



Title	Electronic chirality inversion of lanthanide complex induced by achiral molecules
Author(s)	Wada, Satoshi; Kitagawa, Yuichi; Nakanishi, Takayuki; Gon, Masayuki; Tanaka, Kazuo; Fushimi, Koji; Chujo, Yoshiki; Hasegawa, Yasuchika
Citation	Scientific reports, 8, 16395 https://doi.org/10.1038/s41598-018-34790-0
Issue Date	2018-11-06
Doc URL	http://hdl.handle.net/2115/72145
Rights(URL)	https://creativecommons.org/licenses/by/4.0/
Type	article
Additional Information	There are other files related to this item in HUSCAP. Check the above URL.
File Information	s41598-018-34790-0.pdf



[Instructions for use](#)

SCIENTIFIC REPORTS



OPEN

Electronic chirality inversion of lanthanide complex induced by achiral molecules

Satoshi Wada¹, Yuichi Kitagawa², Takayuki Nakanishi³, Masayuki Gon⁴, Kazuo Tanaka⁴, Koji Fushimi², Yoshiki Chujo⁴ & Yasuchika Hasegawa²

A novel mechanism for chiroptical activity inversion based on the electronic structure of metal complexes without Λ - or Δ -type structure change was demonstrated spectroscopically and theoretically. To demonstrate the mechanism, a europium (Eu(III)) complex with chiral (+)-3-(trifluoroacetyl)camphor (+tfc) and achiral triphenylphosphine oxide (tppo) was prepared. The steric and electronic structures of the Eu(III) complex were adjusted by additional achiral tppo and coordinating acetone molecules, and were characterised by ¹H NMR, photoluminescence, and emission lifetime measurements. The optical activity of the Eu(III) complex in solution was evaluated by circularly polarized luminescence (CPL) measurements. CPL sign inversion, which was independent of Λ - or Δ -type structure changes from the spectroscopic viewpoint, and a drastic CPL intensity enhancement were observed depending on the external achiral molecules around Eu(III) ion. These phenomena provide the first clarification of optical activity change associated with electronic structure rather than chiral coordination structure-type (Λ or Δ) under external environments.

Chirality is prevalent in our life. Chirality is attributed to asymmetric molecular structure, which arises from the molecular level to the macroscopic helical scale, like a DNA double helix^{1,2}. Chiral structures also play a critical role in chiral properties such as enzyme activity and asymmetric synthesis using chiral catalysts^{3–5}. The chiral structures are susceptible to external environments such as solvents⁶, temperature⁷, light⁸, or pH⁹, causing chiral inversion phenomena in large variety of systems from biological macromolecules like B- or Z-form DNA¹⁰ to artificial soft matter and liquid crystals^{11–14}. Hence, the influence of external environment is of importance to control the molecular chirality.

Generally, chiral molecules exhibit optical activities with absorption (circular dichroism: CD) or emission (circularly polarized luminescence: CPL) related to the molecular structure¹⁵. Optical activities are estimated from the different electronic transition probabilities between left- and right-handed circularly polarized light. The CD and CPL signals are dependent on the molecular chirality, which is influenced by the external field. The inversion of optical activity induced by the external field has been reported extensively^{16–25}. Ishii and Verbiest achieved the CD signal inversion of a chiral polythiophene derivative by controlling the molecular aggregation speed¹⁶. George observed the tuneable helicity inversion of Zn(II)-coordinated naphthalenediimide assemblies with chiral adenosine phosphates using CD measurements¹⁷. Meskers and Swager described the solvent polarity-dependent CPL signal switching of a self-assembled chiral poly(*p*-phenylenevinylene) derivative¹⁸. The magnitude of optical activity is estimated by a dissymmetry factor (*g*), which is composed of electric and magnetic field components related to the electronic transition¹⁵. Their components are governed by the molecular structure in the surrounding environment.

Here, we targeted chiral lanthanide complexes to clarify the influence of electric and magnetic field components on optical activities. Their sharp luminescence bands comprise electric and magnetic field components of 4f-4f transitions²⁶. Their components in the electronic transitions are dominated by external organic molecules around the lanthanide ion. In particular, the magnetic dipole transition of the europium (Eu(III)) complex

¹Graduate School of Chemical Sciences and Engineering, Hokkaido University, N13 W8, Kita-ku, Sapporo, Hokkaido, 060–8628, Japan. ²Faculty of Engineering, Hokkaido University, N13 W8, Kita-ku, Sapporo, Hokkaido, 060–8628, Japan. ³Faculty of Industrial Science and Technology, Tokyo University of Science, 6-3-1 Nijjuku, Katsushika-ku, Tokyo, 125-8585, Japan. ⁴Graduate School of Engineering, Kyoto University, Katsura, Nishikyo-ku, Kyoto, 615-8510, Japan. Correspondence and requests for materials should be addressed to Y.K. (email: y-kitagawa@eng.hokudai.ac.jp) or Y.H. (email: hasegaway@eng.hokudai.ac.jp)

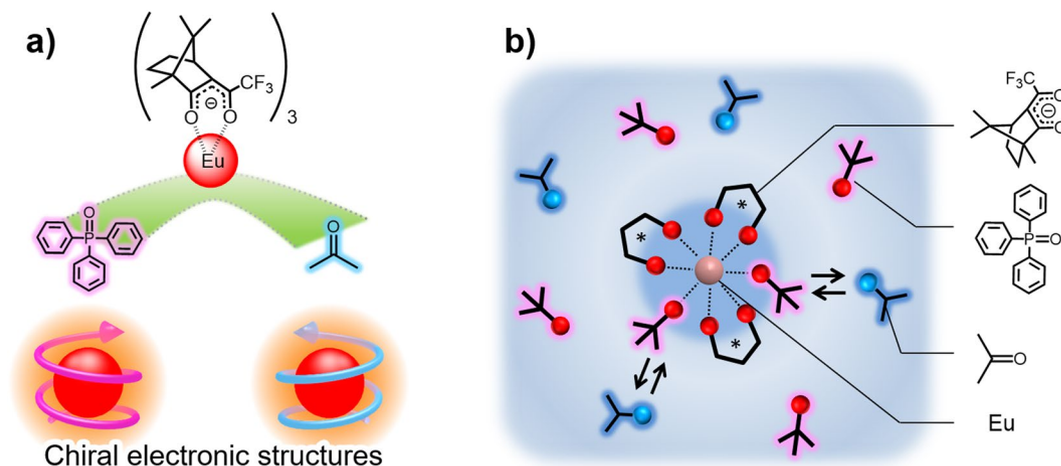


Figure 1. Schematic images of (a) chiral electronic structures depending on the external molecules around the Eu(III) ion and (b) the coordination structures of the Eu(III) complex depending on external achiral molecules around the Eu(III) ion in solution.

