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Engineering of a zero cross-talk fluorescent polymer nanoprobe for self-referenced ratiometric imaging of lysosomal hypochlorous acid in living cells†

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The rational design of a zero cross-talk self-referenced ratiometric fluorescent probe for accurate molecular imaging in living cells is still a major challenge. Herein, we report a zero cross-talk fluorescent polymer nanoprobe (named ZC-FPN) with a self-referenced ratiometric effect for lysosomal hypochlorous acid (HClO) imaging using single-wavelength excitation. The newly designed polymer nanoprobe (ZC-FPN) is prepared by using a simple co-precipitation method through self-assembly of two amphiphilic block copolymers, which are covalently linked with a porphyrin fluorophore (HClO-responsive unit) and a naphthalimide fluorophore (reference unit), respectively. Remarkably, the ZC-FPN displays an unprecedented separation of two emission peaks (~191 nm), which can avoid the spectral cross-talk problem of the emission-shift type ratiometric probes and is highly favorable for high-resolution bioimaging. In addition, the as-prepared ZC-FPN also shows good water dispersibility, high selectivity, excellent long-term fluorescence stability (>10 weeks) and low cytotoxicity. More importantly, the positive charge of the amino moiety enhances the efficient uptake of the nanoprobe by cancer cells and enables the nanoprobe to selectively accumulate in lysosomes, achieving endogenous HClO imaging in living cells through fluorescence ratiometric imaging.

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Introduction

Fluorescent probes integrated with an optical imaging technique have played vital roles in studying biological events owing to several unique features.^{1–19} Among the various strategies to construct fluorescent probes, polymer nanoparticle-based fluorescent nanoprobe (named polymer nanoprobe) have aroused great attention in many fields due to their intrinsic optical properties.^{20–29} To date, a series of currently developed polymer nanoprobe are

primarily based on either “one-emission signal” or “dual-emission signals (ratiometric)”.^{30–35} Compared to the one-emission signal-based polymer nanoprobe, dual-emission signal-based polymer nanoprobe can enable more accurate detection and imaging through self-calibrating dual emission signals, thus providing more reliable quantitative analysis results.³⁵

Generally, the commonly adopted strategies for designing ratiometric polymer nanoprobe are mainly dependent on either Intramolecular Charge Transfer (ICT) or Fluorescence Resonance Energy Transfer (FRET) mechanisms.^{36–43} Unfortunately, most of the ICT or FRET-based ratiometric polymer nanoprobe often show serious cross-talk between two emission bands, resulting in a low-contrast resolution performance for bioimaging.⁴⁴ Therefore, the rational design of novel ratiometric fluorescent polymer nanoprobe with zero cross-talk of two emission signals for high-resolution imaging is still urgently demanded.

Porphyrin and its derivatives have been widely utilized for fluorescent probes due to their excellent features of high photostability, large Stokes shift (>100 nm), near-infrared (NIR) emission (>650 nm), etc.^{45–47} More importantly, our major new finding is that the NIR fluorescence of 4-(10,15,20-triphenylporphyrin-5-yl)phenol (TPP-OH) derivatives can be sensitively and selectively quenched when exposed to hypochlorous acid (HClO) in

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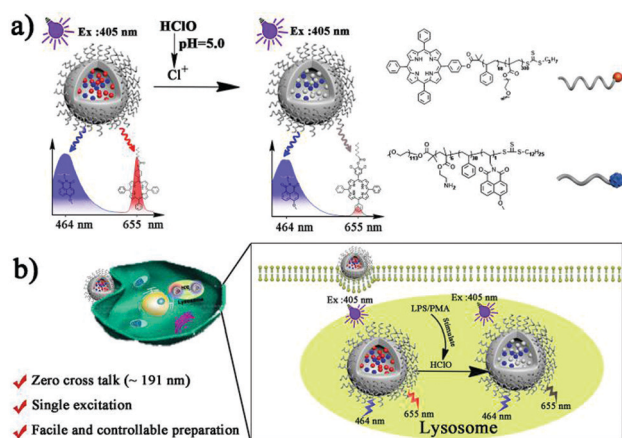
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an acid (pH 5.0) environment, making TPP-OH suitable for visualizing HClO in living cells (especially for lysosomal HClO). On the other hand, naphthalimide and its derivatives have also been demonstrated to be promising probes for bioimaging because they possess high quantum yield, photostability and so on.^{48–50} Previously, Li *et al.* developed a naphthalimide-porphyrin based ratiometric fluorescence probe for imaging intracellular Hg^{2+} with a fixed excitation wavelength (415 nm).⁵¹ This probe displays two well-resolved (zero cross-talk) emission spectra (~ 125 nm), which is beneficial for intracellular imaging with high resolution. Inspired by the work of Li and coworkers, we assume that both porphyrins and naphthalimides can be utilized to construct a zero cross-talk ratiometric fluorescent probe with single-wavelength excitation.

