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Bright near-infrared circularly polarized electrochemiluminescence from Au₉Ag₄ nanoclusters†

Metal nanoclusters are excellent electrochemiluminescent luminophores owing to their rich electrochemical and optical properties. However, the optical activity of their electrochemiluminescence (ECL) is unknown. Herein, we achieved, for the first time, the integration of optical activity and ECL, *i.e.*, circularly polarized electrochemiluminescence (CPECL), in a pair of chiral Au_9Ag_4 metal nanocluster enantiomers. Chiral ligand induction and alloying were employed to endow the racemic nanoclusters with chirality and photoelectrochemical reactivity. $S-Au_9Ag_4$ and $R-Au_9Ag_4$ exhibited chirality and brighted emission (quantum yield = 4.2%) in the ground and excited states. The enantiomers showed mirrorimaged CPECL signals at 805 nm owing to their highly intense and stable ECL emission in the presence of tripropylamine as a co-reactant. The ECL dissymmetry factor of the enantiomers at 805 nm was calculated to be $\pm 3 \times 10^{-3}$, which is comparable with that obtained from their photoluminescence. The obtained nanocluster CPECL platform shows the discrimination of chiral 2-chloropropionic acid. The integration of optical activity and ECL in metal nanoclusters provides the opportunity to achieve enantiomer discrimination and local chirality detection with high sensitivity and contrast.

Electrochemiluminescence (ECL), a property that combines electrochemistry and chemiluminescence (CL), may be observed when electroactive species are reduced or oxidized and the generated redox species undergoes chemical reactions to produce excited luminophores that emit light.^{1,2} ECL is a versatile analytical, immunological, and imaging method owing to its advantages of enhanced temporal and spatial controllability compared with CL, as well as lower background interference higher signal-to-noise ratio compared photoluminescence.3-5 The long-term goals of researchers in this field include the development of novel luminophores,6 exploration of new ECL mechanisms,7-13 and exploitation of new detection14-16 and imaging17-20 platforms. ECL spectroscopy to determine the emission intensity and spectra of luminophores has been applied in emission mechanism studies and practical applications to sense trace amounts of target molecules. 21-23

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Circularly polarized luminescence (CPL) spectroscopy is an analytical technique that provides important information about chiral materials in the excited state.²⁴⁻²⁶ Chiral materials demonstrating CPL have unique applications in displays,²⁷ polarized lasers,²⁸ chiral sensing,²⁹⁻³¹ and optical imaging.³² The combination of CPL and ECL in a single analytical technique, that is, circularly polarized electrochemiluminescence (CPECL) spectroscopy, is expected to integrate the merits of CPL and ECL to achieve chiral detection with high sensitivity and contrast and low background interference. Although pyrene-18C6 and [Ru(phen)₃][ClO₄]₂ complexes have been reported to exhibit CPECL performance,^{33,34} studies on the CPECL of luminophores remain limited.

Metal nanoclusters protected by ligands have received wide attention owing to their definite structure, intense quantum-size effect, and rich physicochemical properties, including chiral, electrochemical, and optical properties.^{35–42} Metal nanoclusters have been demonstrated to be excellent ECL luminophores.^{23,43,44} The ECLs of Au₂₅, Au₃₈, and Au₁₄₄ nanoclusters in the co-reactant pathway were previously reported by Ding and coworkers.^{45–47} We recently demonstrated the intense ECL of rod-shaped bimetallic Au₁₂Ag₁₃ nanoclusters *via* the self-annihilation and co-reactant pathways.⁴⁸ However, the achiral or racemic properties of these materials limit research on the CPECL of metal nanoclusters. Thus, whether the ECL produced by chiral nanoclusters is circularly polarized and whether

CPECL could be generalized and detected in metal nanoclusters are unknown. These issues may be attributed to the difficulty of simultaneously achieving chiral structures, intense emission, and good electrochemical performance.