exhibits a large dissymmetry factor on the CPL (g_{CPL}) by the introduction of chiral organic ligands; this is established by the effective contribution of the magnetic field component^{27–30}. The optical activity signal of the lanthanide complex with chiral ligands has been evaluated by Λ - or Δ -type coordination structures previously^{31,32}. Yuasa and Parker reported the chiroptical activity inversion of chiral Eu(III) complexes influenced by the different steric structures of achiral molecules^{33,34}, which implied steric inversion between Λ - and Δ -type structures. Note that chiroptical activity is related to the electronic transition of the Eu(III) complex; therefore, the magnitude and sign could be also affected by the chiral electronic structure of the Eu(III) ion surrounded by external ligands (Fig. 1a).

In order to demonstrate the chiroptical activity based on the chiral electronic structure in external environments, we choose the chiral Eu(III) complex with bidentate (+)-3-(trifluoroacetyl)camphor (+tfc) and monodentate triphenylphosphine oxide (tppo) as the chiral and external achiral ligands, respectively. The Eu(III) complex $[\text{Eu}(+\text{tfc})_3(\text{tppo})_2]$ (Δ -type structure, as determined by X-ray single-crystal analysis) shows a large g_{CPL} (-0.47) in acetone- d_6 ³⁵. The chiral electronic structure of the Eu(III) complex in solution was adjusted by additional achiral tppo and acetone molecules (Fig. 1b). The coordination and electronic structures in liquid media were characterised using ¹H NMR, photoluminescence, and emission lifetime measurements. The chiroptical activities of the Eu(III) complex under external environments were evaluated using CPL measurements. In this study, we demonstrate a mechanism for the optical activity inversion based on the chiral electronic structure of the Eu(III) complex without Λ - or Δ -type structure change spectroscopically and theoretically, for the first time.

Results

Coordination structures in solution. X-ray crystallography measurements indicated that the coordination structure of the Eu(III) complex in the solid is an eight-coordinated Δ -type structure composed of three chiral +tfc and two tppo ligands³⁵. To evaluate the conformation of $[\text{Eu}(+\text{tfc})_3(\text{tppo})_2]$ (**Eu**(+)) in acetone, ¹H NMR spectra with additional n equivalents ($n = 0, 8, 28, 48,$ and 98 relative to **Eu**(+)) of tppo molecules, namely **Eu**(+)-Ex n , were acquired (Fig. 2 and Supplementary Table S1). In the low-magnetic-field side, protons of tppo molecules in **Eu**(+)-Ex0 show broad peaks in Fig. 2 (black; A). The line-broadening and chemical shifts originate from the exchange reaction and paramagnetic effect on the metal complex^{36,37}. The paramagnetic effect of the Eu(III) ion generally induces little broadening (bandwidth; nearly 10 Hz)³⁷. Therefore, the large broadening of tppo signals in our experiment (Fig. 2, black; A, bandwidth; nearly 300 Hz) is mainly caused by their exchange reaction in acetone- d_6 . The lower-magnetic-field shift in **Eu**(+)-Ex0 (Fig. 2, black; A) is influenced by the direct coordination of tppo ligands with the Eu(III) ion.

NMR peaks of the chiral +tfc ligands in the Eu(III) complex were observed in the high-magnetic-field side (Fig. 2, signals B–I). **Eu**(+)-Ex8 (green) provided effective chemical shifts of +tfc ligands compared with those of **Eu**(+)-Ex0 (black). We also observed gradual shifts at the B, E, G, and I peaks of **Eu**(+)-Ex28 (purple), -Ex48 (red), and -Ex98 (blue). The effective shifts of the tppo and +tfc signals indicate that the Eu(III) complex with tppo molecules is rearranged by additional tppo molecules, resulting in the formation of several equilibrium states in acetone- d_6 .

Luminescence properties. Photophysical properties of Eu(III) complexes are affected by the coordination geometry³⁸. The emission spectra of **Eu**(+)-Ex0 and -Ex498 in acetone (1×10^{-3} M) are shown in Fig. 3 (black; a, and red; b). The Eu(III) complexes show sharp emission peaks in the region of 570–630 nm, which are attributed to the ⁵D₀ → ⁷F_{*J*} ($J = 0, 1,$ and 2) transitions of Eu(III) ions. The spectra were normalised with respect to the integrated intensities of the magnetic dipole transition (⁵D₀ → ⁷F₁). Their spectral shapes in liquid media were different from that of $[\text{Eu}(+\text{tfc})_3(\text{tppo})_2]$ in the solid state (Supplementary Fig. S1). The emission spectra for the ⁵D₀ → ⁷F₁ and ⁵D₀ → ⁷F₂ transitions were also changed in response to the concentration of the tppo molecules in

