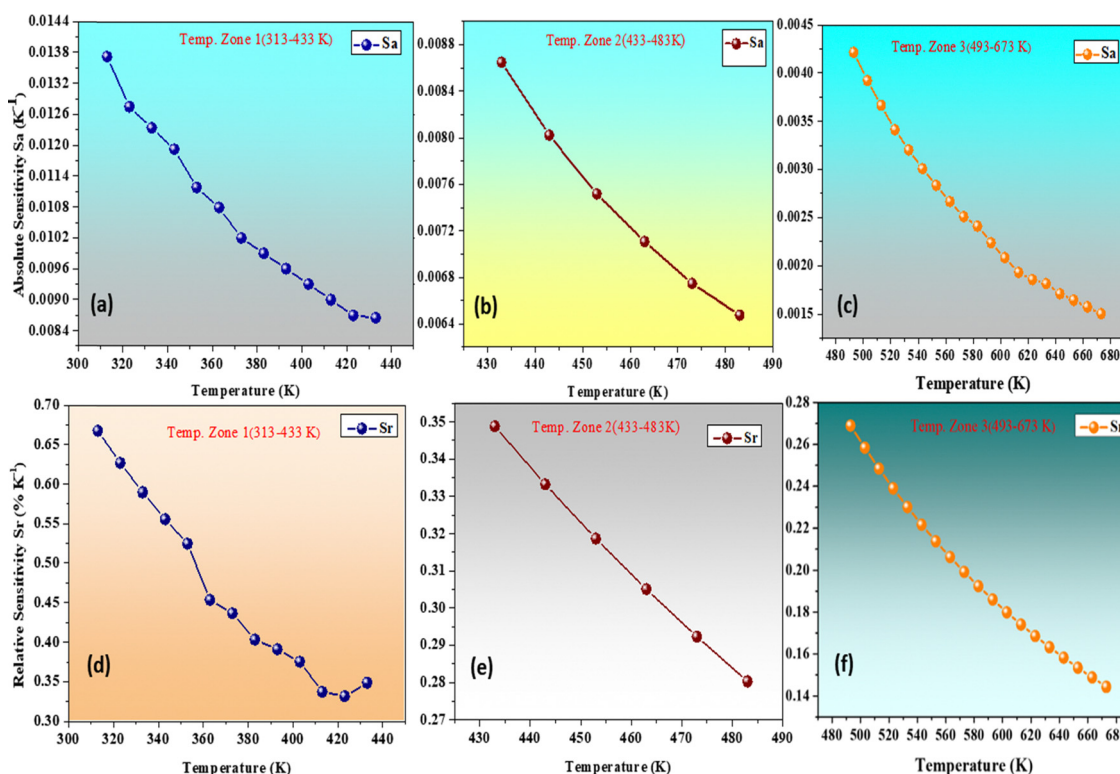


Fig. 7 The absolute sensitivities (S_a) of the TCL pair, $11\,655\text{ cm}^{-1}$ (858 nm) and $11\,454\text{ cm}^{-1}$ (873 nm) in the three temperature regions in: (a) temp. zone 1 (313–433 K), (b) temp. zone 2 (443–483 K), and (c) temp. zone 3 (493–673 K), and the relative sensitivities (S_r) in the three temperature regions in: (d) temp. zone 1, (e) temp. zone and (f) temp. zone 3, respectively.



Table 1 Relative temperature sensitivity of different nanothermometers

Host	λ_{em} (nm)	Temperature range (K)	S_r (Max) (% K ⁻¹)	Ref.
PbZrTiO ₃ :Er,Yb	858, 878	300–673	1.56	This work
α -NaYF ₄ :Yb/Tm@CaF ₂	802, 820	313–373	0.43	31
SrF ₂ :Gd,Nd	858, 866	293–338	0.61	32
YNbO ₄ :Nd	893, 930	303–473	0.28	33
Fluoroin-dogallate glass:Nd	820, 890	288–323	0.3	30
Y ₂ O ₃ :Er	800	293–353	2.02 \pm 0.06	26
Gd ₂ O ₃ :Nd	825, 890	288–323	1.75	34
NaYF ₄ :Er	837, 980	113–443	2.15	35

