The ability to tune and enhance the properties of luminescent materials is essential for enlarging their application potential. Recently, the modulation of the photoluminescence emission of lanthanide-doped ferroelectric perovskites by applying an electric field has been reported. Herein, we show that the ferroelectric order and, more generally the polar order, has a direct effect on the photoluminescence of Eu$^{3+}$ in the model BaZr$_x$Ti$_{1-x}$O$_3$ perovskite even in the absence of an external field. The dipole arrangement evolves with increasing $x$ from long-range ferroelectric order to short-range order typical of relaxors until the non-polar paraelectric BaZro$_3$ is achieved. The cooperative polar interactions existing in the lattice ($x < 1$) promote the off-center displacement of the Eu$^{3+}$ ion determining a change of the lanthanide site symmetry and, consequently, an abrupt variation of the photoluminescence emission with temperature. Each type of polar order is characterized by a distinct photoluminescence behaviour.
The BaZrTi1−xO3 (BZT) perovskite solid solution has been selected as a model host because the polar order can be tuned by changing x. The end-member BaTiO3 (x = 0) is one of the most carefully investigated ferroelectric compounds and, thanks to its high dielectric constant and low losses, is widely used as a ceramic material for the fabrication of multilayer ceramic capacitors in the microelectronic industry and other applications.10,17 BaTiO3 exists in different crystallographic variants, depending on temperature. At the Curie temperature, TC = 125–130 °C, it turns from the high temperature paraelectric cubic form (C, Pm-3m) to the ferroelectric tetragonal structure (P4mm). On further cooling, two other ferroelectric phases are observed: orthorhombic (O, Amm2), below 5–15 °C, and rhombohedral (R, R3m), below ~80 – 90 °C. All polar forms originate from a small deformation of the prototype cubic lattice arising from the off-center displacement of the Ti ion (about 0.1 Å at room temperature) in the TiO6 octahedron. The cooperative, long-range ordering of the resulting electrical dipoles gives rise to macroscopic spontaneous polarization, spontaneous lattice strain and ferroelectricity.24–26 Differently from BaTiO3, the crystal structure of the end-member BaZrO3 (x = 1) corresponds to the prototype Pm-3m cubic perovskite up to the melting point.26,27 Barium zirconate is a nonpolar, paraelectric solid which has attracted attention in recent years as an efficient proton conductor when doped with acceptor impurities.22,23 BaTiO3 and BaZrO3 show complete solid solubility. In the BZT solid solution the extent of polar order decreases with increasing x.24–28 As x increases, the R/O transition (TC = 50 °C) and, consequently, the small deviation of the polar structure from the cubic structure (TC = 125–130 °C) to the ferroelectric tetragonal (TC = 125–130 °C) to the ferroelectric tetragonal structure (P4mm) at about ~10 °C. A single and broad permittivity peak at 50 °C, corresponding to a diffuse ferroelectric to paraelectric transition. Relaxor or dipolar glass state with only short-range polar order is observed for higher values of x, up to 0.95, and finally weak paraelectric behavior dominates in BaZrO3. The relaxor state is characterized by the presence of polar nanoregions (PNRs) embedded in a paraelectric matrix and an average cubic structure.27,28 According to recent investigations, the PNRs in BZT essentially correspond to frozen BaTiO3 clusters with Ti off-centering.

Eu3+ was chosen as a dopant because of the well-known bright emissions and for its powerful ability to act as a spectroscopic local probe for the determination of the symmetry of the first coordination sphere of a lanthanide ion in organometallic molecules and crystals.29 Furthermore, Eu3+ has an ionic radius (c.n. 12) of 1.226 Å and, therefore, its crystalchemical behavior can be considered as representative of other trivalent lanthanide ions with intermediate size, from Sm3+ to Er3+.

Six Eu-doped Ba(Ti, Zr)O3 ceramics with composition Ba1−yEuyTi1−x−y/4ZrxO3 (y = 0.01; x = 0, 0.05, 0.15, 0.30, 0.50, 0.70) and Ba1−yEuyZr1−y/4O3 (y = 0.01) were investigated. The different samples are labelled as BZX, where x = 100x.