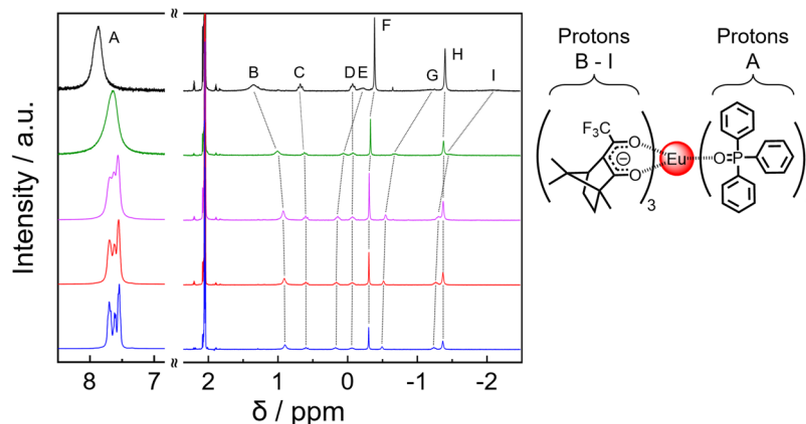


Figure 2. ^1H NMR spectra of **Eu(+)-Ex0** (black), **Eu(+)-Ex8** (green), **Eu(+)-Ex28** (purple), **Eu(+)-Ex48** (red), and **Eu(+)-Ex98** (blue) in acetone- d_6 (**Eu(+)**; 1×10^{-3} M).

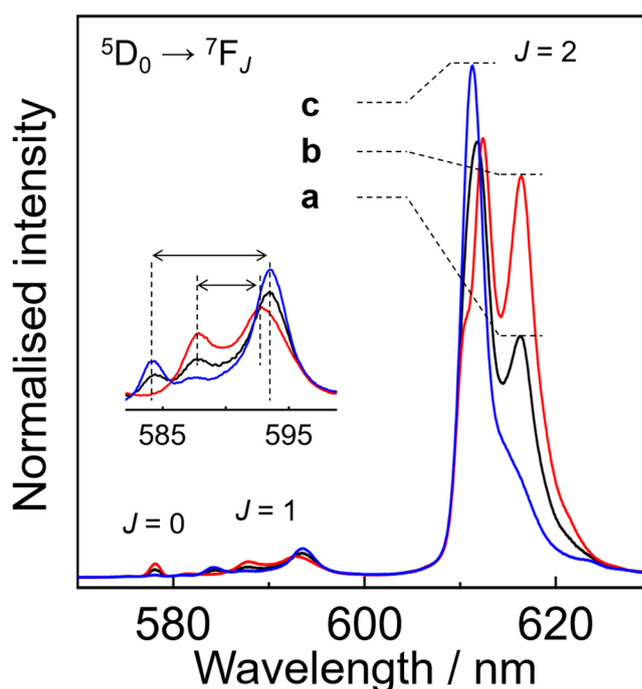


Figure 3. Photoluminescence spectra of (a) **Eu(+)-Ex0** (1×10^{-3} M, black), (b) **Eu(+)-Ex498** (1×10^{-3} M, red), and (c) **Eu(+)-Ex0** (1×10^{-5} M, blue) excited at 350 nm in acetone.

solution. In particular, the $^5\text{D}_0 \rightarrow ^7\text{F}_1$ transition band is composed of three Stark sublevels under the electric field (crystal field). **Eu(+)-Ex0** showed three peaks at 584.5, 588, and 593.5 nm in the $^5\text{D}_0 \rightarrow ^7\text{F}_1$ transition (Fig. 3 inset, black; a), whereas **Eu(+)-Ex498** showed two peaks at 588 and 593 nm (Fig. 3 inset, red; b). The $^5\text{D}_0 \rightarrow ^7\text{F}_1$ transition of **Eu(+)-Ex0** at a lower concentration (1×10^{-5} M, Fig. 3, blue; c) also showed three peaks at 584, 587.5, and 593.5 nm, which are similar to that of **Eu(+)-Ex0** at a higher concentration. The small peak at 587.5 nm can be attributed to the $^5\text{D}_1 \rightarrow ^7\text{F}_3$ transition, which is sometimes observed in the same energy region as the $^5\text{D}_0 \rightarrow ^7\text{F}_1$ transition³⁸. The emission bands at around 612 nm are attributed to hypersensitive electric dipole transitions ($^5\text{D}_0 \rightarrow ^7\text{F}_2$), which are strongly dependent on the local symmetry of the Eu(III) ion. The change of spectral shape is influenced by the rearrangement of coordination geometries of the Eu(III) complex depending on additional tppo molecules. In case of the non-coordinating toluene solution, the emission spectra of **Eu(+)-Ex0** and **-Ex48** were similar in shape to that of **Eu(+)-Ex498** in acetone, irrespective of the amount of additional tppo molecules (Supplementary Fig. S2). We propose that the inner coordination structure of **Eu(+)-Ex498** in acetone is composed of one Eu(III) ion, three +tfc ligands, and two tppo ligands.

The time-resolved emission profiles of **Eu(+)-Ex0** and **-Ex498** in acetone (1×10^{-3} M) were measured to clarify their coordination structures. The emission lifetimes were estimated using triple (for **Eu(+)-Ex0**) or double (for **Eu(+)-Ex498**) exponential functions to analyse several conformations in solution (Fig. 4a,b). The estimated