Herein, we constructed metal nanocluster enantiomers with chirality and strong emission, namely, S-Au₉Ag₄ and R-Au₉Ag₄ (S-Au₉Ag₄ = [Au₉Ag₄(S-DIOP)₄Cl₄]Cl; R-Au₉Ag₄ = [Au₉Ag₄(R-DIOP)₄Cl₄]Cl; S/R-DIOP = 1,4-bis(diphenylphosphino)-2,3-o-isopropylidene-2,3-butanediol) to study the optical activity of their ECL. Intense red-emissive and chiral S-Au₉Ag₄ and R-Au₉Ag₄ nanoclusters were constructed by alloying and chiral ligand induction, and the optical and electrochemical performance of the resultant molecules was investigated. A pair of mirrored CPECL peaks at 805 nm was observed in the oxidative reduction pathway of S-Au₉Ag₄ and R-Au₉Ag₄ when tripropylamine (TPrA) was used as a co-reactant. The CPECL dissymmetry factor of the nanoclusters was calculated to be g_{ECL} = \pm 5 \times 10⁻³, which is comparable with the g_{PL} (\pm 3 \times 10⁻³) obtained from their PL.

The chirality of nanoclusters usually originates from the inherent chirality of the metal core, chiral arrangement of motifs, and chiral ligands.⁴² The introduction of chiral ligands endows achiral or racemic nanoclusters with optical activity.⁴⁹⁻⁵¹ Alloying can improve the PL of nanoclusters.^{52,53} In this study, racemic Au₉Ag₄ protected by bisdiphenylphosphinobutane (DPPB) was first prepared, and its composition was characterized by ESI-MS, XPS (Fig. S1†), ¹H NMR (Fig. S2†), ³¹P NMR (Fig. S3†), and TGA (Fig. S4†). The molecular formula of the Au₉Ag₄ nanocluster was determined to be [Au₉Ag₄(DPPB)₄Cl₄][†] by ESI-MS analysis (Fig. 1b, black line). The structure of [Au₉Ag₄(DPPB)₄Cl₄]Cl was determined using single-crystal XRD, as shown in the inset of Fig. 1a and detailed in Table S1.† A pair of

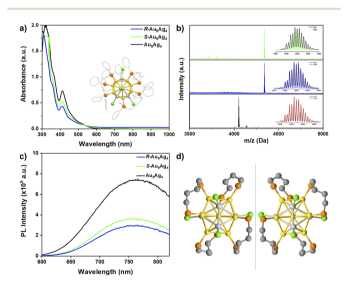


Fig. 1 (a) UV-vis absorption spectra of Au_9Ag_4 (black), R- Au_9Ag_4 (blue), and S- Au_9Ag_4 (green). The inset shows the structure of a Au_9Ag_4 nanocluster. (b) ESI-MS spectra of Au_9Ag_4 (black), R- Au_9Ag_4 (blue), and S- Au_9Ag_4 (green). (c) PL spectra of Au_9Ag_4 (black), R- Au_9Ag_4 (blue), and S- Au_9Ag_4 (green). (d) Au_9Ag_4 enantiomer pairs. Color label: gold = Au_4 , white = Au_4 , green = Cl_4 , orange = P_4 , gray = Cl_4 ; all benzene rings were omitted for clarity.

enantiomers was observed in a unit cell (Fig. 1d), indicating that the nanoclusters are racemic. The chirality of the nanoclusters originated from the chiral arrangement of the protecting motif. The chiral ligand S/R-DIOP was used to obtain optically pure Au₉Ag₄ nanoclusters, and chiral S-Au₉Ag₄ and R-Au₉Ag₄ nanoclusters were prepared. The ESI-MS spectra of S-Au₉Ag₄ and R-Au₉Ag₄ showed peaks at 4339.9, consistent with the formula [Au₉Ag₄(DIOP)₄Cl₄]⁺ (Fig. 1b, blue and green lines). The UV-vis spectra of S-Au₉Ag₄ and R-Au₉Ag₄ were similar to that of [Au₉-Ag₄(DPPB)₄Cl₄]Cl, indicating the similarity of their structures (Fig. 1a). XPS further confirm the composition of [Au₉Ag₄(-DIOP)₄Cl₄]⁺ (Fig. S5†). The PL spectra of Au₉Ag₄, S-Au₉Ag₄, and R-Au₉Ag₄ were centered at 760 nm, and their lifetimes were 0.21, 0.80, and 0.94 μs, respectively (Fig. S6†). The PL quantum yields of Au₉Ag₄, S-Au₉Ag₄, and R-Au₉Ag₄ were 2.96%, 3.56% and 4.20%, respectively.

