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Synthesis, micellar structures and emission mechanisms of an AIE and DDED-featured fluorescent pH- and thermo-meter

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A new nanoprobe, the luminescent diblock copolymer PNIPAM(MAh-4)-*b*-P4VP (PN4P), with pH- and thermo-responsive deprotonation-driven emission decay (DDED) and aggregation-induced emission (AIE) features was designed and synthesized. The nanoprobe PN4P can form micellar structures in water with reversible dual-responsive fluorescence (FL) behavior within a wide pH range of 2–11. The critical solution temperature was found at about 32, 30 and 27 °C as the pH switched from 2, 7 to 11. The critical pH value of the probe was about 4.0, and the micelles showed a core–shell inversion in response to pH and thermal stimuli, accompanied by a desirable emission tunability. P4VP as the micellar shell at pH = 2 was more easily dehydrated with the increase in temperature as compared to PNIPAM as the micellar shell at pH > 4. The strongest dehydration of the P4VP shell would make PN4P the most strongly aggregated and the most AIE-active, which supports the 2.10-fold most distinguished thermal-responsive emission enhancement at pH = 2. Moreover, a dramatic acidochromic redshift of the emission band from 450 (pH > 4) to 490 nm (pH = 2) was observed, and the maximum emission at pH = 2 was enhanced by about 2.07-fold as compared with that at pH = 7. Therefore, the probe displays the desired dual responses and good reversibility. AIE and DDED are the two major mechanisms responsible for the dual-responsive emission change, with AIE playing a more important role than DDED. This work offers a promising approach to interpreting temperature (range from 28 to 40 °C) and pH changes (range from 2 to 7) in water.

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1. Introduction

Stimuli-responsive polymers are powerful tools in drug delivery,^{1–4} tissue engineering,⁵ biological sensors,^{6–8} and fluorescence (FL) probes.^{9–11} In recent years, many stimuli-responsive polymers, whose properties change in response to ionic strength, light, pH and temperature, have been studied,¹² and molecular level solubility changes of segments could be triggered by such stimuli as temperature,¹³ pH,¹⁴ CO₂ (ref. 15 and 16) and electric/magnetic field.¹⁷ Among the diverse stimuli, thermal- and pH-responsive polymers have been widely explored due to their potential *in vivo* applications,^{11,18} and polymers with both thermal and pH stimuli are extensively studied dual responsive systems.^{19–23} As is known, poly(*N*-

isopropylacrylamide) (**PNIPAM**) is a thermal-responsive polymer with a lower critical solution temperature (LCST) of about 32 °C.²⁴ Owing to its thermo-sensitivity and desired biocompatibility, **PNIPAM**-containing materials have been extensively employed in temperature-targeted therapy systems and drug delivery materials.^{18,25} **PNIPAM** was also used to regulate the fluorophore emission *via* thermally-induced aggregations and micro-environmental changes.^{26,27} Polymers with end-capped dye moieties²⁶ or at the junction between two blocks have been reported.²⁷ Guo *et al.*²⁶ synthesized a fluorescent nano-thermometer **PNIPAm-MAh-4** with aggregation-induced emission (AIE) character, which showed enhanced emission in water with a temperature increase and the maximum emission at 50 °C was about 7-fold stronger than that at 15 °C. In contrast, a carbazole-containing **PNIPAM** copolymer solution²⁷ exhibited 22 times emission enhancement along with a temperature drop from 50 to 25 °C. Besides, a few double-hydrophilic diblock copolymers with both thermo and pH responses were reported, together with a core–shell inversion,²³ which included poly(*N*-isopropylacrylamide)-*block*-poly(4-vinylpyridine) (**PNIPAM-*b*-P4VP**), **PNIPAM-*b*-PAA** and **PPO-*b*-PDEA**. Among these, **PPO-*b*-PDEA** micelles were unstable at ambient temperature and the **PNIPAM-*b*-PAA** counterpart had strong hydrogen bonds

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between PAA and **PNIPAM** blocks, which greatly limited their further applications.²³ **PNIPAM-*b*-P4VP** seemed to be the most desirable dual-responsive copolymer because there were no strong hydrogen bonds between **P4VP** and **PNIPAM** blocks, and **PNIPAM** covers a critical aggregation temperature range that is especially suitable for biomedical applications.