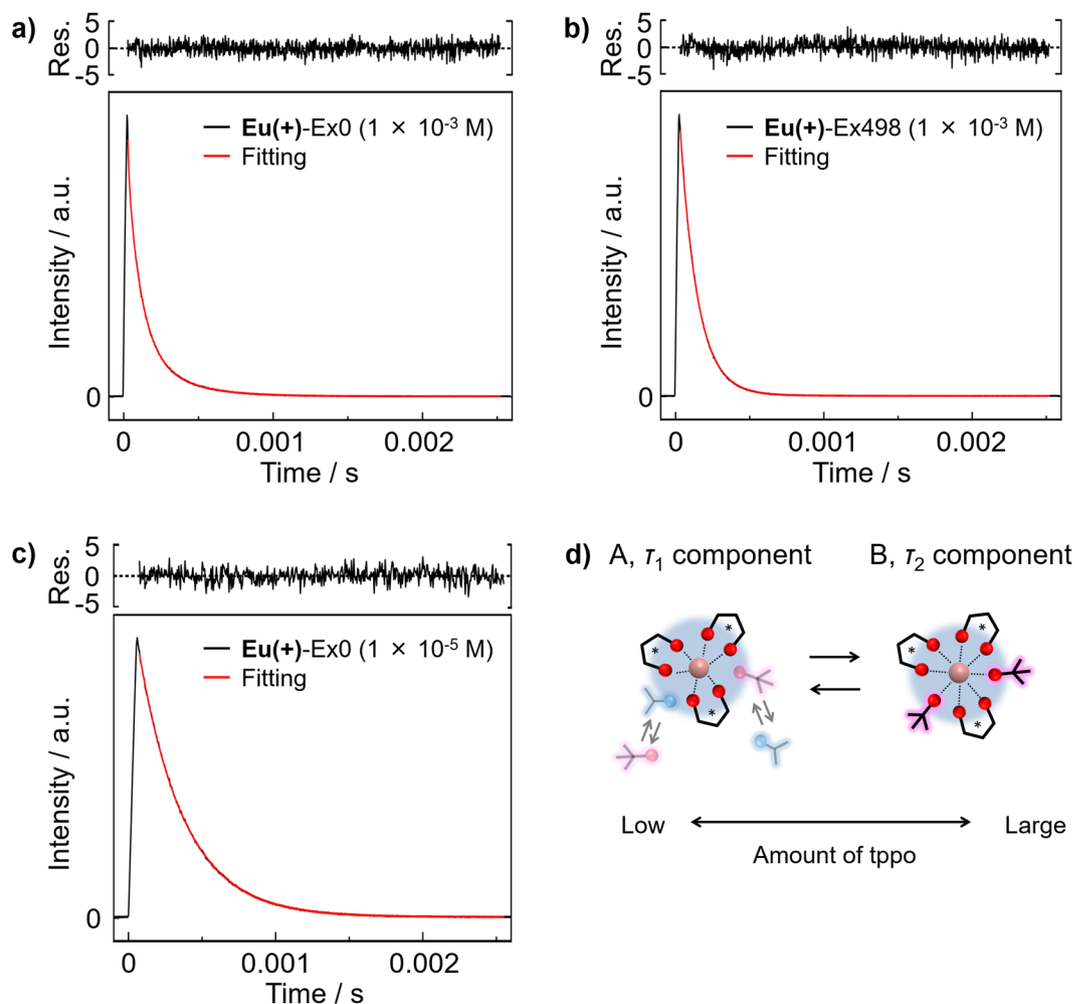


Figure 4. Emission decay profiles, fittings, and residuals (Res.) of (a) **Eu(+)-Ex0** (1×10^{-3} M), (b) **Eu(+)-Ex498** (1×10^{-3} M), and (c) **Eu(+)-Ex0** (1×10^{-5} M) at 612 nm excited at 356 nm in acetone. (d) The coordination image of the τ_1 and τ_2 components of the **Eu(III)** complex in acetone.

	Concentration [M]	τ_1 [ms]	τ_2 [ms]	τ_3 [ms]	g_{CPL}
Eu(+)-Ex0	1×10^{-3}	0.27 (50%)	0.09 (49%)	0.01 (1%)	-0.44
Eu(+)-Ex498	1×10^{-3}	0.37 (3%)	0.12 (97%)	—	+0.013
Eu(+)-Ex0	1×10^{-5}	0.31 (100%)	—	—	-1.0

Table 1. Luminescence properties of **Eu(+)-Ex n** excited at 356 nm in acetone^a. ^aEmission decay curves were analysed by multi-exponential curve fittings [$I(t) = \sum A_i \exp(-t/\tau_i)$]. The ratio of each component denotes $100 \times A_i \tau_i / \sum A_i \tau_i$.

emission lifetimes are summarised in Table 1. For **Eu(+)-Ex0**, the emission lifetimes (τ_1 , τ_2 , and τ_3) and their ratios were calculated to be 0.27 ms (50%), 0.09 ms (49%), and 0.01 ms (1%), respectively. The longer component τ_1 in higher concentration (1×10^{-3} M, 0.27 ms) was similar to the single component τ_1 in lower concentration (1×10^{-5} M, 0.31 ms, Fig. 4c). We consider that the τ_1 component is the **Eu(III)** complex with coordinating acetone molecules (Fig. 4a,d). The τ_1 and τ_2 component ratios decreased and increased, respectively, with increasing amount of tppo molecules. The main τ_2 value of **Eu(+)-Ex498** was found to be 0.12 ms (97%). The lifetime τ_2 in acetone was similar to the single-lifetime component in toluene (Supplementary Table S2), indicating that the **Eu(III)** ion is attached with three +tfc and two tppo ligands (Fig. 4b,d). We revealed that the two types of steric structures with τ_1 and τ_2 components were reorganized in response to the external tppo and acetone molecules.

Chiroptical properties. The CPL spectra and dissymmetry factors of **Eu(+)-Ex0** and -Ex498 are shown in Fig. 5 and Table 1, respectively. The CPL signals for the $^5D_0 \rightarrow ^7F_1$ transition were composed of two peaks at

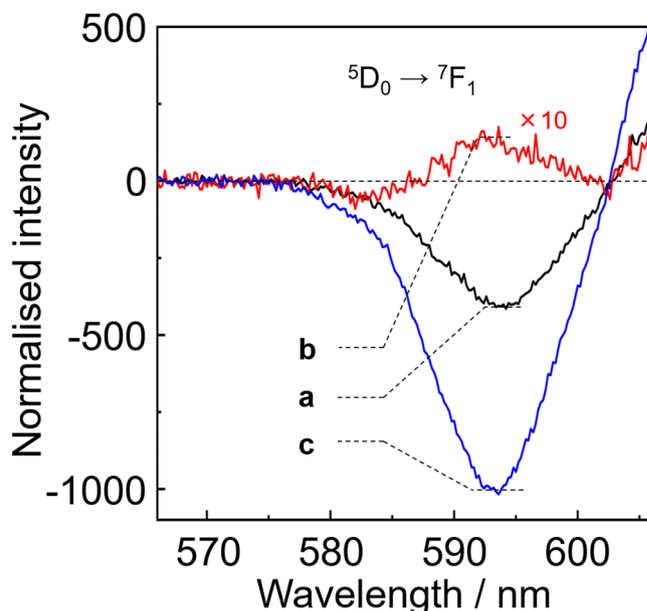


Figure 5. CPL spectra of (a) **Eu(+)-Ex0** (1×10^{-3} M, black), (b) **Eu(+)-Ex498** (1×10^{-3} M, red), and (c) **Eu(+)-Ex0** (1×10^{-5} M, blue) excited at 350 nm in acetone.