Results and Discussion

Dielectric properties and polar order. The dielectric constant (real part of permittivity) of the BZ0-BZ30 ceramics is reported in Fig. 1. The phase transitions are indicated by anomalies of the dielectric constant. When x = 0 (Fig. 1a), there are three peaks at ~70, 15 and 125 °C corresponding to R/O, O/T and T/C transitions, respectively.

The sharp permittivity peak at TC is typical of conventional ferroelectrics with first-order ferro/para transition. For ceramic BZ5 (Fig. 1b), TC decreases to 108 °C and the temperature of the O/T transition increases to 50 °C. The R/O transition is no longer detected in Fig. 1b but can still be identified from the anomaly of the dielectric loss (the ratio of the imaginary and real part of dielectric permittivity, see Figure S1, Supporting Information) at about ~10 °C. A single and broad permittivity peak at 50 °C, corresponding to a diffuse R/C phase transition, is observed in Fig. 1c when x = 0.15. Accordingly, the dielectric loss also reveals a single transition (Figure S1, Supporting Information). The sample BZ30 (Fig. 1d) displays a typical relaxor behavior: the temperature corresponding to the maximum permittivity shifts at higher temperatures with increasing frequency. The permittivity maximum no longer corresponds to a phase transition but is determined by the relaxation dynamics of the PNRs.33 Comparison with literature data indicates that the dielectric properties of the Eu-doped samples are rather similar to those reported for BZT materials likely due to the low concentration of the dopant. The phase transition temperatures of ferroelectric samples are very close (within a few degrees) to those of the undoped ceramics.

The polarization - electric field (E) loops reported in Fig. 1e confirm the results of dielectric measurements. BZ0, BZ5 and BZ15 show typical hysteretic ferroelectric behavior. Both the maximum polarization and the remnant polarization (intercept with the vertical axis) decrease with increasing x as expected from the progressive lowering of TC. BZ30 sample shows a non-linear behavior without hysteresis, a feature representative of relaxors.

Crystal structure. The splitting of the (200) peak in the XRD pattern of the BZ0 ceramic (Figure S2, Supporting Information) indicates a typical P4/mmm tetragonal structure and the refinement gives a c/a ratio of the lattice parameters of 1.0096, only slightly smaller than that of undoped BaTiO3 (1.0109). The unit cell volume, 64.35 Å3, is virtually the same of undoped barium titanate (64.34 Å3) being the difference within the experimental error. Although peak splitting is no longer observed, the broadening of some lines (compare the 111 and 200 peaks) indicates a non-cubic structure for BZ5 (Figure S2, Supporting Information), in agreement with the existence of long-range ferroelectric order at room temperature. The assignment of the crystal symmetry is not straightforward given the absence of splitting. Refinement with P4/mmm tetragonal structure with a lower c/a ratio of 1.0041 results in a marginally better fit than orthorhombic Pmmm structure. XRD pattern of BZ15 indicates a cubic Pm-3m symmetry (Figure S2) though this composition is expected to be ferroelectric at room temperature as indicated by the dielectric measurements (Fig. 1c). This apparent discrepancy is determined by the proximity of the phase transition (TC = 50 °C) and, consequently, the small deviation of the polar structure from the cubic arrangement not detectable with a conventional diffractometer. The ceramics with x = 0.30, 0.50 and 0.70 show the cubic Pm-3m structure, as expected from their relaxor nature. Relaxors possess an average cubic structure but their local symmetry is lower and characterized by the existence of PNRs with short-range polar order. The
unit cell volume increases linearly with $x$ (Figure S2, Supporting Information) as a result of the larger ionic radius of $\text{Zr}^{4+}$ (0.720 Å, c.n. 6) in comparison to $\text{Ti}^{4+}$ (0.605 Å, c.n. 6). The unit cell volume of the ceramic with $x = 1$ is 73.71 Å$^3$ to be compared with 73.83 Å$^3$ of undoped $\text{BaZrO}_3$.

**Photoluminescence properties.** PL spectra of the investigated ceramics are reported in Fig. 2. Panel a) displays the stability range of crystal phases and polar variants as determined from the study of the dielectric properties. The drawings serve as a quick eye guide to correlate structure and polar order with luminescence properties.