The circular dichroism (CD) and CPL spectra of S-Au₉Ag₄ and R-Au₉Ag₄ were investigated to determine their optical activities. As shown in Fig. 2a, the CD spectra of S-Au₉Ag₄ and R-Au₉Ag₄ are mirror-imaged and show main peaks at 325, 360, and 375 nm, minor peaks at 345, 420, 460, and 490 nm, and a shoulder peak at 550 nm. These results reflect the groundstate chirality of S-Au₉Ag₄ and R-Au₉Ag₄. The concentrationindependent anisotropy factor $(g_{abs} = 2(A_L - A_R)/(A_L + A_R))$ of the nanoclusters was calculated from their CD spectra (Fig. 2b) and found to be 2.8×10^{-3} at 600 nm. To understand the optical activity of S-Au₉Ag₄ and R-Au₉Ag₄ in the excited state further, their CPL spectra were recorded. As shown in Fig. 2c, S-Au₉Ag₄ and R-Au₉Ag₄ manifested strong CPL activity at \sim 760 nm, with negative and positive signals, respectively. The anisotropy factors (g-factor) of S/R-Au₉Ag₄ were calculated to be $g_{\rm PL}(R) = 2.84 \times 10^{-3}$ and $g_{\rm PL}(S) = -2.33 \times 10^{-3}$, respectively.

S-Au₉Ag₄ and R-Au₉Ag₄ were subjected to cyclic voltammetry (CV) to determine the appropriate potential for ECL generation.

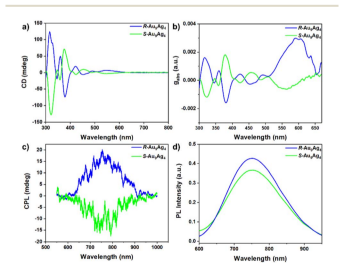


Fig. 2 (a) CD spectra of R-Au₉Ag₄ (blue) and S-Au₉Ag₄ (green). (b) Anisotropy factors (g-factors) of R-Au₉Ag₄ (blue) and S-Au₉Ag₄ (green) nanoclusters. (c) CPL spectra of R-Au₉Ag₄ (blue) and S-Au₉Ag₄ (green). (d) Corresponding PL spectra of R-Au₉Ag₄ (blue) and S-Au₉Ag₄ (green) during CPL detection. R-Au₉Ag₄ and S-Au₉Ag₄ nanoclusters were dissolved in DCM for the CD and CPL experiments.

Experiments were performed in dichloromethane (DCM) with a glassy carbon working electrode. As shown in Fig. 3a, S-Au₉Ag₄ and R-Au₉Ag₄ exhibit similar voltammetric features, which indicates that they have the same electronic state. Two-step reduction bands centered at $E_{\rm R1}=-0.819$ V and $E_{\rm R2}=-1.432$ V and a one-step oxidation band centered at $E_{\rm O1}=1.106$ V were also observed. The reduction currents in the first and second reduction steps were identical, and the oxidation current was two times greater than the reduction current at each step. Thus, two electrons were injected to generate ${\rm Au_9Ag_4}^0$ and ${\rm Au_9Ag_4}^1$ during reduction scanning, and two electrons were removed to generate ${\rm Au_9Ag_4}^{3+}$ during oxidation scanning in the voltammetry experiment. The electrochemical gaps were calculated to be 1.925 V, which is consistent with their optical gaps 1.952 eV (Fig. S7†).

The ECL performance of the samples in the co-reactant pathway was explored because luminophores usually present relatively strong and stable signals in the presence of a co-reactant. Fig. 3b and c show the ECLs of S-Au₉Ag₄ and R-

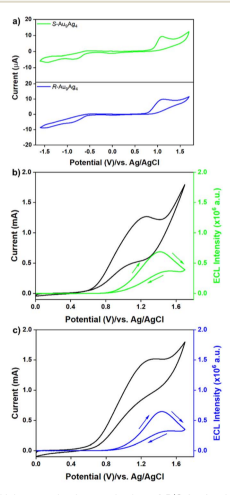


Fig. 3 (a) Voltammetric characterization of R/S-Au₉Ag₄ in degassed DCM with 0.1 M tetrabutylammonium perchlorate (TBAP) as the supporting electrolyte. (b) and (c) CV (black curve) and corresponding ECL signal (blue/green curves) recorded in DCM containing 0.23 mM R/S-Au₉Ag₄, 700 mM TPrA, and 0.1 M TBAP. A platinum mesh electrode and Ag/AgCl wire were used as the working and reference electrodes, respectively. Scan rate $= 0.1 \text{ V s}^{-1}$.