583 and 594 nm. The CPL signals at 594 nm were inverted by the addition of tppo molecules. The CPL spectrum of **Eu(+)-Ex0** (1×10^{-3} M) shows a large negative peak at 594 nm, the g_{CPL} value of which (-0.44) is similar to a previously reported g_{CPL} value ($g_{\text{CPL}} = -0.47$)³⁵ in acetone- d_6 (1×10^{-3} M). The CPL spectrum of **Eu(+)-Ex498** (1×10^{-3} M, excess amount of tppo) exhibits a small positive peak ($g_{\text{CPL}} = +0.013$). The enantiomer [**Eu(-tfc)**₃(tppo)₂] also exhibits CPL sign inversion at the transition (Supplementary Fig. S3). The CPL intensity of **Eu(+)-Ex0** in lower concentration (1×10^{-5} M, $g_{\text{CPL}} = -1.0$) is much larger than that in higher concentration (1×10^{-3} M, $g_{\text{CPL}} = -0.44$). In contrast, the CPL signals of **Eu(+)-Ex0** and **-Ex498** at around 583 nm in the ${}^5\text{D}_0 \rightarrow {}^7\text{F}_1$ transition exhibit negative CPL signals in these conditions. The CPL sign inversion behaviours depending on the external environments are summarised in Supplementary Fig. S4. Law and Dai reported similar CPL sign inversion phenomena of chiral **Eu(III)** complexes in the ${}^5\text{D}_0 \rightarrow {}^7\text{F}_1$ transition depending on solvents (Supplementary Fig. S4b)³⁹.

Considering the presence of τ_1 and τ_2 components in the emission lifetime measurements, the observed CPL spectra of **Eu(+)-Ex0** and **-Ex498** in acetone (1×10^{-3} M) were attributed to several equilibrium states of the **Eu(III)** complex in acetone. The large negative g_{CPL} of **Eu(+)-Ex0** is dominated by the τ_1 component related to coordinating acetone molecules (Supplementary Figs S5–S7). The small positive g_{CPL} of **Eu(+)-Ex498** is related to the τ_2 component of the eight-coordinated **Eu(III)** complex with two inner tppo ligands; this was supported by the similar positive g_{CPL} ($+0.036$) in toluene (Supplementary Fig. S8).

Discussion

The dissymmetry factor g_{CPL} is expressed in terms of the transition electric dipole moment $\vec{\mu}$ and transition magnetic dipole moment \vec{m} as follows²⁸,

$$g_{\text{CPL}} = 4 \cdot \frac{|\vec{\mu}| |\vec{m}| \cos \theta}{|\vec{\mu}|^2 + |\vec{m}|^2} = 4 \cdot \frac{\left(\frac{|\mu|}{|\vec{\mu}|}\right) \cos \theta}{\left(\frac{|\mu|}{|\vec{\mu}|}\right)^2 + 1} \quad (1)$$

where θ is the angle between $\vec{\mu}$ and \vec{m} . When $\vec{\mu} = \vec{m}$ ($\theta = 0^\circ$), equation (1) provides the largest g_{CPL} value ($=2$) mathematically (Fig. 6a). In the region $|\vec{\mu}|/|\vec{m}| < 1$ (Fig. 6a,b, orange regions), the **Eu(III)** complex with a large $|\vec{\mu}|$ provides a large g_{CPL} value. In general, the $|\vec{m}|$ value in the ${}^5\text{D}_0 \rightarrow {}^7\text{F}_1$ transition is larger than the $|\vec{\mu}|$ value ($|\vec{\mu}|/|\vec{m}| < 1$)⁴⁰. The intensity of $|\vec{\mu}|$ in the ${}^5\text{D}_0 \rightarrow {}^7\text{F}_1$ transition depends on the crystal field around the **Eu(III)** ion^{41,42}. The ${}^7\text{F}_1$ energy level of the **Eu(III)** ion in a typical eight-coordinate structure (C_{4v} or D_{2d}) splits into two Stark sublevels (Fig. 6b)³⁸. The two bands at 583 and 594 nm in the CPL spectra are assigned to the $A_1 \rightarrow A_2$ and $A_1 \rightarrow E$ transitions⁴², respectively, in Fig. 6b,c. The observed CPL signal in the $A_1 \rightarrow E$ transition was inverted from minus to plus, while that in the $A_1 \rightarrow A_2$ transition retained the minus sign. In C_{4v} or D_{2d} symmetry, the direct product $A_2 (=A_1 \times A_2)$ is expressed in terms of the electric dipole (ED) forbidden and magnetic dipole (MD) allowed transitions (R_2) on the character table in group theory (Fig. 6c, Supplementary Tables S3 and S4). On the other hand, the direct product $E (=A_1 \times E)$ produces ED and MD allowed transitions ((x, y) ; (R_x, R_y), Fig. 6c, Supplementary Tables S3 and S4). The CPL sign at 583 nm (ED forbidden $A_1 \rightarrow A_2$ transition) reflects the intrinsic Λ - or Δ -type structure, because of insensitive electronic state mixing. Considering the same CPL sign at 583 nm