Polymeric micelles self-assembled by amphiphilic diblock copolymers have gained popularity as carriers for the fabrication of multifunctional fluorescent nanoprobe because of their unique advantages including tunable structure design, good water dispersibility, high photostability and excellent biocompatibility, *etc.*⁵² In particular, the covalent incorporation of large π - π conjugate chromophores (eg. porphyrins and naphthalimides) into the micelle core can not only effectively avoid the leakage problem of small dyes physically encapsulated in the micelle core, but also largely improve their brightness.⁴⁴

In view of the good performance of TPP-OH for sensitive and selective HClO detection, we herein design a zero cross-talk fluorescent polymer nanoprobe (named ZC-FPN) with a self-referenced ratiometric detection effect, obtained by self-assembling two fluorophore-contained amphiphilic block copolymers through a simple co-precipitation method (Scheme 1). The as-prepared ZC-FPN displays a single excitation wavelength (405 nm) and two well-resolved emission properties (~ 191 nm), one from the HClO-responsive fluorophore (TPP-OH) and the other from the reference fluorophore (naphthalimide). Furthermore, the nanoprobe with a positive charge can especially stain lysosomes and further be used to detect endogenous HClO in living cells through fluorescence ratiometric imaging.



Scheme 1 Schematic illustration of ZC-FPN for self-referenced ratiometric detection of HClO (a) and ratiometric monitoring of endogenous HClO in lysosomes of living cells (b).

Experimental section

Materials and instruments

The relevant chemical reagents were obtained from commercial companies. The detailed synthesis, characterization, instruments, cytotoxicity and living cell imaging are given in the ESI.[†]

Results and discussion

Synthesis and characterization of ZC-FPN

The selection of dyes was the key point for the fabrication of a zero cross-talk ratiometric fluorescent nanoprobe. The NIR dye (TPP-OH) served as the response unit for HClO recognition as HClO can induce the quenched fluorescence of TPP-OH, while naphthalimide was utilized as the internal reference unit based on its good stability when exposed to HClO. Significantly, the two dyes displayed a single-wavelength excitation (405 nm) and two well-resolved emission peaks at 464 and 655 nm, thereby enhancing the detection accuracy.

Next, two amphiphilic diblock copolymers (TPP-*b*-PS₆₈-*b*-PEGMA₃₀₀ and PEO₁₁₃-*b*-P(AEMH₆-*co*-EANI₁-*co*-PS₃₀)) containing TPP-OH and naphthalimide were synthesized by Reversible Addition-Fragmentation Chain Transfer Polymerization (Schemes S1 and S2, ESI[†]). The relative intermediate and target compounds were identified by ¹H NMR, ¹³C NMR, Mass spectrum, Infrared (IR) spectrum and Gel Permeation Chromatograph (Fig. S1–S9 and Tables S1, S2, ESI[†]).

Finally, by using a simple co-precipitation method, the above two amphiphilic diblock copolymers can be easily self-assembled together into a stable polymeric micelle (ZC-FPN) with an average particle size of about 22 nm, which was measured by Dynamic Light Scattering (DLS, Fig. 1A). In addition, the spherical morphology of ZC-FPN with the corresponding average diameter was further analysed by Transmission Electron Microscopy (TEM, Fig. 1B). Remarkably, no obvious changes of diameter of ZC-FPN were visualized even after being stored at 25 °C for more than 30 days (Fig. 1C), and clear and stable aqueous dispersions of ZC-FPN can also be observed (Fig. 1D). The above results confirmed the excellent colloidal stability of ZC-FPN.