Au₉Ag₄ with TPrA as the co-reactant when the potential was scanned from 0 to 1.7 V. Both nanoclusters displayed similar ECL profiles. TPrA oxidation occurs at 0.7 V, but the ECL signals were observed at an onset potential of 0.9 V, thus implying that ECL formation requires the oxidation of both the luminophore and co-reactant by the electrode reactions. Step-ECL has been demonstrated to generate more durable and stable signals than CV-ECL, which is highly desirable for ECL optical activity testing.48 S-Au₉Ag₄ and R-Au₉Ag₄ displayed weaker ECL signals in the presence of benzoyl peroxide (BPO) as a co-reactant compared with TPrA (Fig. S8†). The ECL of the nanoclusters in the oxidative reduction pathway was explored to select a suitable co-reactant that could generate strong ECL signals and achieve high contrast. The concentration of the TPrA coreactant was optimized in step ECL experiments to obtain higher and more stable signals (Fig. S9 and S10†). The intensity of the step ECL signals of S-Au₉Ag₄ and R-Au₉Ag₄ increased with increasing co-reactant concentration, peaked at 700 mM, and remained stable thereafter.

Because the S-Au₉Ag₄ and R-Au₉Ag₄ nanoclusters exhibited similar ECL behaviors, we selected S-Au₉Ag₄ as an example and explored its ECL spectra in the presence of different concentrations of TPrA. The ECL spectra of S-Au₉Ag₄ and R-Au₉Ag₄ under a holding potential of 0-1.7 V applied cyclically every 5 s were determined. The exposure time was set to 5 s. One spectrum was obtained from each nanocluster at a holding potential of 1.7 V. The ECL spectrum of S-Au₉Ag₄ shifted when different amounts of TPrA were used. As shown in Fig. S12 (right panel),† in 50 mM TPrA, the peak position of the ECL spectrum can be observed at 760 nm, which is consistent with the PL spectral position observed in Fig. 1c. As the TPrA concentration increased, the ECL spectrum of S-Au₉Ag₄ red-shifted. When the concentration of TPrA was 700 mM, the ECL emission peak of S-Au₉Ag₄ was detected at 805 nm. To assign the ECL spectra, we collected the PL spectra of S-Au₉Ag₄ electrolyzed at different potentials. According to the reduction peaks of S-Au₉Ag₄ (Fig. S11†), we collected the PL spectra of S-Au₉Ag₄ electrolyzed at -1.2, -1.6, and -2.4 V. As shown in the left panel in Fig. S12,† electrolyzed S-Au₉Ag₄ showed peaks at 780, 785, and 795 nm, corresponding to the emissions of $(S-Au_9Ag_4)^0$, $(S-Au_9Ag_4)^0$ Au_9Ag_4)¹⁻, and $(S-Au_9Ag_4)^{x-}$ ($x \ge 2$), respectively. The emission intensity of the nanoclusters decreases as the potential reaches more negative potentials (Fig. S13†). Although we attempted to excite only the electrolyzed nanoclusters near the electrode and collect their PL spectrum, the excitation light passing through the sample cell inevitably excited a few non-electrolyzed nanoclusters. Thus, the PL spectrum collected under electrolysis is actually the sum of the PL spectra of the electrolyzed sample and a small amount of the unelectrolyzed sample. That is, the PL spectra of $(S-Au_9Ag_4)^0$, $(S-Au_9Ag_4)^{1-}$, $(S-Au_9Ag_4)^{x-}$ may be more red-shifted than shown in the left panel in Fig. S12.† The ECL spectral peaks centered at 780, 790, and 805 nm can be respectively assigned to the ECL emissions of $(S-Au_9Ag_4)^0$, $(S-Au_9Ag_4)^0$ Au₉Ag₄)¹⁻, combinations of clusters of multiple valences (S- $\operatorname{Au}_{9}\operatorname{Ag}_{4}^{x-}$ ($x \ge 2$). Similar ECL emission peak shifts have been observed in Au₂₅ nanoclusters. 23,54 The possible mechanism and major reaction processes of S-Au₉Ag₄ nanoclusters are

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presented in the ESI (Schemes S1–S4†). TPrA is oxidized by the electrode to produce strongly reductive TPrA radical after deprotonation. When a more positive potential is applied, the nanoclusters are oxidized. The strongly reductive TPrA reduces the nanoclusters to generate an excited-state nanoclusters that relaxes to the ground state and produces ECLs.