The shape of room temperature emission spectra (Fig. 2b), obtained exciting the $^5\text{L}_6 \leftrightarrow ^7\text{F}_0$ transition at 395 nm, strongly depends on sample composition. The PL spectrum of $\text{BZ0}$ shows four groups of bands that are characteristics for europium emission from the $^5\text{D}_0$ excited state to the $^7\text{F}_j$ ($j = 0, 1, 2, 3, 4$) manifolds located at: 578 nm ($^7\text{F}_0$), 590–596 nm ($^7\text{F}_1$), 613–623 nm ($^7\text{F}_2$) and 680–720 nm ($^7\text{F}_4$). The $^5\text{D}_0 \rightarrow ^7\text{F}_2$ band, if any, is quite faint. HR spectra of the $^5\text{D}_0 \rightarrow ^7\text{F}_1$ (Figure S3) band evidence four components indicating the presence of at least two nonequivalent Eu$^{3+}$ centers. In fact, considering the degeneration of the $j = 1$ level, a maximum of three components can be observed for a single site.

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**Figure 1.** (a–d) Real part of permittivity (relative dielectric constant) of $\text{Ba}_{1-y}\text{Eu}_y\text{Ti}_{1-x-y/4}\text{Zr}_x\text{O}_3$ ($y = 0.01$, $x = 0.0, 0.05, 0.15$ and $0.30$) ceramics at $10^2$–$10^6$ Hz. (e) Ferroelectric hysteresis loops at room temperature for different samples.
To verify the absence in the emission spectra of contributions coming from interface/foreign sites, sample BZ0 was annealed at 1450 °C for 48 h. This prolonged thermal treatment determined the exaggerated growth of some grains up to a size of 300–400 μm, thus allowing the intrinsic luminescence of Eu³⁺ in the perovskite lattice to be separated from other extrinsic contributions, such as the segregation of secondary phases (e.g. Eu₂Ti₂O₇) and/or glassy phase films at grain boundaries, if any. Exaggerated grain growth during prolonged sintering is quite common in pure and slightly doped BaTiO₃ ceramics and related to the formation of a liquid phase due to local stoichiometry variations or localized impurities.

PL spectra collected by means of a microscope on a polished cross-section at the center of a large grain and on the fine-grained matrix (Figure S4) are virtually identical. Therefore, the non-equivalent Eu³⁺ sites are associated with the perovskite lattice. Although the ionic radius of Eu³⁺ in octahedral coordination (0.947 Å) is rather large and the stoichiometry was designed for exclusive Ba-site substitution, a small amount of the rare-earth could be incorporated at the B site. The probability of Eu³⁺ incorporation in perovskite B-site increases with increasing Zr content, as the unit cell volume expands of about 15% moving from BaTiO₃ to BaZrO₃. Alternative and complementary explanations for the existence of multiple sites will be discussed further on studying the emission spectra as a function of temperature.

As x increases from 0 to 0.7, the intensity of the ⁵D₀ → ⁷F₂ band decreases, the ⁵D₀ → ⁷F₁ emission gets stronger and overcomes the intensity of the ⁵D₀ → ⁷F₂ transition in BZ50 and BZ70. The ⁵D₀ → ⁷F₂ emission is strongly reduced in sample with composition x = 1, and the spectrum is dominated by the sharp ⁵D₀ → ⁷F₁ transition. Noteworthy, a significant band broadening is observed for BZ30, BZ50 and BZ70 ceramics. These shapes closely resemble those of Eu³⁺ doped glasses in which Eu³⁺ ions occupy multiple sites having small differences in the coordination environment. Indeed, for intermediate compositions a significant contribution to PL is given by several sites corresponding to a different number of ZrO₆ (TiO₆) octahedra surrounding the Eu³⁺ ion. Moreover,
The heterogeneous local structure of the relaxor ceramics, corresponding to Ti-rich PNRs embedded in a cubic paraelectric matrix\textsuperscript{26,27}, further increases the disorder of the europium sites.