In addition, amphiphilic block copolymers can form micelles *via* self-assembly in selective solvents, and various dye-functionalized amphiphilic block copolymers have been prepared to realize further FL property control through the regulation of their self-assembled nanostructures in solutions.²⁷⁻³² Cheng *et al.*^{28,31} reported their work on the structure and emission behavior tuning of self-assembled fluorescent composites (SAFC) between dye molecule and **PS-*b*-P4VP** in CO_2 -expanded liquids. The morphology and emission behaviors of SAFC were considerably pressure dependent and the maximum FL intensity of SAFC at 5.41 MPa was enhanced 14-fold.²⁸ Tan *et al.*³² designed FL vesicles *via* the complexation of FL poly-oxometalate and stimuli-responsive triblock copolymer **PEO-*b*-PS-*b*-PDMAEMA** by electrostatic interactions, which showed on-off switchable emission behavior along with pH tuning of the reversible micelle-to-vesicle change. Unfortunately, these polymers had no temperature response and their micellization process often involved bio-incompatible organic solvents.

To date, AIE-featuring materials have received great attention owing to their potentials in stimuli-responsive materials,^{33,34} life sciences^{11,35,36} and biomedical engineering.^{4,37,38} For example, TPE polymers³³ were synthesized with AIE character, pH response and solvatochromic behavior, and the P1-P3 based sensors exhibited pH response through the protonation of the amino groups and could identify twelve different nitroarenes in

water; however, the above TPE polymers are not temperature responsive. Very recently, AIE-active nanoprobes **PNIPAM-*b*-P(DPA-*co*-TPE)** and **P(NIPAM-*co*-TPE)-*b*-PDPA** with pH and thermo responses were reported,¹¹ and the respective FL intensities were enhanced 7- and 3-fold for **PNIPAM-*b*-P(DPA-*co*-TPE)** and **P(NIPAM-*co*-TPE)-*b*-PDPA** by solution pH or temperature tuning. Although **PNIPAM-*b*-P4VP** is a desired thermo- and pH- dual-responsive copolymer,²³ little attention has been paid to the luminescence performance of chromophores regulated by **PNIPAM-*b*-P4VP**-based polymers, and it appears that thermal and pH responsive fluorescent sensor materials remain a highly unexplored area. Therefore, our objectives include the design of a novel biocompatible dye-labeled polymer probe with pH and thermo dual-responses suitable for biomedical applications, showing entirely different pH-driven emission trends against temperature, and finding the dual-responsive micellar structures and emission mechanisms of the probe in water, uncovering the dominant factors that decide the pH- or thermal responses, as well as obtaining the sensitivity range or suitable working temperature and pH range of the probe. The innovation of this work is to ensure that the developed pH and temperature-responsive probe has the features of totally different emission mechanisms such as AIE/deprotonation-driven emission decay (DDED) against temperature within different pH ranges. Thus, a **PNIPAM-*b*-P4VP**-based polymer labeled with a dye unit possessing both AIE and DDED features appears to be an ideal choice. Accordingly, in this work, we focus on the synthesis and dual-responsive emission behaviors of a double-hydrophilic diblock copolymer **PNIPAM(MAh-4)-*b*-P4VP** (**PN4P**, Fig. 1). Herein, **PNIPAM** and **P4VP** blocks, as well as **4** moieties,²⁶ endow the **PN4P** with thermal-, pH-, and FL-