The optical activity of the ECL emissions of S-Au₉Ag₄ and R-Au₉Ag₄ was explored in two ways. In the first method, a commercial CPL instrument was used (Fig. S14†). It is used to collect the circularly polarized emissions, and the sample is excited externally under the condition of constant potential electrolysis. This method is simple but has high detection limits because it works with PL systems that typically have a stronger light signal than ECL systems. In the second method, a CCD with left- and right-handed circular polarizers applied to the optical path was used. Such method has higher sensitivity and detection limit due to the high sensitivity of the instrument for detection of weak light. As shown in Fig. S15 and S16,† the ECL spectra of S-Au₉Ag₄ and R-Au₉Ag₄ are centered at 805 nm. The CPECL spectra of S-Au₉Ag₄ and R-Au₉Ag₄ were collected using a commercial CPL instrument with triggering by an electrochemical reaction. As shown in Fig. 4a, a pair of mirror-imaged CPECL signals of S-Au₉Ag₄ and R-Au₉Ag₄ were observed. The ECL dissymmetric factors (g_{ECL}) of S-Au₉Ag₄ and R-Au₉Ag₄ were calculated to be $\pm 3 \times 10^{-3}$, which is comparable with the $g_{\rm PL}$ obtained from their CPL. The dissymmetry factors (g_{ECL} or g_{PL})

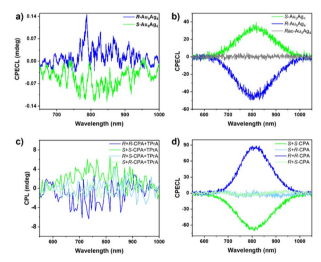


Fig. 4 (a) CPECL of *R*-Au₉Ag₄ (blue) and *S*-Au₉Ag₄ (green) collected by commercial CPL instrument. 0.23 mM *R/S*-Au₉Ag₄ dissolved in DCM were subjected to CPECL testing under 1.7 V electrolysis. (b) CPECL of *R*-Au₉Ag₄ (blue) and *S*-Au₉Ag₄ (green) collected by adding circular polarizers. ECL spectra were collected in DCM containing 0.23 mM *R/S*-Au₉Ag₄, 700 mM TPrA, and 0.1 M TBAP. A platinum mesh electrode and Ag/AgCl wire were used as the working and reference electrodes, respectively. The CPECL spectra were obtained by subtracting the ECL spectrum obtained by inserting a right-handed circular polarizer from the ECL spectrum obtained by inserting a left-handed circular polarizer. (c) CPL spectra after adding 1 equivalent *R*-CPA or *S*-CPA to *R*-Au₉Ag₄ or *S*-Au₉Ag₄ dissolved in DCM. (d) CPECL spectra after adding *R*-CPA or *S*-CPA to *R*-Au₉Ag₄. One equivalent *R*-CPA or *S*-CPA was added into *R*-Au₉Ag₄ or *S*-Au₉Ag₄, which is subjected to the CPECL experiments.

of the S-Au₉Ag₄ and R-Au₉Ag₄ nanoclusters are small, likely because of the relatively large nanocluster body and small chiral unit.

In the second detection method, circular polarizers were used as filters to demonstrate the circular polarization of ECL produced by S-Au₉Ag₄ and R-Au₉Ag₄ (Fig. S17†). In general, a left-handed polarizer converts left-handed circularly polarized light into linearly polarized light. When right-handed circularly polarized light enters the polarizer from this direction, the output is zero. The opposite output is obtained when a righthanded polarizer is applied; that is, left-handed circularly polarized light cannot pass through a right-handed circular polarizer. Two spectra can be obtained when left- and righthanded circular polarizers are added to the light path. The spectrum obtained from the difference in the spectra collected after filtration by the left- and right-handed polarizers is the circularly polarized light produced by the luminophore. The CPECL spectrum of each enantiomer was obtained by subtracting the spectrum filtered by the right-handed circular polarizer from that filtered by the left-handed circular polarizer.