Spectral response of ZC-FPN towards HClO

With the ZC-FPN in hand, the HClO response of ZC-FPN ($48 \mu\text{g mL}^{-1}$) was firstly investigated in a pH = 5.0 simulated physiological environment (PBS). The addition of HClO can result in an obvious decrease in absorption at 418 nm of ZC-FPN, indicating an efficient reaction between the TPP molecules and HClO (Fig. S10, ESI[†]). Different from TPP molecules, naphthalimide molecules (EANI) were inert to HClO, and thus the absorbance at 380 nm of ZC-FPN displayed no distinct change. As can be seen from the normalized excitation spectra of EANI and TPP-OH (Fig. S11, ESI[†]), they share a common excitation wavelength at 405 nm. Upon excitation at 405 nm, a significantly decreased fluorescence (at 655 nm) of TPP-OH can be observed in the presence of HClO, while the fluorescence (at 464 nm) of EANI exhibited no appreciable distinct change when exposed to HClO (Fig. S12, ESI[†]). Significantly, the two well-resolved emission peaks

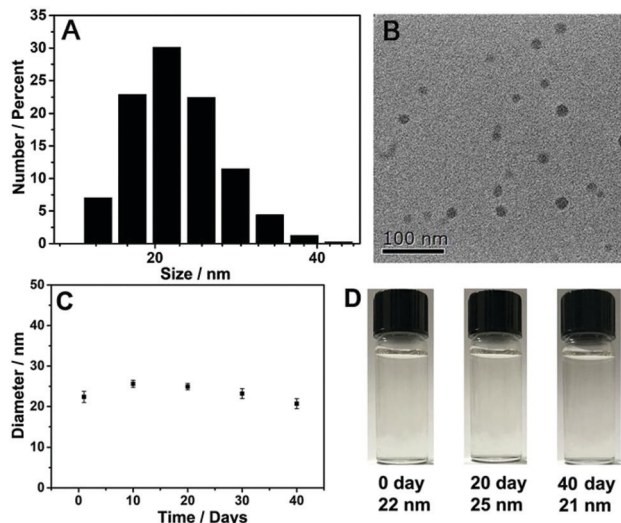


Fig. 1 (A) Average diameter of ZC-FPN; (B) TEM of ZC-FPN; (C) average particle sizes and (D) images of ZC-FPN (1.44 mg mL^{-1}) with increasing time.

($\sim 191 \text{ nm}$) with a zero cross-talk property suggested an improved resolution for intracellular ratiometric imaging.

Subsequently, the capability of ZC-FPN for self-referenced ratiometric detection of HClO was carried out, as shown in Fig. 2A. When incubated with different concentrations of HClO, the ZC-FPN displayed gradually quenched fluorescence at 655 nm, while the reference blue band at 464 nm remained almost unchanged. Notably, the quenched fluorescence of ZC-FPN at 655 nm may be ascribed to the intramolecular heavy atom effect of a chlorinated porphyrin structure (Scheme S3, ESI[†]).^{53–55} Both the $^1\text{H NMR}$ (Fig. S13, ESI[†]) and mass spectrum (Fig. S14, ESI[†]) also confirmed this response mechanism. In addition, the fluorescence intensity ratios (I_{464}/I_{655}) of ZC-FPN illustrated a good linear relationship with increasing HClO concentration (Fig. 2B), and the detection limit was determined to be $1.99 \mu\text{M}$. The time response of ZC-FPN towards HClO was also investigated (Fig. S15, ESI[†]). When exposed to HClO, the I_{464}/I_{655} ratio of ZC-FPN increased remarkably and then reached a plateau in 7 min. Therefore, 7 min was selected as a suitable time for sensing HClO.

Additionally, the selectivity experiments of ZC-FPN were carried out in the presence of various analytes commonly existing in biological matrices, and the results are given in Fig. 2C. Fortunately, only HClO could trigger a remarkable fluorescence intensity ratio change, while other potential interfering substances exhibited no perceptible effect, suggesting a good selectivity of ZC-FPN to HClO. Moreover, the pH influence of the fluorescence intensity ratio (I_{464}/I_{655}) of ZC-FPN treated with and without HClO was also visualized. As shown in Fig. 2D, the I_{464}/I_{655} ratio of ZC-FPN itself was low and unchanged obviously from pH 3.0 to 9.0, while a remarkably enhanced I_{464}/I_{655} ratio of ZC-FPN treated with HClO can be visualized from pH 3.0 to 8.0, suggesting the good capability of ZC-FPN for the ratiometric detection of HClO in complex biosystems. Moreover, the preferable photostability and excellent long-term