The narrowing of the $^{5}D_{0}\rightarrow^{7}F_{1}$ line and the drop of the intensity of $^{5}D_{0}\rightarrow^{7}F_{2}$ transition observed in the emission spectra of BZ100 suggests a decrease of the number of spectroscopic sites determined by the simpler composition and the disappearance of the PNRs. Very similar spectra are reported for BaZrO$_3$ and BaSnO$_3$ (cubic $Pm-3m$) in which Eu$^{3+}$ is incorporated at the Ba site\textsuperscript{36–39}. Indeed, the $^{5}D_{0}\rightarrow^{7}F_{1}$ transition is the most intense when Eu$^{3+}$ occupies a centrosymmetric position such as $Oh$ site of Ba in BaZrO$_3$ for which all transitions but $^{5}D_{0}\rightarrow^{7}F_{1}$ are forbidden.

Not only the composition but also the temperature has a strong influence on the Eu$^{3+}$ emission. At $-20^\circ$C (Fig. 2b) the spectra of the ferroelectric ceramics (BZ0, BZ5 and BZ15) are dominated by the $^{5}D_{0}\rightarrow^{7}F_{1}$ bands whose integrated intensity is three and a half times larger compared to the $^{5}D_{0}\rightarrow^{7}F_{4}$ transition that is the most intense at room temperature. All other bands, and in particular the $^{5}D_{0}\rightarrow^{7}F_{1}$ one, have a much lower intensity. The $^{5}D_{0}\rightarrow^{7}F_{4}$ transition is the strongest even in sample BZ30 in comparison to RT. The spectra of BZ50, BZ70 and BZ100 are instead very similar to those observed at room temperature. At $-100^\circ$C (Fig. 2c), the $^{5}D_{0}\rightarrow^{7}F_{4}$ transition is the most intense for all ceramics except BZ100, for which it has the same order of magnitude as $^{5}D_{0}\rightarrow^{7}F_{2}$ and $^{5}D_{0}\rightarrow^{7}F_{1}$. No significant changes in the shape of PL spectra of ferroelectric ceramics are observed between $-20$ and $-100^\circ$C. It is worth noting that band broadening in the PL spectra of relaxors is unaffected by temperature variations.

The detailed evolution of the PL spectra for some representative samples (BZ0, BZ15, BZ50 and BZ100) over the whole investigated temperature range, $-100$ to $140^\circ$C, is illustrated in the color maps reported in Fig. 3. The maps of BZ5, BZ30 and BZ70 samples are shown in Figure S5.

Among other features, in ferroelectric Ti-rich ceramics (BZ0, BZ5 and BZ15) the main temperature-induced spectral modification is represented by the inversion of the intensity of the $^{5}D_{0}\rightarrow^{7}F_{1}$ and $^{5}D_{0}\rightarrow^{7}F_{4}$ transitions...
The ionic radius (c.n. 12) of Eu$^{3+}$ (1.226 Å) is much smaller than that of Ba$^{2+}$ (1.610 Å) and, consequently, the lanthanide ion is over-coordinated when it is positioned at the Ba site and the possibility of occupying a lower symmetry site has to be considered. Very recent investigations including careful crystal structure determination from synchrotron X-ray diffraction data and first-principle calculations on Gd- and Dy-doped BaTiO$_3$ as well as the study of dielectric relaxations in Gd- and Dy-doped SrTiO$_3$ have provided some evidence about the off-center displacement of trivalent lanthanide ions with average size at A-site of these perovskites.

As illustrated in Fig. 5, and referring to the cubic structure for the sake of simplicity, a displacement of about 1 Å along one of the six [100] directions (towards the center of any of the cube faces) drives Eu$^{3+}$ ions off-center displacement of trivalent lanthanide ions with average size at A-site of these perovskites. The off-center displacement of Eu$^{3+}$ ions is not affected by the off-center displacement of Ti$^{4+}$ ions.

According to the above results (see Figs 2, 3 and S6), the ferroelectric ceramics show PL emission dominated by the $^{3}D_0 \rightarrow ^{7}F_4$ transitions over a broad temperature range, from $-100^\circ$C up to room temperature or even higher (up to 60–80$^\circ$C for $x = 0$). Emission spectra characterized by an abnormally high intensity of $^{3}D_0 \rightarrow ^{7}F_4$ transition are reported for Eu$^{3+}$ occupying $D_{ad}$ (or distorted $D_{ad}$) sites with coordination geometry close to square antiprism as well as distorted $D_{ad}$ sites. For an undistorted $D_{ad}$ symmetry the $^{3}D_0 \rightarrow ^{7}F_4$ transition is forbidden whereas the transitions $^{3}D_0 \rightarrow ^{7}F_1$ and $^{3}D_0 \rightarrow ^{7}F_4$ are allowed. This situation is uncommon and occurs only for the point groups $D_{ad}$ and O.