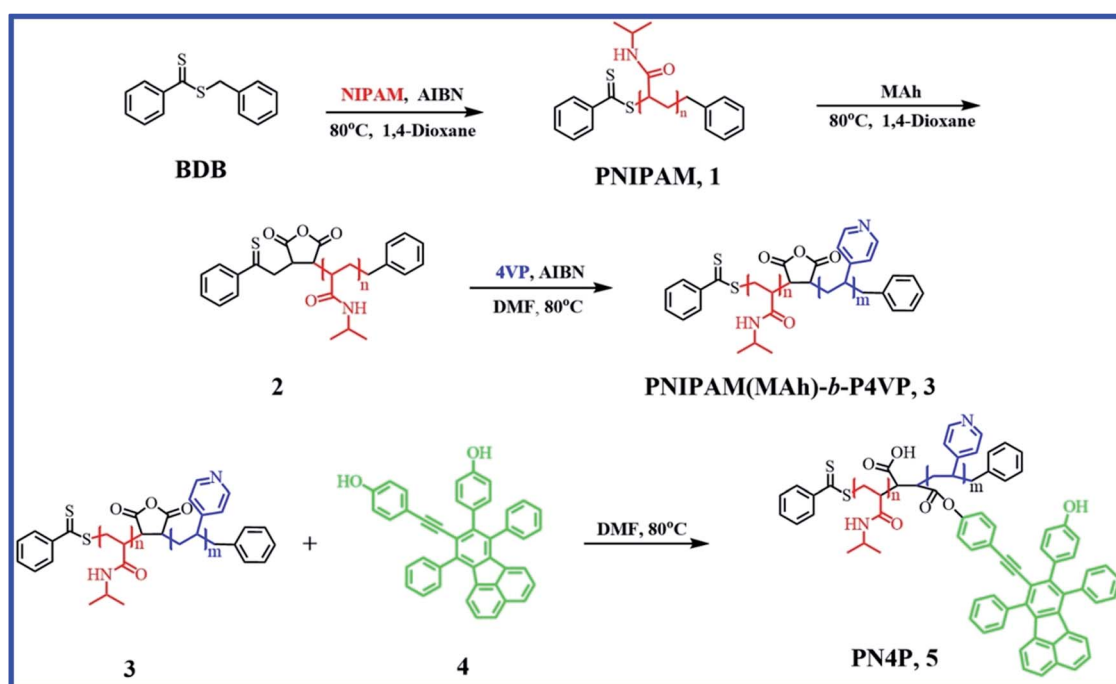


Fig. 1 Synthetic route for PN4P (or compound 5).



responsive properties. Particularly, the dye unit **4**, with partially propeller-like moieties and phenolic OH groups as an archetypal AIE/DDED module, was used to label the **PNIPAM-*b*-P4VP** and was expected to serve as an intriguing building block for AIE/DDED-active luminogenic polymers. The micellar/clustering behavior of the probe in water was studied by solution dispersibility, DLS, NMR, and TEM measurements, and the emission performance was systematically examined by temperature and pH tuning. As expected, the **PN4P** probe demonstrates obvious DDED and AIE features in different microenvironments, and simultaneously displays good reversibility and is more sensitive within the pH range of 2 to 7 and temperature range from 28 to 40 °C.

2. Experimental section

2.1. Materials

4-Vinyl pyridine (4VP, Alfa Aesar, Beijing, 96%) was distilled under reduced pressure and stored in a refrigerator. *N*-Isopropylacrylamide (**NIPAM**, Shanghai Chemical Reagents Co., 98%) was recrystallized from toluene and *n*-hexane (1 : 3). Maleic anhydride (MAh, Beijing Chemical Reagents Co., 99%) was recrystallized from anhydrous chloroform. 2,2-Azobis(isobutyronitrile) (AIBN, Beijing Chemical Reagents Co., 98%) was recrystallized from ethanol twice. 1,4-Dioxane (Tianjin Chemical Reagents Co., 99%) was distilled in the presence of sodium slices. Tetrahydrofuran (THF, Beijing Chemical Reagents Co., 99%) was distilled from sodium benzophenone ketyl under argon immediately prior to use. Magnesium powder, potassium hydroxide, iodine, MgSO₄, CuI and PdCl₂, dichloromethane, chloroform, ethyl acetate, methanol, ethanol, petroleum ether and *ortho*-xylene (Beijing Chemical Reagents Co., analytical pure grade) were used as received. The ethanol for absorption (Abs) was HPLC grade (≥99.9%), obtained from Aladdin, and was used as received. 4-(9-(2-(4-Hydroxyphenyl)ethynyl)-7,10-diphenylfluoranthen-8-yl)phenol (**4**) and benzyl dithiobenzoate (BDB) were synthesized in our lab according to a reference procedure.²⁶ **PNIPAM** ($M_n = 16\,200\text{ g mol}^{-1}$) and copolymer **PNIPAM(MAh-*b*-P4VP** ($M_n = 26\,200\text{ g mol}^{-1}$) were synthesized according to a procedure described previously,^{26,28} and the detailed reaction conditions for the preparation of **PNIPAM** and **PNIPAM(MAh-*b*-P4VP**, the number average molecular weight (M_n) and polydispersity index (M_w/M_n) of **PNIPAM** and **PNIPAM(MAh-*b*-P4VP** determined by gel permeation chromatography (GPC) are tabulated in Table 1. The **PNIPAM(MAh-*b*-P4VP** (**PN4P**) probe was synthesized according to a reference

Table 2 PN4P results^a

Sample	PNIPAM/ 4 ^b	NIPAM/ 4 ^c	$M_{n, GPC}^d$	M_w/M_n^d
PN4P	1 : 5	1600 : 1	26 000	1.34

^a The reaction carried out at 80 °C for 24 h. Sample **PN4P** was prepared from sample 2 shown in Table 1. ^b Molar feed ratio of **PNIPAM** to **4**.