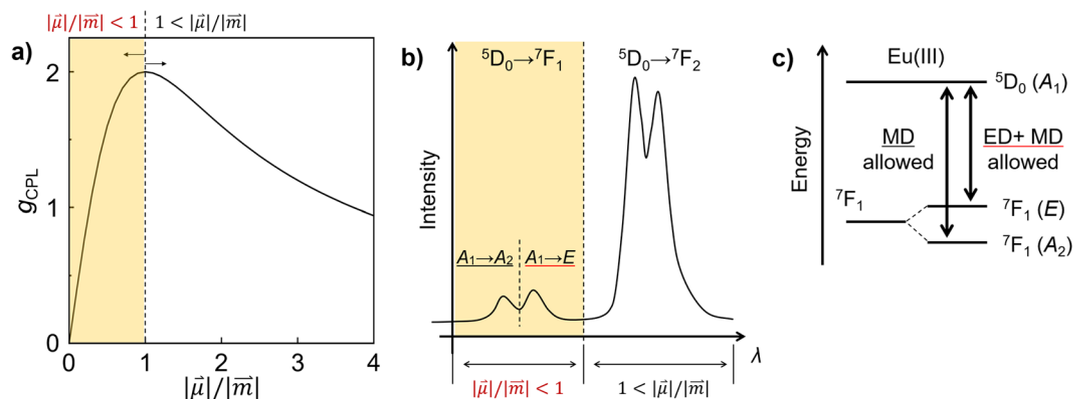


Figure 6. (a) A simulated g_{CPL} curve against $|\vec{\mu}|/|\vec{m}|$ ratio in equation (1) with $\theta = 0^\circ$. (b) An attribution of photoluminescence spectrum and (c) an energy diagram in the ${}^5\text{D}_0 \rightarrow {}^7\text{F}_1$ transition of Eu(III) complex (C_{4v} or D_{2d}) based on the electronic structure and group theory.

of the τ_1 and τ_2 components, the structure-type (Λ or Δ) with the τ_1 component is the same as that with the τ_2 component in our experiments. The coordination symmetry of τ_1 component was much similar to that of τ_2 component, which was supported by magnetic circular dichroism (MCD) measurements and DFT calculation (Supplementary Figs S9–S11). In contrast, the CPL sign in the ED and MD allowed $A_1 \rightarrow E$ transition is sensitive to electronic state mixing even for the same chiral structure-type (Λ or Δ). The effective sign inversion and drastic intensity change of the CPL signal in the $A_1 \rightarrow E$ transition should be caused by the change of $\vec{\mu}$ based on electronic state mixing.

In the ${}^5\text{D}_0 \rightarrow {}^7\text{F}_1$ transition, $\vec{\mu}$ is mainly altered by the J -mixing of ${}^7\text{F}_2$ or ${}^7\text{F}_3$ sublevels into ${}^7\text{F}_1$ ^{38,43}. In the photoluminescence spectra, the Stark splitting energy of the τ_1 component (270 cm^{-1} , Fig. 3 inset, blue; c) was larger than that of the τ_2 component (160 cm^{-1} , Fig. 3 inset, red; b). The large Stark splitting energy suggests the large J -mixing in the $A_1 \rightarrow E$ transition of the τ_1 component⁴³. The J -mixing increases the $|\vec{\mu}|$ value at the ED allowed $A_1 \rightarrow E$ transition relative to that at the ED forbidden $A_1 \rightarrow A_2$ transition, which is consistent with relatively large emission intensity at the $A_1 \rightarrow E$ transition (593.5 nm) of the τ_1 component (Fig. 3 inset, blue; c). The $|\vec{\mu}|$ increase leads to the large g_{CPL} value in equation (1).

The angle θ between $\vec{\mu}$ and \vec{m} of the τ_1 component is larger than 90° , whereas that of the τ_2 component is smaller than 90° , suggesting that the angle is established by $\vec{\mu}$ vector change due to J -mixing. The extra-large enhancement of g_{CPL} from +0.013 to -1.0 also indicates that J -mixing promotes the direction of $\vec{\mu}$ and \vec{m} to antiparallel, leading to the large g_{CPL} . We demonstrated that the CPL sign and intensity are strongly influenced by the chiral electronic structure depending on the $\vec{\mu}$ under J -mixing in the same chiral structure-type.

Conclusion

We successfully observed a CPL sign inversion with a drastic g_{CPL} change from +0.013 to -1.0 for the Eu(III) complex with the same chiral coordination structure-type. The CPL phenomena were attributed to the chiral electronic structure depending on the $\vec{\mu}$ under J -mixing. We also achieved an extra-large g_{CPL} (-1.3) of the Eu(III) complex with chiral +tfc ligands in DMSO, suggesting that the g_{CPL} of the Eu(III) complex could be enhanced by J -mixing with a small 4f-5d mixing character (Supplementary Figs S12 and S13, Table S5). The results provide a novel aspect for the optical activity of metal complexes and molecular design of chiral lanthanide complex by maximising the g_{CPL} value.

Methods

Synthesis of Tris(3-trifluoroacetyl-(+)-camphorato)europium(III) bis(triphenylphosphine oxide) ([Eu(+tfc)₃(tppo)₂]). Europium(III) acetate n -hydrate (0.36 g) was dissolved in distilled water (150 mL), and a few drops of 28% ammonia solution were added. (+)-3-trifluoroacetyl camphor (+tfc, 0.50 g, 2.0 mmol) in methanol (20 mL) was added to the solution, and the mixture was stirred for 3 h at room temperature. The obtained powder was washed with distilled water, and the powder was dried in vacuo (0.42 g). The powder (0.19 g) and triphenylphosphine oxide (tppo, 0.11 g, 0.40 mmol) were dissolved in methanol (20 mL) and refluxed for 3 h. The reaction solution was evaporated using a rotary evaporator. The obtained powder was recrystallised from a hot acetonitrile solution and gave yellow crystals. Yield: 47%. Elemental analysis: Calculated for $\text{C}_{72}\text{H}_{72}\text{EuF}_9\text{O}_8\text{P}_2$: C, 59.63%, H, 5.00%. Found: C, 59.54%, H, 4.92%³⁵.