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As illustrated in Fig. 5, and referring to the cubic structure for the sake of simplicity, a displacement of about 1 Å along one of the six [100] directions (towards the center of any of the cube faces) drives Eu$^{3+}$ from the
is similar to the Eu-O distance in Eu2O3 and Eu2Ti2O7, 2.2–2.3 Å). In principle, the lanthanide can also move in other directions. Off-center displacement of Eu3+ suggests that [100] displacement is preferred over [111]. Opposite, the position of Sr cations is largely central. The different behavior of the two earth-alkaline cations is essentially determined by their different ionic radius (c.n. 12), 1.34 Å for Ca2+ and 1.44 Å for Sr2+, being their chemical properties similar. It is worth noting that Ca2+ in CaTiO3 (orthorhombic Pnma perovskite structure) occupies 8-coordinated distorted sites with Ca-O distances between 2.36 and 2.67 Å. The remaining 4 distances are > 3 Å.

Following the same reasoning, the additional driving force determining off-centering of Eu3+ in ferroelectric ceramics is the enhancement of polar order and ferroelectricity resulting from the correlated movement of Eu3+. 

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**Figure 5.** Evolution of local symmetry during the off-center displacement of Eu3+ along one of the [100] directions of cubic BaTiO3. See text for details. Red spheres: O2−. Green sphere Eu3+. The images were realized using VESTA56.
and Ti$^{4+}$ ions. The off-center displacement of Eu$^{3+}$ will give rise to an electrical dipole. If Eu$^{3+}$ preferentially moves along one of the [100] directions in such a way that the component of the associated dipole moment will positively contribute to the dipole moment corresponding to the [111] relaxation of the Ti ions, the europium displacement will concur to the overall polarization and ferroelectricity. This mechanism can be amplified by the deformation of the TiO$_6$ octahedra nearest to Eu$^{3+}$ as happens with Ca$^{2+}$. Opposite, if Eu$^{3+}$ moved randomly, there would be no net contribution to ferroelectricity and no extra energy gain determined by off-centering. Hence, the absence of polar order in BaZrO$_3$ prevents a correlated displacement of Eu$^{3+}$ and Zr$^{4+}$ and the lanthanide will largely stay in central position. It should be kept in mind that the concentration of Eu$^{3+}$ ions is as low as 1% and, consequently, their mutual interaction will be negligible thus hampering a cooperative behavior of the lanthanide ions alone.

The evolution of the PL spectra with temperature is mainly ascribed to the change of the population of the different Eu$^{3+}$ sites. For ferroelectric ceramics, the off-centering of Eu$^{3+}$ occurs over a broad temperature range (up to 60–80 °C for BaTiO$_3$) by the cooperative polar interactions discussed above and made possible by the existence of ferroelectric order. The polar interactions counteract the thermal motion which would favor the occupation of the central sites or more disordered distributions. In the absence of polar interactions, as in BaZrO$_3$, off-centering of some Eu$^{3+}$ can only occur at low temperature (≤80 °C), as indicated by the increasing contribution of the $^5D_0 \rightarrow ^3F_4$ transition to the overall emission (Figs 2 and 3), thanks to the reduced thermal motion. In relaxors (BZ30, BZ50, BZ70), Eu$^{3+}$ off-center displacement should be confined within PNRs, whereas a behavior similar to that exhibited by BaZrO$_3$ is expected in the paraelectric matrix. However, the PL spectra of BZ30 below −40°C are very similar to those of the ferroelectric samples (Figs 2, 3 and S5). This observation supports the idea that the ferroelectric to relaxor crossover in BaZr$_{1−x}$Ti$_x$O$_3$ occurs over the composition range 0.25 ≤ x < 0.35 with coexistence of PNRs and ferroelectric domains, the latter originated by percolation of polar clusters on cooling.$^{57, 60}$ The rapid increase of the $^5D_0 \rightarrow ^3F_4$ emission at lower temperature in the other two relaxors (Figs 2–4 and S5) is again attributed to the suppression of thermal motion and off-centering of some Eu$^{3+}$ ions.