0.357 \pm 0.02% K⁻¹ at 433 K in temperature zone 2, and 0.286 \pm 0.09% K⁻¹ at 483 K in temperature zone 3. These S_r values are the highest reported for Er³⁺ ions within the context of NIR TCLs, highlighting the exceptional sensitivity achievable in this system. The PbZrTiO₃:Er³⁺/Yb³⁺ combination in the current C1:PZT phosphor is significant as a nanothermometer ($S_{r\text{max}}$ = 0.68% K⁻¹ at 313 K) among the many luminous thermometers based on Er³⁺/Yb³⁺ ion dopant hosts. Indeed, several papers^{12,25–30} exploiting Er³⁺ as a dopant exhibit larger sensitivities exploiting energy levels that are not thermally coupled and emission in the NIR region as well. Table 1 illustrates that the use of different dopants can yield varying sensitivity values. The table also indicates that among different dopants, the present work exhibits higher maximum sensitivities ($S_{r\text{max}}$) using NIR-I (800–980 nm) TCL bands, and also maintains stability over a wide temperature range. The significance of the present work lies in two main areas: first, although larger sensitivities have been reported with

Er³⁺/Yb³⁺ ion-doped hosts, optical thermometry performed using NIR bands, with detailed explanations of their origins from Stark sublevel splitting, is crucial and seldom discussed. Second, the Stark sublevel split NIR bands demonstrate effective optical thermometry across a broad temperature range, spanning both low (12–313 K) and high temperatures (313–673 K). However, these materials have excitation wavelengths that are not in line with the biological windows, limiting their applicability in *in vivo* measurements due to poor spectral resolution and image quality. Additionally, their thermal sensitivity is optimized for temperatures higher than those typically required for biomedical applications. Interestingly, the current phosphor could be employed in temperature sensing using more than one TCL pair, a salient feature uncommon among other luminescent centers. Moreover, the relative sensitivity of the NIR emitting TCL pair remains good at higher temperatures and maintains good thermal sensitivity from the low temperature region (12 K) to the higher order temperatures (673 K), exhibiting promising capabilities as a higher yield thermometry phosphor.

3.3 NIR temperature sensor

To demonstrate the practical application of PbZrTiO₃:Er³⁺/Yb³⁺ (C1:PZT), as an optical thermometer, the temperature-dependent NIR emission spectra were collected over the temperature range of 313–673 K, as shown in Fig. 8(a)–(c). A test of the temperature sensitivity of the C1:PZT sensor was conducted by measuring temperatures using a thin-film phosphor sensor (C1:PZT deposited on a Si-100 substrate) and an infrared thermal camera under 980 nm excitation. The initial