The ECL spectra of S-Au₉Ag₄ and R-Au₉Ag₄ collected before and after the addition of the left/right-handed circular polarizers are shown in Fig. S15 and S16,† respectively. Six experiments were performed for each enantiomer to evaluate the reproducibility of the results. The ECL intensity decreased when the circular polarizers were added to the light path; this result may be attributed to the influence of the intrinsic transmittance of the device. Similar results were obtained in all other experiments for each enantiomer. The ECL intensity of S-Au₉Ag₄ after filtration by the left-handed circular polarizer was higher than that obtained after filtration by the right-handed circular polarizer (Fig. S15†), resulting in positive CPECL signals (Fig. 4b, green line). Negative CPECL signals were observed for R-Au₉Ag₄ (Fig. 4b, blue line). The CPECL signal of R-Au₉Ag₄ was mirror-imaged with that of S-Au₉Ag₄. Differences in the intensity of the CPECL signals of the different groups are due to differences in the angle between the light and polarizer in these groups. Interestingly S-Au₉Ag₄ showed negative signals in its CPL but positive ones in its CPECL spectrum obtained by adding the circular polarizers. By contrast, R-Au₉Ag₄ showed positive signals in its CPL spectra but negative ones in its CPECL spectra obtained by adding the circular polarizers. The contradicting signals in the CPL and CPECL spectra are due to differences in the instruments and measurement procedure. The ECL spectra of racemic rac-Au₉Ag₄ (a mixture of S-Au₉Ag₄ and R-Au₉Ag₄ at a 1:1 ratio) before and after passing through the left/right-handed circular polarizers demonstrated random features (i.e., some signals were positive, some signals were negative, and some signals were zero); thus, the averaged CPECL of racemic rac-Au₉Ag₄ showed a straight line (Fig. 4b, gray line).

In addition, discrimination against enantiomers was further investigated. Chiral carboxylic acid (*S/R*-2-chloropropanoic acid, *S/R*-CPA) were employed for the discrimination of this CPECL detection platform due to the binding ability of oxygen atoms to the silver atoms. ^{55,56} The ECL intensity of *S*-Au₉Ag₄ after filtration by the left-handed circular polarizer was lower than that

obtained after filtration by the right-handed circular polarizer after addition of S-CPA (Fig. S18†), indicating a negative CPECL signal (Fig. 4d, green line). The CPECL of S-Au₉Ag₄ is inverted after addition of S-CPA. For the R-Au₉Ag₄, the CPECL changed from an initial negative signal (Fig. 4b, blue line) to a positive signal (Fig. 4d, blue line) after adding the R-CPA (Fig. S19†). While adding the R-CPA into S-Au₉Ag₄ or adding the S-CPA into R-Au₉Ag₄, the ECL intensity showed comparable after filtration by the left-handed and right-handed circular polarizers (Fig. S20 and S21†), resulting in zero signals for CPECL (Fig. 4d, light green and light bule lines). The discrimination of nanoclusters against enantiomers was further studied using CPL. The CPL of the S-Au₉Ag₄ changed from negative (Fig. 2c, green line) to positive (Fig. 4c, green line) after adding the S-CPA and TPrA. While the CPL of the R-Au₉Ag₄ changed from positive (Fig. 2c, blue line) to negative (Fig. 4c, blue line) after adding the R-CPA and TPrA. The CPLs are zero when adding R-CPA to S-Au₉Ag₄ or adding S-CPA to R-Au₉Ag₄ (Fig. 4c, light green and light blue lines). However, addition of the chiral enantiomer or TPrA alone did not result in a change in the CPL signal (Fig. S22 and S23†). It may be that the nanoclusters formed adducts with chiral acids and TPrA leading to chiral changes.⁵⁷

In conclusion, we unambiguously demonstrated that intense CPECL in the near-infrared region can be generated and detected from chiral Au₉Ag₄ metal nanoclusters. Chiral ligand induction and alloying were employed to obtain chiral and luminescent S/R-Au₉Ag₄ metal nanoclusters, and the photoelectrochemical performance of these nanoclusters was explored. The ECL emissions of the metal nanoclusters in the presence of TPrA as a co-reactant were centered at 805 nm. The mirror-imaged CPECL signals of S-Au₉Ag₄ and R-Au₉Ag₄ indicate that the construction and detection of CPL triggered by electrochemistry was successfully achieved. The obtained nanocluster CPECL platform shows the discrimination of chiral 2-chloropropionic acid. Our further work aims to explore and address the interactions between nanoclusters luminophore, co-reactant and discriminating enantiomers. Overall, this work establishes an integration platform of chirality and electrochemiluminescence in metal nanoclusters for discriminating enantiomers and developing novel chiral detection platforms.

Data availability

All the data are shown in the ESI. CCDC 2165721 contains the supplementary crystallographic data for this paper.

Author contributions

L. J. and M. J. performed the synthesis, characterizations, optical and electrochemical tests, and ECL experiments. S. C. and L. J. analysed the data. B. Y., X. W., W. D., and Y. L. assisted in the synthesis and characterizations. S. C. and M. Z. directed the study. All authors prepared the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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