One can wonder if the evolution of the PL spectra with temperature might be determined by a change of defect chemistry, for example by a redistribution of Eu$^{3+}$ ions between A and B sites in the perovskite. However, owing to the very low mobility of cation and oxygen vacancies in the investigated temperature range, the defect chemistry of BaTiO$_3$ and BaZrO$_3$ is essentially frozen. The most mobile kind of ionic defects, start to be mobile above 400 °C.$^{61}$ A change in the distribution of Eu$^{3+}$ between Ba and Ti sites is unlikely because this would imply a complete re-equilibration of the bulk defect chemistry and in particular the concentration of the charge compensating defects (cation vacancies for Ba-site incorporation, oxygen vacancies for B-site substitution) which can only occur by long-range diffusional transport. In contrast, the change of the Eu$^{3+}$ position from the center of the dodecahedron to a nearby acentric site occurs over a very short distance (≈1 Å) and does not require diffusion.

Conclusions

An effective tuning of the photoluminescence of rare-earth ions incorporated in ferroelectric perovskites has recently been achieved by exploiting the strong coupling between the lattice strain and the electric field (or the mechanical stress) typical of ferroelectric materials. However, as shown in the present paper, the polar order and, in particular, the ferroelectric order, has a direct and remarkable impact on the photoluminescence of Eu$^{3+}$ in the BaZr$_{1−x}$Ti$_x$O$_3$ perovskite even in the absence of an external field.

In the ferroelectric ceramics (x = 0–0.15), the photoluminescence spectra display a crossover from a dominating $^3D_0 \rightarrow ^3F_4$ emission above room temperature to a dominating $^5D_0 \rightarrow ^3F_4$ emission at lower temperature. This behavior provides strong evidence of the off-center displacement of Eu$^{3+}$ along one of the [100] directions of the lattice into a site with approximate D$_4h$ symmetry with a coordination polyhedron close to a square antiprism with decreasing temperature. The off-centering of the lanthanide from the regular dodecahedral Oh site of Ba$_2$O$_3$ can be attributed to the cooperative polar interactions with the ferroelectric lattice, i.e. the correlated displacement of Eu$^{3+}$ and Ti$^{4+}$ ions aided by the smaller ionic radius of Eu$^{3+}$ (1.226 Å) in comparison to Ba$^{2+}$ (1.610 Å). Conversely, in BaZrO$_3$ the absence of ferroelectric order prevents a correlated displacement of Eu$^{3+}$ and Zr$^{4+}$ and the lanthanide will largely occupy the central dodecahedral Oh site in spite of the larger unit cell of the zirconate in comparison to BaTiO$_3$. Only at low temperature (~100 °C) the suppression of thermal motion allows off-centering of some Eu$^{3+}$ ions.

Materials with compositions x = 0.30–0.70 have a short-range polar order typical of relaxors and their photoluminescence shows a distinct crossover but a more complex behavior. The complexity partly arises from the heterogeneous nature of the relaxors at the nanoscale, corresponding to BaTiO$_3$ polar nanoregions embedded in a paraelectric matrix. Moreover, for these intermediate compositions several Eu$^{3+}$ sites corresponding to a different number of ZrO$_6$ (TiO$_6$) octahedra surrounding the lanthanide will give a significant contribution to the luminescence.

It is expected that other RE$^{3+}$ ions with intermediate size (Sm$^{3+}$ to Er$^{3+}$), when incorporated at the Ba site of the perovskite, can exhibit a behavior similar to that of Eu$^{3+}$ as the underlying mechanism is rather general and not restricted to a specific ion. However, the magnitude of the off-center displacement is likely to depend on the ionic radius of the dopant producing a systematic trend. Furthermore, a ferroelectric/polar order control of photoluminescence could be observed in other perovskites.