Synthesis of Tris(3-trifluoroacetyl(-)-camphorato)europium(III) bis(triphenylphosphine oxide) ([Eu(-tfc)₃(tppo)₂]). [Eu(-tfc)₃(tppo)₂] was prepared using the same method for [Eu(+tfc)₃(tppo)₂], starting from (-)-3-trifluoroacetyl camphor, yielding yellow powder. Yield: 64%. Elemental analysis: Calculated for $\text{C}_{72}\text{H}_{72}\text{EuF}_9\text{O}_8\text{P}_2$: C, 59.63%, H, 5.00%. Found: C, 59.49%, H, 4.94%.

Materials

Europium(III) acetate *n*-hydrate, acetone- d_6 (99.9%), acetone (spectroscopic grade), toluene (spectroscopic grade), dimethyl sulfoxide (spectroscopic grade), and 28% ammonia solution were purchased from Wako Pure Chemical Industries Ltd. (+)-3-(trifluoroacetyl)camphor and (–)-3-(trifluoroacetyl)camphor were purchased from Sigma-Aldrich Co. Triphenylphosphine oxide was purchased from Tokyo Chemical Industry Co., Ltd. All other chemicals and solvents were of reagent grade and were used without further purification.

Apparatus. Elemental analyses were performed on an Exeter Analytical CE440. Proton nuclear magnetic resonance (^1H NMR) spectra were recorded in acetone- d_6 on an auto-NMR JEOL ECS 400 MHz; Acetone ($\delta_{\text{H}} = 2.05$ ppm) was used as an internal reference. Emission spectra and emission lifetimes were measured using a Horiba/Jobin-Yvon FluoroLog-3 spectrofluorometer. CPL spectra were measured using a JASCO CPL-200 spectrofluoropolarimeter.

References

- Hembury, G. A., Borovkov, V. V. & Inoue, Y. Chirality-sensing supramolecular systems. *Chem. Rev.* **108**, 1–73 (2008).
- Pérez-García, L. & Amabilino, D. B. Spontaneous resolution under supramolecular control. *Chem. Soc. Rev.* **31**, 342–356 (2002).
- Yoon, T. P. & Jacobsen, E. N. Privileged chiral catalysts. *Science* **299**, 1691–1693 (2003).
- Saville, C. K. *et al.* Biocatalytic asymmetric synthesis of sitagliptin manufacture. *Science* **329**, 305–310 (2010).
- Noyori, R., Kitamura, M. & Ohkuma, T. Toward efficient asymmetric hydrogenation: Architectural and functional engineering of chiral molecular catalysts. *Proc. Natl. Acad. Sci.* **101**, 5356–5362 (2004).
- Mimura, Y. *et al.* Solvent-sensitive sign inversion of excimer origin circularly polarized luminescence in bipyrrenyl peptides. *ChemistrySelect* **2**, 7759–7764 (2017).
- Yao, J. *et al.* Temperature-driven planar chirality switching of a pillar[5]arene-based molecular universal joint. *Angew. Chem. Int. Ed.* **56**, 6869–6873 (2017).
- Kim, J. *et al.* Induction and control of supramolecular chirality by light in self-assembled helical nanostructures. *Nat. Commun.* **6**, 1–8 (2015).
- Lin, R. *et al.* pH-switchable inversion of the metal-centered chirality of metallabenzenes: Opposite stereodynamics in reactions of ruthenabenzene with L- and D-cysteine. *Chem. Eur. J.* **17**, 2420–2427 (2011).
- Choi, J. & Majima, T. Conformational changes of non-B DNA. *Chem. Soc. Rev.* **40**, 5893–5909 (2011).
- Zhang, L., Qin, L., Wang, X., Cao, H. & Liu, M. Supramolecular Chirality in Self-Assembled Soft Materials: Regulation of Chiral Nanostructures and Chiral Functions. *Adv. Mater.* **26**, 6959–6964 (2014).
- Mateos-Timoneda, M. A., Crego-Calama, M. & Reinhoudt, D. N. Supramolecular chirality of self-assembled systems in solution. *Chem. Soc. Rev.* **33**, 363–372 (2004).
- Katsonis, N., Lacaze, E. & Ferrarini, A. Controlling chirality with helix inversion in cholesteric liquid crystals. *J. Mater. Chem.* **22**, 7088–7097 (2012).
- Pieraccini, S., Masiero, S., Ferrarini, A. & Piero Spada, G. Chirality transfer across length-scales in nematic liquid crystals: Fundamentals and applications. *Chem. Soc. Rev.* **40**, 258–271 (2011).
- Mason, S. F. *Molecular optical activity and the chiral discriminations*. (Cambridge University Press, 1982).
- Hattori, S., Vandendriessche, S., Koeckelberghs, G., Verbiest, T. & Ishii, K. Evaporation rate-based selection of supramolecular chirality. *Chem. Commun.* **53**, 3066–3069 (2017).
- Kumar, M. *et al.* A dynamic supramolecular polymer with stimuli-responsive handedness for *in situ* probing of enzymatic ATP hydrolysis. *Nat. Commun.* **5**, 1–8 (2014).
- Satrijo, A., Meskers, S. C. J. & Swager, T. M. Probing a conjugated polymer's transfer of organization-dependent properties from solutions to films. *J. Am. Chem. Soc.* **128**, 9030–9031 (2006).
- Okazaki, M. *et al.* Solvent-controlled sign inversion of circularly polarized luminescent binaphthylacetic acid derivative. *J. Photochem. Photobiol. A* **331**, 115–119 (2016).
- Kimoto, T. *et al.* Control of solid-state circularly polarized luminescence of binaphthyl organic fluorophores through environmental changes. *Asian J. Org. Chem.* **2**, 404–410 (2013).
- Suzuki, N., Fujiki, M., Kimpinde-Kalunga, R. & Koe, J. R. Chiroptical inversion in helical Si-Si bond polymer aggregates. *J. Am. Chem. Soc.* **135**, 13073–13079 (2013).
- Guo, S., Suzuki, N. & Fujiki, M. Oligo- and polyfluorenes meet cellulose alkyl esters: Retention, inversion, and racemization of circularly polarized luminescence (CPL) and circular dichroism (CD) via intermolecular C-H/O=C interactions. *Macromolecules* **50**, 1778–1789 (2017).
- Nishikawa, T., Nagata, Y. & Sugimoto, M. Poly(quininoxaline-2,3-diyl) as a multifunctional chiral scaffold for circularly polarized luminescent materials: Color tuning, energy transfer, and switching of the CPL handedness. *ACS Macro Lett.* **6**, 431–435 (2017).
- Sheng, Y. *et al.* Reversal circularly polarized luminescence of AIE-active chiral binaphthyl molecules from solution to aggregation. *Chem. Eur. J.* **21**, 13196–13200 (2015).
- Hutin, M. & Nitschke, J. Solvent-tunable inversion of chirality transfer from carbon to copper. *Chem. Commun.* **2**, 1724–1726 (2006).
- Bünzli, J.-C. G. On the design of highly luminescent lanthanide complexes. *Coord. Chem. Rev.* **293–294**, 19–47 (2015).
- Richardson, F. Selection rules for lanthanide optical activity. *Inorg. Chem.* **19**, 2806–2812 (1980).
- Francesco, Z. & Di Bari, L. Lanthanide circularly polarized luminescence: Bases and applications. *Chirality* **27**, 1–13 (2015).
- Lunkley, J. L., Shirota, D., Yamanari, K., Kaizaki, S. & Muller, G. Chiroptical spectra of a series of tetrakis((+)-3-heptafluorobutylrylcamphorato)lanthanide(III) with an encapsulated alkali metal ion: circularly polarized luminescence and absolute chiral structures for the Eu(III) and Sm(III) complexes. *Inorg. Chem.* **50**, 12724–12732 (2011).
- Kumar, J., Marydasan, B., Nakashima, T., Kawai, T. & Yuasa, J. Chiral supramolecular polymerization leading to eye differentiable circular polarization in luminescence. *Chem. Commun.* **52**, 9885–9888 (2016).
- Carr, R., Evans, N. H. & Parker, D. Lanthanide complexes as chiral probes exploiting circularly polarized luminescence. *Chem. Soc. Rev.* **41**, 7673–7686 (2012).
- Kono, Y., Hara, N., Shizuma, M., Fujiki, M. & Imai, Y. Complexes of Eu(III)(hfa)₃ with a planar chiral P(III) ligand (Phanephos): solvent-sensitive sign inversion of circularly polarized luminescence. *Dalton Trans.* **46**, 5170–5174 (2017).
- Yuasa, J., Ueno, H. & Kawai, T. Sign reversal of a large circularly polarized luminescence signal by the twisting motion of a bidentate ligand. *Chem. Eur. J.* **20**, 8621–8627 (2014).
- Smith, D. G., Pal, R. & Parker, D. Measuring equilibrium bicarbonate concentrations directly in cellular mitochondria and in human serum using europium/terbium emission intensity ratios. *Chem. Eur. J.* **18**, 11604–11613 (2012).
- Harada, T. *et al.* Circularly polarized luminescence of Eu(III) complexes with point- and axis-chiral ligands dependent on coordination structures. *Inorg. Chem.* **48**, 11242–11250 (2009).
- Kleckner, I. R. & Foster, M. P. An introduction to NMR based approaches for measuring protein dynamics. *Biochim. Biophys. Acta* **1814**, 942–968 (2011).