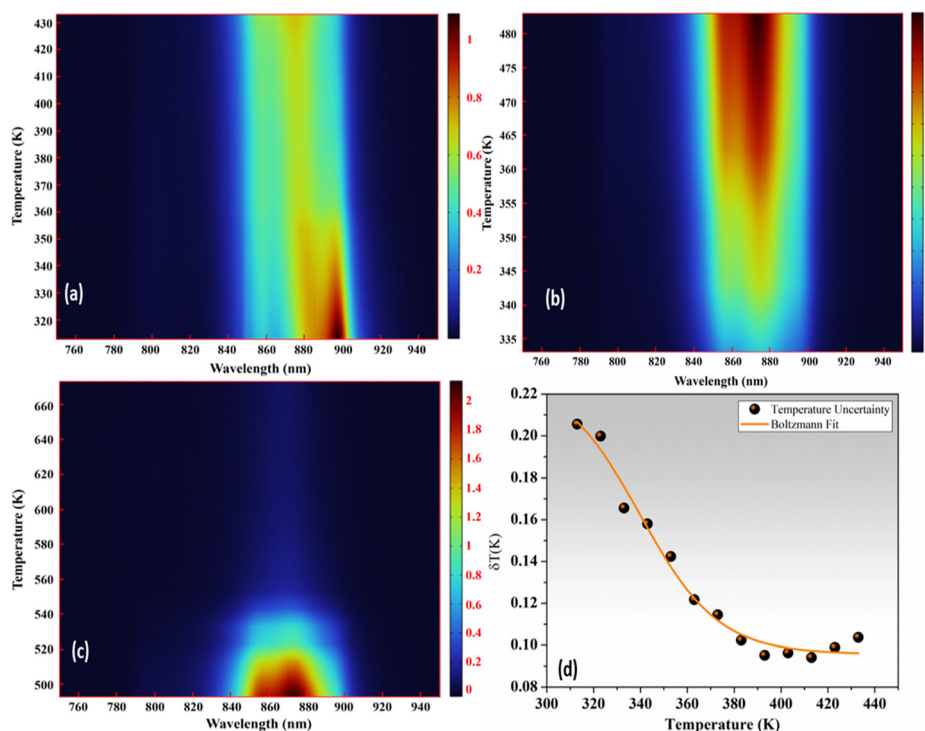


Fig. 8 The near-infrared UC spectra (temperature dependent) of the C1:PZT phosphor using NIR TCL emission bands in (a) temp. zone 1 (313–433 K), (b) temp. zone 2 (443–483 K), and (c) temp. zone 3 (493–673 K) and (d) temperature uncertainty of the C1:PZT sample while measuring relative sensitivity.



procedures for fabricating the thermometer device were followed as previously described.¹⁴ A NIR sensor and a thermal infrared imager, with varying irradiation times, were used to measure the temperature of electronic components on a printed circuit board (PCB) using FIR technology at 980 nm. The measurement process was as follows: first, the UC spectra of the as-prepared thin-film FIR thermometer were collected under continuous wave (CW) excitation with a 980 nm laser. The temperature of the electronic components was then calculated by correlating the FIR curves to the temperature values. Fig. 9(a) shows the temperatures of the electronic components, while Fig. 9(b) presents a comparison between the temperatures measured using an infrared thermal camera (x-axis) and those obtained using the FIR method (y-axis). The recorded temperatures from the NIR thermal imaging camera were found to be in close agreement with the FIR-based measurements, as the NIR thermal imaging captured the surface temperature of the samples, which closely reflects the actual temperature at the measurement location.

One of the other significant characteristics, expressed in terms of S_r values, is the temperature uncertainty (δT), which may be explained by,

$$\delta T = \frac{1}{S_r} \left| \frac{\delta \Delta}{\Delta} \right| \quad (9)$$

where $\delta \Delta$ could be calculated by error propagation theory.³⁶ Fig. 7 shows the temperature uncertainty of the TCL bands referred to. For these samples, the computed values of δT were 0.10–0.21 K (313–433 K) [refer to Fig. 8(d)]. Comparing this sample to other oxides previously reported in the literature, it demonstrated superior temperature uncertainty and sensitivity.³⁷

3.4 Correlation of Stark sublevel splitting in Er^{3+} ions and optical temperature sensing

In order to explain the correlation between NIR upconverted emission with optical thermometry within a broad temperature spectrum, we need to analyze the erbium ion energy diagrams