Experimental Section

Sample preparation. Six Eu-doped Ba(Ti,Zr)O$_3$ ceramics with composition Ba$_{1−x}$Eu$_x$Ti$_{1−y}$Zr$_y$O$_3$ (y = 0.01; x = 0, 0.05, 0.15, 0.30, 0.50, 0.70) and Ba$_{1−x}$Eu$_x$Zr$_{1−y}$O$_3$ (y = 0.01) were prepared by the classical solid-state route using industrial electronic grade precursors: BaCO$_3$ (Solvay Bario e Derivati, Italy), TiO$_2$ (Evonik Degussa grade P25, Germany), ZrO$_2$ (Toho grade TZ0, Japan) and Eu$_2$O$_3$ (Metall Rare Earth Ltd, China) powders as raw materials. The compositions correspond to Eu$^{3+}$ substitution at the Ba site with Ti vacancy compensation (0 ≤ x ≤ 0.7) and Zr vacancy compensation (x = 1). The different samples are labelled as BZx, where X = 100x. Predominant substitution at the Ba site of BaTiO$_3$ is reported for Ba/Ti < 1 up to 2–3 at.% europium.$^{62, 63}$
the solubility limit of trivalent europium. The low dopant concentration adopted (1 mol.%) guarantees that the structural and physical properties of the materials are not significantly affected. Precursor powders were wet-mixed in polypropylene jars using water as liquid and a solution of ammonium polyacrylate (pH = 10) as dispersant. After drying, the mixed powder was calcined for 4 h at 1000 °C and then compacted in cylinders (length: 1 cm, diameter: 1 cm) by isostatic pressing at 1500 bar. The resulting green were sintered in air for 4 h at different temperatures depending on composition: 1450 °C (BZ0-BZ30), 1550 °C (BZ50-BZ70) and 1600 °C (BZ100). The relative density of the final samples was determined by the Archimedes’ method. The ceramics are well densified with relative density ≥94%. Ceramic disks with a thickness of about 1 mm were cut from the sintered body and characterized by different techniques.

**Crystal structure, dielectric and ferroelectric properties.** The phase composition and crystal structure was investigated by X-ray diffraction (XRD) using a Cubix diffractometer (Panalytical, The Netherlands) with Cu Kα radiation (30 kV, 30 mA). The lattice parameters were determined using FullProf 2000. For dielectric measurements, Ag-Pd electrodes were deposited on the plane-parallel polished surfaces of the disks followed by annealing in air at 500 °C for 12 h. The dielectric properties were measured by an impedance bridge E4980A Precision LCR Meter (Agilent, Santa Clara, CA) and a dielectric spectrometer CONCEPT40 (Novocontrol Technologies, Hundsangen, Germany) in the temperature range −150–150 °C at 102–106 Hz. Polarization – electric field ferroelectric loops were recorded at room temperature on the electroded ceramics immersed in transformer oil bath by a Sawyer–Tower modified circuit fed by triangular high voltage wave (frequency: 10 Hz) by using a TREK amplifier.

**Photoluminescence.** Room temperature luminescence spectra were recorded on solid samples in a front-face acquisition geometry with a spectrofluorimeter (Fluorolog-3, Horiba JobinYvon) equipped with double-grating monochromator in both the excitation and emission sides, coupled to a R928P Hamamatsu photomultiplier and a 450 W Xe arc lamp as the excitation source. The emission spectra were corrected for detection and optical spectral response of the spectrofluorimeter supplied by the manufacturer.

HR spectra and temperature dependent experiments (~100–140 °C) were carried out in backscattering geometry using a Horiba T64000 triple spectrometer equipped with a Peltier-cooled charge-coupled device detector (Horiba Synapse). The scattered radiation was collected through a 10 × microscope objective (Olympus MPLAN, 10 ×/0.25). The spectrograph, equipped with 2400 lines/mm gratings for high resolution and 300 lines/mm for temperature dependent experiments, was used as a single stage spectrograph. Temperature dependent experiments were performed by means of a Linkam THMS600 heating/cooling microscope stage having temperature stability <0.1 °C over −196 °C to 600 °C temperature range.

**Data Availability**
All data generated or analysed during this study are included in this published article (and its Supplementary Information files).