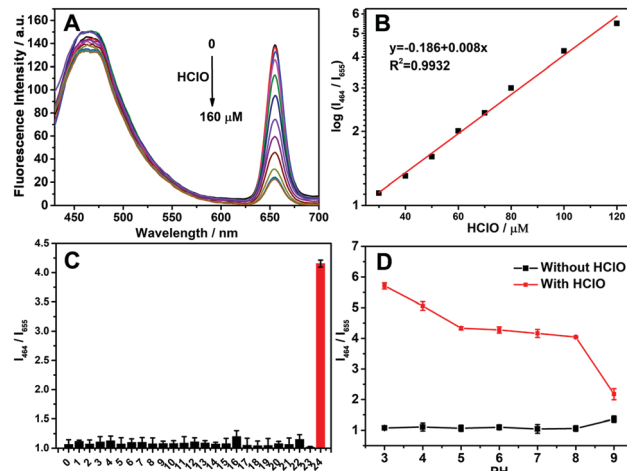


Fig. 2 (A) Ratiometric emission spectra of ZC-FPN ($48 \mu\text{g mL}^{-1}$) in the presence of HClO (0– $160 \mu\text{M}$) in pH 5.0 PBS buffer solution ($\lambda_{\text{ex}} = 405 \text{ nm}$); (B) working curve of ZC-FPN to HClO (30– $120 \mu\text{M}$); (C) the fluorescence intensity ratio (I_{464}/I_{655}) of ZC-FPN ($48 \mu\text{g mL}^{-1}$) in the presence of various analytes ($100 \mu\text{M}$ for HClO and 1 mM for others): (0) blank, (1) Na^+ , (2) Mg^{2+} , (3) Ni^{2+} , (4) Ca^{2+} , (5) K^+ , (6) Zn^{2+} , (7) Cu^{2+} , (8) PO_4^{3-} , (9) HPO_4^{2-} , (10) H_2PO_4^- , (11) SO_4^{2-} , (12) NO_3^- , (13) SO_3^{2-} , (14) Cl^- , (15) HS^- , (16) H_2O_2 , (17) Cys, (18) GSH, (19) Hcy, (20) $^1\text{O}_2$, (21) Bu-OOH, (22) $\cdot\text{OH}$, (23) $t\text{-BuO}^\bullet$, (24) ONOO^- , (25) HClO; (D) effects of pH on the fluorescence intensity ratio (I_{464}/I_{655}) for ZC-FPN ($48 \mu\text{g mL}^{-1}$) without (black curve) and with (red curve) HClO ($100 \mu\text{M}$). Values are mean \pm s.d. for $n = 5$.

fluorescence stability can make ZC-FPN more suitable for further intracellular imaging applications (Fig. S16 and S17, ESI[†]).

Ratiometric imaging in living cells

In view of the above good results, we further intend to evaluate the performance of ZC-FPN for exogenous/endogenous HClO imaging in lysosomes. Before imaging, the degree of toxicity of ZC-FPN toward the representative HeLa cells was evaluated. As can be seen from Fig. S18 (ESI[†]), minimal cytotoxicity was visualized even at high concentrations ($100 \mu\text{g mL}^{-1}$), indicating the excellent biocompatibility of ZC-FPN.

Some references suggested that nanoprobe with a positive charge can selectively localize to the lysosomes (Fig. S19, ESI[†]).^{53,56} The co-localization experiments were carried out by using ZC-FPN and a lysosomal marker (Lyso-Tracker Green) to confirm the location of ZC-FPN in lysosomes. As shown in Fig. 3, the blue fluorescence image of ZC-FPN merged well with that of Lyso-Tracker Green, and the high Pearson's coefficient and overlap coefficient were 0.9727 and 0.9748, suggesting that ZC-FPN mainly stained lysosomes.

Next, ZC-FPN was utilized to ratiometrically detect exogenous HClO in lysosomes of HeLa cells by using NaClO as the HClO donor (Fig. 4). The red fluorescence intensities gradually decreased with increasing HClO concentrations, while no obvious changes can be visualized from the blue channel. Correspondingly, the ratio images ($F_{\text{blue}}/F_{\text{red}}$) also supported this variation tendency (Fig. 4M–P). In view of the good ability of ZC-FPN for exogenous HClO imaging, we then evaluated its performance for endogenous HClO detection in Raw264.7 macrophage cells by stimulating cells