including Stark levels. Stark splitting significantly affects the shape and peak shifts of energy levels in many optical transitions within rare-earth ion matrices. Typically, the intra-configurational 4f–4f transitions in Er^{3+} ions are forbidden by the LaPorte rule due to parity conservation. However, the interaction of rare-earth ions with the crystal field or lattice vibrations can mix states of different parities into the 4f states, making these forbidden transitions observable. To explain the observed NIR emissions in the 850–900 nm range, along with their variable peak positions, we must consider the effect of small perturbations that shift and split the free ion energy levels. Additionally, these perturbed electronic states may interact in a variety of ways with the crystal lattice's phonon states. As a result, thermally coupled levels in Er^{3+} ions are observed. For instance, Huang *et al.*³⁸ previously reported the splitting of the $^4\text{I}_{15/2}$ level at $T = 12$ K, identifying eight distinct Stark components with a spread of 341 cm^{-1} . Identical results were also proposed by Bigot *et al.*³⁹ where the $^4\text{I}_{13/2} \rightarrow ^4\text{I}_{15/2}$ transition ($T = 1.5$ K) in the IR regime has a ground state spreading of 314.4 cm^{-1} with the simultaneous creation of several Stark splittings. However, limited reports are available on the near-infrared $^4\text{S}_{3/2} \rightarrow ^4\text{I}_{13/2}$ transition within the first biological window. To precisely determine the location and splitting of each Stark level within the $^4\text{S}_{3/2}$ multiplet of Er^{3+} , upconversion (UC) spectra can be utilized. The J -manifolds divide into $(2J + 1)/2$ Stark splitting in erbium ions in the presence of a crystal field, which results in two manifolds from the $^4\text{S}_{3/2}$ level and seven Stark split components⁴⁰ *i.e.* 6879, 6818, 6800, 6602, 6596, 6522 and 6233 cm^{-1} , through the $^4\text{I}_{13/2}$ level, respectively. Generally, this Stark sub-level splitting assisted transition in the Er^{3+} ion can be seen by excitations close to $1.5 \mu\text{m}$, also in 1.532 , 1.470 or $1.455 \mu\text{m}$. Considering a quasi-two-level laser process where a higher laser level represents a Stark sublevel the same as the pump level and the lower level belongs to the electrical ground state manifold. The low quantum defect leads to reduced heat deposition in the gain

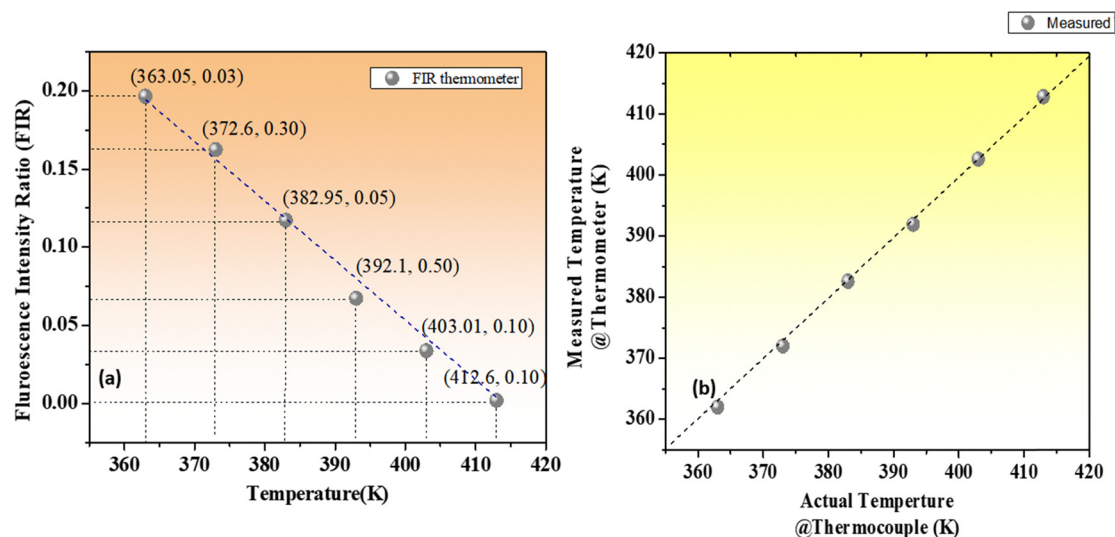


Fig. 9 (a) Temperature measurements via the calibrated FIR curves; (b) temperature of electronic components using FIR and an infrared thermal imager.



medium and enhances the laser process's high quantum efficiency. However, the UC mechanism, assisted by excited state absorption (ESA) and visible through the crystal's green emission, significantly influences the laser process. Consequently, maintaining an erbium concentration below 1 at% is essential to optimize the emission and minimize undesired effects. While the accuracy of the number and width of Stark level splitting depends on the crystal's symmetry, the intensity is influenced by variations in the crystal field,⁴¹ as noted in previous studies. Due to these variations, differing crystal fields across ions result in distinct Stark splitting, thereby broadening the observed width of Stark levels when measured across an ion ensemble. However, in this case, the broadening and shifting of emission peaks with temperature correspond to various transition energies, which can be described using the following nomenclatures.