**References**
1. Feldmann, C., Justel, T., Ronda, C. R. & Schmidt, P. J. Inorganic luminescent materials: 100 Years of research and application. *Adv. Funct. Mater.* 13, 511–516 (2003).
2. Eliseeva, S. V. & Bünzli, J. C. G. Lanthanide luminescence for functional materials and bio-sciences. *Chem. Soc. Rev.* 39, 189–227 (2010).
3. Bai, G., Tsang, M. K. & Hao, J. Tuning the luminescence of phosphors: Beyond conventional chemical method. *Adv. Opt. Mater.* 3, 431–462 (2015).
4. Hao, J., Zhang, Y. & Wei, X. Electric-induced enhancement and modulation of upconversion photoluminescence in epitaxial BaTiO3:Yb/Er thin films. *Angew. Chemie - Int. Ed.* 50, 6876–6880 (2011).
5. Yao, Q. *et al.* Electric field-induced giant strain and photoluminescence–enhancement effect in rare-earth modified lead-free piezoelectric ceramics. *ACS Appl. Mater. Interfaces* 7, 5066–5075 (2015).
6. Sun, H. L., Wu, X., Chung, T. H. & Kwok, K. W. In-situ Electric Field-Induced Modulation of Photoluminescence in Pr-doped Ba0.85Ca0.15Ti0.90Zr0.10O3 Lead-Free Ceramics. *Sci. Rep.* 6, 1–8 (2016).
7. Wang, F. *et al.* In situ reversible tuning of photoluminescence of an epitaxial thin film via piezoelectric strain induced by a Pb(Mg0.53Nb0.47)O3–PbTiO3 single crystal. *J. Mater. Chem.* C 5, 9115–9120 (2017).
8. Sun, H., Wu, X., Peng, D. F. & Kwok, K. W. Room-temperature large and reversible modulation of photoluminescence by in situ electric field in ergodic relaxor ferroelectrics. *ACS Appl. Mater. Interfaces* 9, 34042–34049 (2017).
9. Tu, D. *et al.* LiNbO3:Pr3+: A Multipiezoelectric Material with Simultaneous Piezoelectricity and Sensitive Piezoluminescence. *Adv. Mater.* 29, 1–4 (2017).
10. Zhang, P. *et al.* Pr5+ photoluminescence in ferroelectric (Ba0.53Ca0.47)TiO3 ceramics: Sensitive to polarization and phase transitions. *Appl. Phys. Lett.* 92, 3–6 (2008).
11. Tian, X. *et al.* Remanent-polarization-induced enhancement of photoluminescence in Pr5+ doped lead-free ferroelectric (Bi0.5Na0.5)TiO3 ceramics. *Appl. Phys. Lett.* 102, 7–10 (2013).
12. Du, P., Luo, L., Li, W., Zhang, Y. & Chen, H. Photoluminescence and piezoelectric properties of Pr-doped NBT-xBZT ceramics: Sensitive to structure transition. *J. Alloys Compd.* 559, 92–96 (2013).
13. Wei, Y. *et al.* Dual-enhancement of ferro-/piezoelectric and photoluminescent performance in Pr5+ doped (K0.5Na0.5)NbO3 lead-free ceramics. *Appl. Phys. Lett.* 105, 1–5 (2014).
14. Yao, Y., Luo, L., Li, W., Zhou, J. & Wang, F. An intuitive method to probe phase structure by upconversion photoluminescence of Er3+ doped in ferroelectric Pb(Mg0.53Nb0.47)O3–PbTiO3. *Appl. Phys. Lett.* 106, 1–5 (2015).
15. Khattaa, D. K., Kalaskar, A. & Ranjan, R. Tuning Photoluminescence Response by Electric Field in Electrically Soft Ferroelectrics. *Phys. Rev. Lett.* 116, 2–6 (2016).
16. Haertling, G. H. Ferroelectric Ceramics: History and Technology. *J. Am. Ceram. Soc.* 82, 797–818 (1999).
17. Hiroshi, K., Youichi, M. & Hirokazu, C. Base-Metal Electrode-Multilayer Ceramic Capacitors: Past, Present and Future Perspectives. *Jpn. J. Appl. Phys.* 42, 1–3 (2003).
18. Kwei, G. H., Lawson, A. C., Billinge, J. L. & Cheong, S.-W. Structure of the Ferroelectric Phase of Barium Titanate. *J. Phys. Chem.* 97, 2368–2377 (1993).
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Author Contributions
All the authors contributed to the experiments. In particular G.C., M.T.B., C.C. and V.B. prepared the samples and made structural characterization. G.B. and L.A. performed the luminescence study. O.C., L.C., L.M. performed the electrical characterization of the samples. G.B. and V.B. wrote the manuscript.

Additional Information
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