^c Molar ratio of **NIPAM** to **4** obtained from **PN4P** using the absorbance calibration, by comparison of absorbance of **PN4P** with **4** ($A_{390\text{ nm}}$) dissolved in ethanol at 25 °C. ^d Determined by GPC in THF at 313.2 K, flow rate of 1 mL min⁻¹.

method,²⁶ the detailed reaction conditions and the number average molecular weights from the GPC measurement are tabulated in Table 2. The degree of labeling (molar ratio of **4** to **NIPAM**) in **PN4P** was determined to be 0.063% (or 1/1600, Table 2) by comparison of the absorbance with compound **4** (A390 nm) dissolved in ethanol at 25 °C. The synthesis route for **PN4P** is summarized in Fig. 1. The pH buffers from Sigma-Aldrich were used as received. The lower pH value of the solvent was adjusted with 0.1 mol L⁻¹ HCl and the higher pH value of the solvent was adjusted with 0.1 mol L⁻¹ NaOH. All other reagents were analytical grade and were used without further purification.

2.2. Synthesis

2.2.1 PNIPAM (1). **PNIPAM** was prepared *via* the reversible addition-fragmentation chain transfer (RAFT) polymerization method as reported previously.²⁶ **NIPAM** (2 g, 0.0176 mol), AIBN (0.002 g, 0.0122 mmol), BDB (0.0209 g, 0.0855 mmol) and 1,4-dioxane (3 mL) were added to a 20 mL Schlenk tube, followed by three freeze–vacuum–thaw cycles. The tube was sealed under vacuum and then immersed in an oil bath at 80 °C with magnetic stirring. After reaction for 9 h, the tube was cooled to room temperature and opened to the air. The polymer was dissolved in a suitable amount of 1,4-dioxane, and then precipitated by dropping the solution into anhydrous ether, followed by filtration. Repeating the procedure of dissolving–precipitation–filtration three times, the obtained pink product was dried under vacuum at 30 °C for 24 h to obtain the **PNIPAM**. ¹H NMR (CDCl₃, 400 MHz): δ 4.00 (s, (CH₃)₂CHNH), 1.13 (s, (CH₃)₂CHNH). The ¹H NMR data, the number average molecular weights and polydispersity index from GPC for **PNIPAM** are also displayed in Fig. 2B and C, respectively.

2.2.2 PNIPAM-MAh (2). **PNIPAM** (1 g, 0.0625 mmol), MAh (0.28 g, 2.85 mmol), and 1,4-dioxane (4 mL) were added to a 20 mL Schlenk tube, followed by three freeze–vacuum–thaw cycles. Afterwards, the vacuum-sealed tube was immersed in an oil bath at 80 °C with magnetic stirring. After reaction for 6 h, the tube was quickly cooled to room temperature. The polymer was dissolved in a suitable amount of THF and then precipitated in anhydrous ether to remove the excess MAh, followed by filtration. The operation of dissolving–precipitation–filtration was repeated three times, and the pink polymer obtained was dried under vacuum at 30 °C for 24 h to obtain the **PNIPAM**.

Table 1 Polymerization of **PNIPAM** (1) and **PNIPAM(MAh-*b*-P4VP** (2) under different reaction conditions

Sample	<i>t</i> (h)	Initiator	BDB/AIBN	<i>T</i> (K)	M_n^a	M_w/M_n^a	Yield (%)
1	9	AIBN	7 : 1	353.2	16 200	1.16	55.6
2	24	AIBN	7 : 1	353.2	26 200	1.24	58.3

^a Number-average molecular weight (M_n) and polydispersity index (M_w/M_n) were determined by GPC in THF at 313.2 K, flow rate 1 mL min⁻¹.