37. Cockerill, A. F., Davies, G. L. O., Harden, R. C. & Rackham, D. M. Lanthanide shift reagents in nuclear magnetic resonance spectroscopy. *Chem. Rev.* **73**, 553–588 (1976).
38. Binnemans, K. Interpretation of europium(III) spectra. *Coord. Chem. Rev.* **295**, 1–45 (2015).
39. Dai, L., Lo, W. S., Coates, I. D., Pal, R. & Law, G. L. New class of bright and highly stable chiral cyclen europium complexes for circularly polarized luminescence applications. *Inorg. Chem.* **55**, 9065–9070 (2016).
40. Richardson, F. S., Berry, M. T. & Reid, M. F. Ligand polarization contributions to lanthanide $4f \rightarrow 4f$ magnetic dipole transition moments and rotatory strengths. *Mol. Phys.* **58**, 929–945 (1986).
41. Görrler-Walrand, C., Fluyt, L., Ceulemans, A. & Carnall, W. T. Magnetic dipole transitions as standards for Judd-Ofelt parametrization in lanthanide spectra. *J. Chem. Phys.* **95**, 3099–3106 (1991).
42. Forsberg, J. H. Complexes of lanthanide (III) ions with nitrogen donor ligands. *Coord. Chem. Rev.* **10**, 195–226 (1973).
43. Ma, C. G., Brik, M. G., Kiisk, V., Kangur, T. & Sildos, I. Spectroscopic and crystal-field analysis of energy levels of Eu^{3+} in SnO_2 in comparison with ZrO_2 and TiO_2 . *J. Alloys Compd.* **509**, 3441–3451 (2011).

Acknowledgements

We thank professor K. Monde and assistant professor T. Taniguchi in Hokkaido University for the discussion of spectroscopic measurements. This work was partly supported by JSPS KAKENHI Grant Number JP17H06375. This work was also supported by Grants-in-Aid for Scientific Research (no. 18H02041, no. 18H04497), Young Scientists (B) (no. 17K1446707), and JSPS Research Fellow (no. 16J01325). S. Wada was also supported by The Ministry of Education, Culture, Sports Science and Technology through Program for Leading Graduate Schools (Hokkaido University 'Ambitious Leader's Program').

Author Contributions

S.W. performed all the syntheses and measurements, and wrote the paper under the supervision of Y.K., T.N., K.F. and Y.H., M.G., K.T. and Y.C. supported the CPL measurements. All authors reviewed the manuscript.

Additional Information

Supplementary information accompanies this paper at <https://doi.org/10.1038/s41598-018-34790-0>.

Competing Interests: The authors declare no competing interests.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2018