1 Parametric model for the shape of the energy levels based on Stark level splitting. A parametric model⁴² characterising the shape of Stark levels should take into account that the shape dictates the dominance of Stark levels in Er^{3+} energy levels. The parameterization involves assigning a Gaussian distribution to every Stark level S_i .

$$f_{is}(E) = x_{is,\text{scale}} + \frac{1}{\sqrt{2\pi}x_{is,\text{width}}} \exp\left(-\frac{1(E - E_{i,\text{center}} + x_{is,\text{splitting}})^2}{2x_{is,\text{width}}^2}\right) \quad (10)$$

here, the scaling factor ($x_{is,\text{scale}}$) determines the relative height of the width ($x_{is,\text{width}}$) and $x_{is,\text{splitting}}$ represents the divergence of the Stark sublevel from the center ($E_{i,\text{center}}$) of the discrete level i , ($^{2S+1}L_J$) in the absence of Stark splitting. The parameters $x_{is,\text{scale}}$, $x_{is,\text{width}}$ and $x_{is,\text{splitting}}$ are the free parameters of the model. The shape of each level (i) is determined by calculating the sum over the Stark levels (s_i):

$$f_i(E) = \sum_s f_{is}(E) \quad (11)$$

Fig. 1 displays the shape of wide bands consisting of seven Stark level transitions including temperature dependent peaks, respectively.

2 The shape of the energy levels and line shape function. The line shape function $g_{ij}(E_\Delta)$ of a transition between the energy levels i and j may be computed using two provided shapes $f_i(E)$ and $f_j(E)$ of the energy levels i and j ,

$$g_{ij}(E_\Delta) = \frac{\int f_i(E)f_j(E + E_\Delta)dE}{\int f_i(E)f_j(E + E_\Delta)dE dE_\Delta} \quad (12)$$

The integral of the denominator in eqn (12) gives the normalization which provides validation by

$$\int g_{ij}(E_\Delta)dE_\Delta = 1 \quad (13)$$

This provides an understanding of first order shifting of line shape function of the highest energy level $f_i(E)$ i.e. $11\,454\text{ cm}^{-1}$ (873 nm) by E_Δ ($\sim 199\text{ cm}^{-1}$), and simultaneously finding the line shape of the lower level i.e. $11\,655\text{ cm}^{-1}$ (858 nm) by calculating the area underneath as a product of two. This is indicative of an evaluation of the intensity of the transition

from the higher to the lower energy level, with the precise transition energy E_Δ , as shown in Fig. 1. It is possible to determine the probability of the transition between certain energy levels by doing a subsequent integration across E_Δ . Fig. 1 shows the peaks of the energy level can be well fitted using a Gaussian distribution and their FWHM is calculated which varies with temperature as plotted in Fig. 4.

3 The Stark sublevel parameters. It is possible to determine the energy level's shape by a reverse optimisation method which uses a model described by Steinkemper *et al.*⁴² and Krammers theorem, and the maximum permissible limit for expanding the FWHM as well as the Stark level splitting can be found under $< 200\text{ cm}^{-1}$ which for the current case is shown in Fig. 5.

All of the described methods were considered in the present case and it was concluded that the Stark sublevel splitting is possible certainly in low temperature measurements and even in the room temperature to higher temperature range. Plausible Stark sublevel splitting of $^4S_{3/2}$ and $^4I_{13/2}$ could be observed and hence can show thermometry performance using those transitions referred to (inset of Fig. 1). The decreasing trend of fluorescence of Stark level transitions with increasing temperature from 12 K to 313 K is caused by the phonon assisted excitation between $^4S_{3/2}(\text{I})$ and $^4S_{3/2}(\text{II})$. The fluorescence intensity of the NIR emission bands is thermally quenched as a result of the Er^{3+} ions in the Stark level $^4S_{3/2}(\text{I})$ being instantly thermally excited to the adjacent sublevel $^4S_{3/2}(\text{II})$ with rising temperature due to an energy gap of 50 cm^{-1} (Fig. 3).