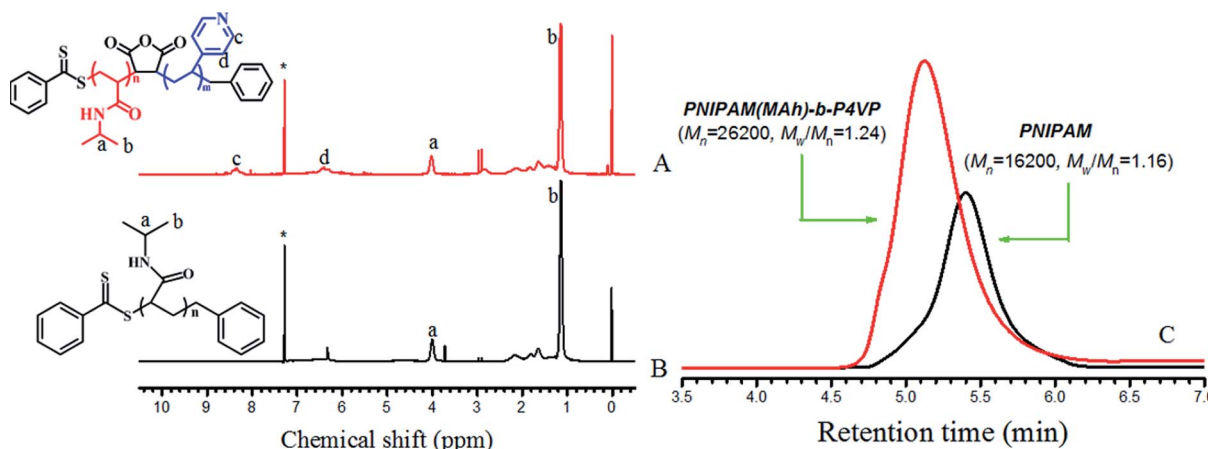


Fig. 2 ¹H NMR spectra of (A) PNIPAM(MAh)-b-P4VP in CDCl₃, (B) PNIPAM in CDCl₃. Solvent signals are marked with asterisks (*); (C) GPC of PNIPAM and PNIPAM(MAh)-b-P4VP.

MAh. IR (KBr, cm⁻¹): 1730.32 ($\nu_{C=O}$); 1652.72 ($\nu_{NH-C=O}$); 1172.53 (ν_{C-S}).

2.2.3 PNIPAM(MAh)-b-P4VP (3). 4VP (0.19 g, 4.66 mmol), PNIPAM-MAh (0.35 g, 0.0218 mmol), AIBN (0.0037 g, 0.0225 mmol), and DMF (3 mL) were added to a 20 mL Schlenk tube with a magnetic bar. After three freeze–vacuum–thaw cycles, the vacuum-sealed tube was immersed in an oil bath at 80 °C and reacted for 24 h. The tube was cooled to room temperature and opened to the air. The polymer was dissolved in DMF, and then precipitated by adding the solution into anhydrous ether dropwise, followed by filtration. The process of dissolving–precipitation–filtration was repeated three times and the obtained pale-yellow product PNIPAM(MAh)-b-P4VP was dried under vacuum at 30 °C for 24 h. ¹H NMR (CDCl₃, 400 MHz): δ 8.0–8.2 (m, 2H), 6.0–6.3 (m, 2H). The ¹H NMR and GPC data for PNIPAM(MAh)-b-P4VP can be found in Fig. 2A and C, respectively.

2.2.4 4-(9-(2-(4-Hydroxyphenyl)ethynyl)-7,10-diphenylfluoranthen-8-yl)phenol (4). Compound 4 was previously prepared in our lab according to the method reported.²⁶ ¹H NMR, δ (CD₃SOCD₃, ppm): 9.84 (s, –OH), 9.30 (s, –OH), 6.30–7.85 (aromatic proton).

2.2.5. PNIPAM (MAh-4)-b-P4VP (5 or PN4P). In a typical polymerization procedure, PNIPAM(MAh)-b-P4VP (0.500 g, 0.0192 mmol), compound 4 (0.050 g, 0.0889 mmol) and DMF (3 mL) were successively added to a dry Schlenk tube, followed by three freeze–vacuum–thaw cycles. Afterwards, the vacuum-sealed tube was immersed in an oil bath at 80 °C with magnetic stirring. After reaction for 24 h, the tube was promptly cooled to room temperature and opened to the air. The polymer was dissolved in the appropriate amount of DMF, and then precipitated in anhydrous ether. The dissolving–precipitation–filtration process was repeated three times and the resultant polymers (PN4P or 5) were dried under vacuum for 24 h to finally obtain the pale-yellow product. The number average molecular weights and polydispersity index from GPC for PN4P, and the labeling degree of the fluorescent moiety 4 in PN4P (molar ratio of NIPAM to 4) from the absorbance measurement

can be found in Table 2. However, considering the low labeling degree of 4 in PN4P (molar ratio of NIPAM/4 = 1600/1), it was somewhat difficult to confirm PN4P by IR or NMR data; we then turned to a thin-layer chromatography (TLC) test to help to identify the PN4P. When comparing PN4P and 4 under the same chromatography conditions using a mixture of ethyl acetate and petroleum ether (3 : 1, volume ratio), we observed two blue-green light-emitting spots at different positions on the same TLC plate under the illumination of a UV lamp, with R_f value of about 7.0 and 0.0 mm for 4 and PN4P, respectively. This proved that PN4P was prepared successfully and demonstrates that PN4P is more polar than 4.