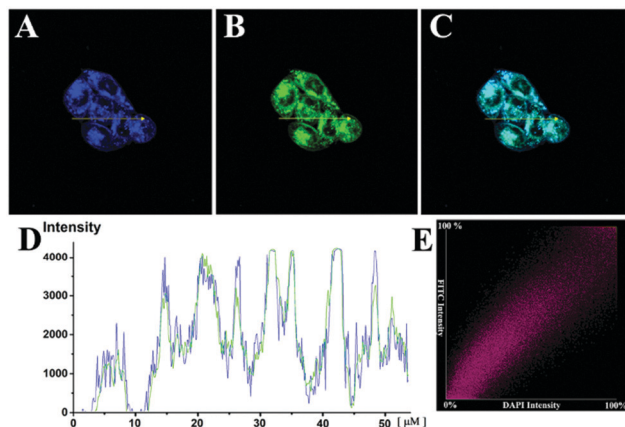


Fig. 3 Co-localization experiments of (A) ZC-FPN ($48 \mu\text{g mL}^{-1}$) and (B) Lyso-Tracker Green (500 nM) in HeLa cells. (C) Overlay of (A) and (B); (D) intensity correlation plot of ROI from (A) and (B); (E) the correlation plot of (A) and (B). Blue channel: $\lambda_{\text{ex}} = 405 \text{ nm}$ and $\lambda_{\text{em1}} = 440\text{--}500 \text{ nm}$; green channel: $\lambda_{\text{ex}} = 504 \text{ nm}$ and $\lambda_{\text{em}} = 520\text{--}540 \text{ nm}$ scale bar: $20 \mu\text{m}$.

with Lipopolysaccharide (LPS) and Phorbol Myristate Acetate (PMA).^{57,58} As can be seen from Fig. 5D and G, there were strong fluorescence signals in both the blue and red channels, indicating the efficient uptake of ZC-FPN by RAW264.7 macrophage cells. In contrast, when the RAW264.7 cells were co-treated with LPS, PMA and ZC-FPN, the blue channel still displayed a bright fluorescence signal (Fig. 5E), while an obvious weak fluorescence signal in the red channel was noted (Fig. 5H). To verify that the endogenously generated HClO were responsible for the intracellular fluorescence

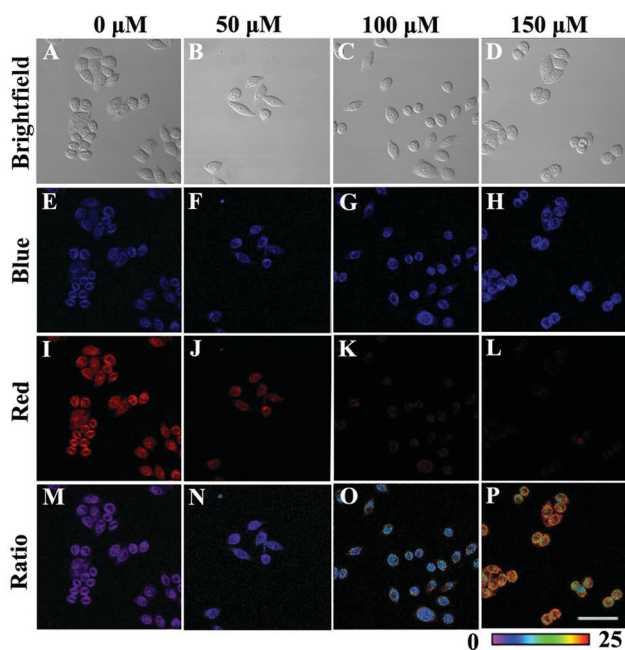


Fig. 4 Confocal fluorescence microscopy images of HeLa cells treated with ZC-FPN ($48 \mu\text{g mL}^{-1}$) followed by incubation with 0 (A, E and I), 50 (B, F and J), 100 (C, G and K) and 150 μM (D, H and L) HClO. (M, N, O and P) The ratio images ($F_{\text{blue}}/F_{\text{red}}$). $\lambda_{\text{ex}} = 405 \text{ nm}$, $\lambda_{\text{em1}} = 440\text{--}500 \text{ nm}$ (blue channel), and $\lambda_{\text{em2}} = 640\text{--}670 \text{ nm}$ (red channel) scale bar: $20 \mu\text{m}$.