3.5 Probability of Stark sublevel splitting in Yb^{3+} ions and anti-Stokes population inversion in Yb^{3+} manifolds

The aforementioned section has established a clear justification for the Stark level splitting among the Er^{3+} ions which leads to the upconverted NIR emissions based on $\text{Yb}^{3+} \rightarrow \text{Er}^{3+}$ ion ET mechanisms. However, it is necessary to further explore the behavior of Yb^{3+} ions at low temperatures and consider revising the energy transfer model to account for potential anti-Stokes population inversion in the Yb^{3+} ion, as well as map this effect accordingly. The anti-Stokes fluorescence process^{43,44} is a phenomenon where a material absorbs photons of lower energy and subsequently emits photons of higher energy, resulting in a net loss of thermal energy from the material. Ytterbium (Yb^{3+}) ions are particularly well-suited for laser-induced fluorescence cooling because they exhibit minimal multiphonon relaxation, concentration quenching, and excited-state absorption. Ytterbium (Yb^{3+}) has two manifolds, the $^2F_{7/2}$ ground state and the $^2F_{5/2}$ excited state, separated by approximately $\sim 10^4\text{ cm}^{-1}$, with moderate electron-phonon coupling. In a glass (ZBLAN & BIG)⁴⁴ host, Stark sub-level splitting occurs approximately by $\sim 200\text{ cm}^{-1}$, enabling thermalization within a manifold when population distributions are perturbed. To investigate and map the possibility of anti-Stokes population inversion in the current sample, the authors synthesized an $\text{Yb}^{3+}:\text{PbZrTiO}_3$ phosphor without Er^{3+} ions using the same experimental procedure outlined in this work.¹⁴ The Yb^{3+} ion concentration was maintained at 3.0 mol%, consistent with previous experiments.



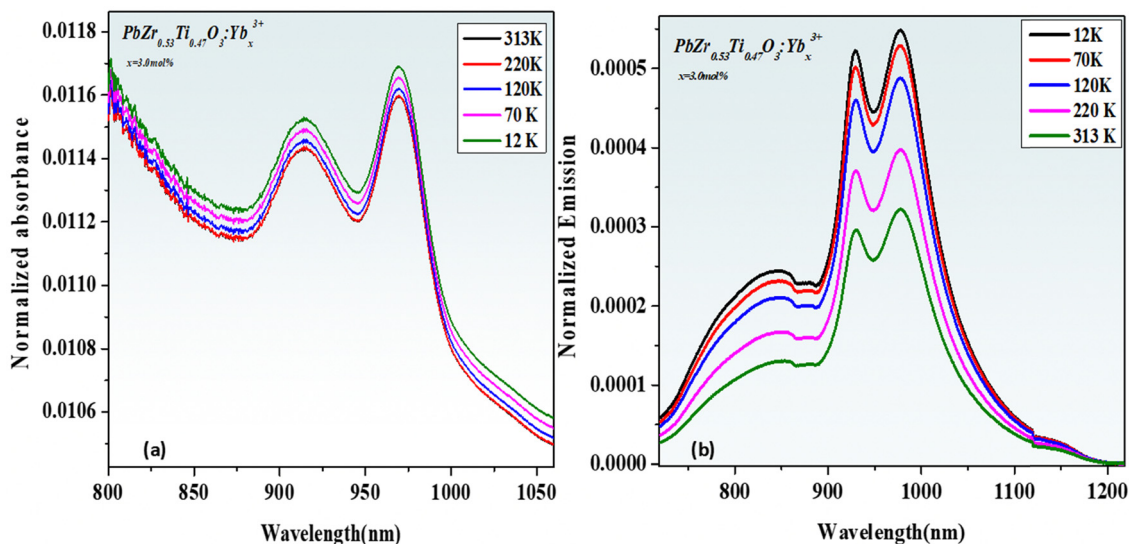


Fig. 10 Normalized (a) absorption and (b) emission spectra of the Yb^{3+} :PZT host at different temperatures.