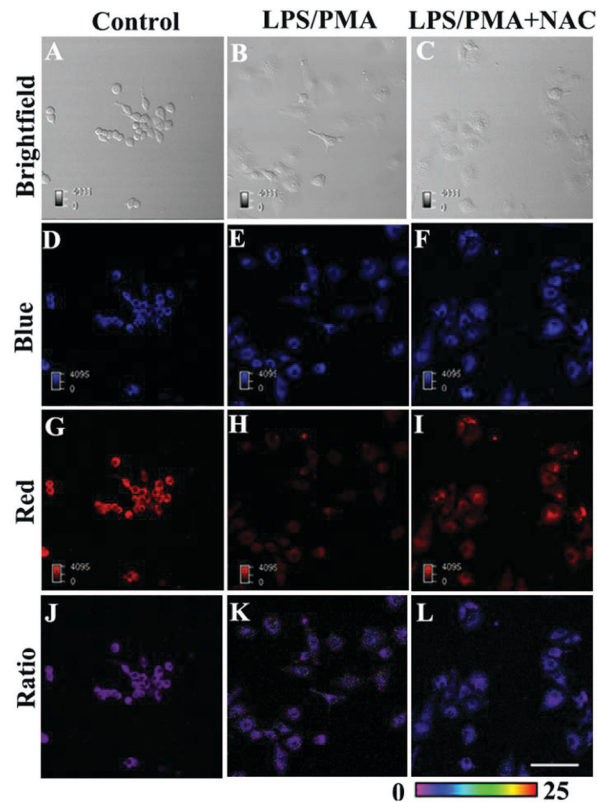


Fig. 5 Confocal fluorescence microscopy images of RAW264.7 macrophage cells. (A, D and G) Cells incubated with ZC-FPN ($48 \mu\text{g mL}^{-1}$); (B, E, and H) cells incubated with LPS (1 mg mL^{-1}), followed by co-incubation with PMA (1 mg mL^{-1}) and ZC-FPN ($48 \mu\text{g mL}^{-1}$); (C, F, and I) cells incubated with LPS (1 mg mL^{-1}) and NAC (2 mM), followed by co-incubation with PMA (1 mg mL^{-1}) and ZC-FPN ($48 \mu\text{g mL}^{-1}$); (J, K and L) the ratio images ($F_{\text{blue}}/F_{\text{red}}$); $\lambda_{\text{ex}} = 405 \text{ nm}$, $\lambda_{\text{em1}} = 440\text{--}500 \text{ nm}$ (blue channel), and $\lambda_{\text{em2}} = 640\text{--}670 \text{ nm}$ (red channel) scale bar: $20 \mu\text{m}$.

signal changes, a control experiment was performed by treating the LPS/PMA stimulated RAW264.7 cells with *N*-acetylcysteine (NAC, an effective ROS scavenger).⁵⁹ Because NAC can efficiently remove the endogenously generated HClO, and thus result in a very low HClO level in RAW264.7 cells. As expected, both the blue and red channels exhibited bright fluorescence signals (Fig. 5F and I), indicating that the quenched fluorescence signals of the red channels indeed originated from the endogenously produced HClO. Additionally, when the RAW264.7 cells were treated with NAC and ZC-FPN, both the blue and red channels exhibited strong fluorescence signals (Fig. S20, ESI[†]), indicating that NAC cannot induce the fluorescence signal change of the red channel. These results revealed the good ability of ZC-FPN for exogenous/endogenous HClO imaging in living cells.

Conclusion

To sum up, we have successfully developed a zero cross-talk fluorescent polymer nanoprobe (ZC-FPN) for HClO imaging in lysosomes with a ratiometric effect. The rationally designed ZC-FPN possesses some prominent advantages over the previously described HClO fluorescent probes with a dual-emission

property (Table S1, ESI[†]): First, the synthesized ZC-FPN can operate in a 100% aqueous solution, whereas the previously described HClO fluorescent probes operated in mixed organic-aqueous solvent systems; second, the ZC-FPN illustrates an ideal single excitation wavelength and well-resolved dual emissions (~191 nm), which can avoid the spectral cross-talk issue of the emission-shift type or the FRET-based ratiometric probes and is highly beneficial for high-resolution bioimaging; third, benefiting from the positive charge of the amino moiety modified nanoparticles, the ZC-FPN with good cell-membrane permeability can selectively locate at lysosomes, and further be utilized to detect exogenous and endogenous HClO in the lysosomes of living cells by using fluorescence ratiometric imaging.

Conflicts of interest

The authors declare no competing financial interest.

Acknowledgements

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