The absorption and emission spectra were recorded from 12–313 K to evaluate the theoretical minimum operating temperatures and cooling efficiencies under optimal conditions as described in the reports.^{43,44} Fig. 10 illustrates the temperature-dependent absorption and emission spectra for the Yb^{3+} : PbZrTiO_3 phosphor sample. The long-wavelength absorption tail, which is indicative of cooling potential, shifts with temperature as the absorption coefficient decreases at lower temperatures. This observation supports the idea that Yb^{3+} ions doped in the current host material may undergo sub-Stark anti-Stokes transitions, as predicted by Murtagh *et al.*⁴⁴ However, the emission and absorption spectra of the Er^{3+} ion doped C1:PZT phosphor sample do not exhibit the broad 976 nm emission, as shown in Fig. 10, when measured using a similar pump-probe spectrophotometer⁴³ system. Although the authors verified the possibility of Yb^{3+} manifold-assisted Stark transitions, the energy transfer (ET) process from Yb^{3+} to Er^{3+} is more likely in the current system. This conclusion is consistent with previous findings on energy bridging mechanisms²⁰ in Er^{3+} -doped phosphors, modelled using rate equations. The emission observations on Yb^{3+} : PbZrTiO_3 phosphor further supports the Yb^{3+} Stark sub level split manifolds at lower temperatures as reported by Mungan *et al.*⁴³ However, in the present work, the observed NIR emission bands around 800–900 nm occur within a broad emission envelope. These emissions are more accurately explained by Stark sublevel splitting of Er^{3+} ions rather than Yb^{3+} ions, which typically emit in the 900–1000 nm range.

4. Conclusion

In summary, optical temperature sensing and real time temperature monitoring were investigated using NIR upconverted emissions from C1:PZT samples, which have shown excellent thermometry using visible upconversion emission in our previous work. The current work on optical thermometry completes the extension of the

Er^{3+} ion optical thermometry performance using visible to NIR upconverted emission bands as TCLs. Detailed analysis of UC emissions under 980 nm excitation across temperatures from 12 K to over 300 K reveals that these NIR emissions arise from Stark sublevel transitions. The observed single NIR emission band, comprising multiple peaks, was deconvoluted using Gaussian fitting and shows temperature-dependent changes from 12 K to 313 K. The FWHM of the emission bands centred at $11\,655\text{ cm}^{-1}$ (858 nm) and $11\,454\text{ cm}^{-1}$ (873 nm) was observed to increase with rising temperature, suggesting its potential as a reliable sensing parameter. By utilizing temperature-dependent FWHM modulation, a peak relative sensitivity ($S_{r,\text{max}}$) around $1.56 \pm 0.09\% \text{ K}^{-1}$ at 673 K was achieved at 673 K. This sensitivity represents the highest reported value for Er^{3+} ions in the context of NIR-TCLs. Moreover, these higher values of sensitivity in a wide temperature range (12–683 K) using NIR upconverted TCL bands were explained and correlated on the backdrops of Stark sublevel splitting. A parametric model nomenclature was demonstrated to confirm the splitting of the $^4\text{I}_{13/2}$ level of the Er^{3+} ion and to determine the line shape function using boundary conditions, shown in the work. These findings indicate that this material demonstrates significant potential for a range of applications, particularly in low-temperature sensing utilizing NIR emissions. The results further indicate that these NIR emissions possess strong sensing capabilities with excellent repeatability for temperature detection in microelectronic devices, as maintaining a sensitivity of $1.56 \pm 0.09\% \text{ K}^{-1}$ at 673 K presents challenges in precision. Additionally, the relative sensitivity S_r is nearly $1.0\% \text{ K}^{-1}$ within the biologically relevant temperature range of 270 K to 320 K, highlighting its potential for biomedical applications, particularly in photo-thermal treatments.

Author contributions

The original idea, design, conceptualization of the work, experiments, data analysis and interpretation and drafting of the



article were carried out by PPS. Data analysis of the article was carried out by YS. Critical review, scientific suggestions and editing were carried out by HCS.

Data availability

Data will be made available on request.

Conflicts of interest

The authors have no conflicts of interest to declare. All co-authors have seen and agree with the contents of the manuscript and there is no financial interest to report. We certify that the submission is original work and is not under review at any other publication.

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