2.3 General information

Fourier transform infrared (FT-IR) spectra were recorded on a NEXUS-470 spectrometer. ¹H NMR spectra were obtained from a DRX-400 NMR instrument with tetramethylsilane as an internal standard. The average molecular weights (M_w and M_n) and polydispersity indexes (PDIs) were measured by an Agilent 1200 Series LC Gel permeation chromatography instrument equipped with a G1310A HPLC Iso Pump, G1362A differential refractive index (RI) detector and four columns at 40 °C, using monodispersed polystyrene (PS) as a calibration standard. THF was used as the eluent at a flow rate of 1.0 mL min⁻¹. UV-vis spectra were taken on a Persee TU-1901 spectrophotometer with a temperature controller. The pH- and thermal-responsive fluorescence measurements were carried out on a Hitachi FL-4600 spectrofluorometer. The emission spectra were recorded at an excitation wavelength of 390 nm. The micellar/clustering morphology of PN4P in water was characterized by transmission electron microscopy (TEM), using a H-800 microscope (Hitachi, Japan). TEM samples were prepared by dropping sample solutions on carbon-coated copper grids, absorbing the solvent on filter paper, and evaporating the solvent at room temperature. Dynamic light scattering (DLS) measurements were carried out to analyze the hydrated diameter of PN4P micelles in aqueous solution with a Nano Plus-3 Particle Size

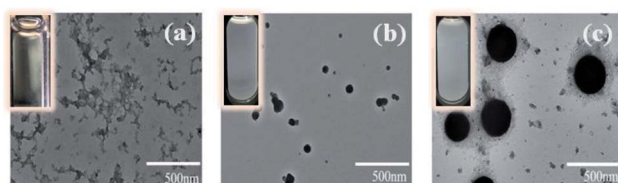


Fig. 3 (a–c) Temperature-responsive TEM images of PN4P in water at pH = 2 and the concentration of 0.5 mg mL^{−1} at (a) 20 °C, (b) 34 °C, (c) 50 °C. Inset: the corresponding dispersibility images of the solution.

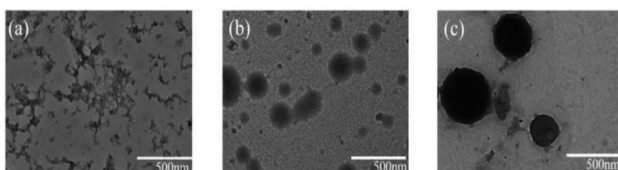


Fig. 4 pH-responsive TEM images of PN4P in water at 20 °C and the concentration of 0.5 mg mL^{−1} at (a) pH = 2, (b) pH = 7, (c) pH = 11.

analyzer (Micromeritics Instruments, USA), equipped with a He–Ne laser (664.4 nm, 70 mW). The Nano Plus version 5.22 software package (Micromeritics Instrument) was used to assist the measurements and data analysis. The hydrodynamic radius (R_h) can be extracted from the Stokes–Einstein equation as follows:

$$R_h = k_B T / 6\pi\eta D, \quad (1)$$

where T denotes the absolute temperature, and the parameters k_B , D and η represent the Boltzmann constant, the diffusion coefficient and the solvent viscosity, respectively. All the pH measurements were made with a PHS-2F precision pH meter

(Inesa Instruments of Shanghai, China), which was calibrated with standard buffer solutions (pH 6.86 and pH 9.18).

3. Results and discussion

3.1 The pH- and thermal-responsive aggregation/micellization of PN4P in water

In this section, the combination of DLS, ¹H NMR, TEM and dispersibility data against temperature/pH were used to help clarify the clustering structure of PN4P in water.

The thermal-responsive solubility/dispersibility and TEM images of PN4P aqueous solutions at pH = 2 are displayed in Fig. 3, and the pH-responsive TEM images at 20 °C are shown in Fig. 4. It was found from the TEM data that the structures of the copolymer clusters appeared as spherical composite micelles under conditions of pH = 2 and temperature ≥ 34 °C (Fig. 3), or at conditions of 20 °C and pH > 2 (Fig. 4). Firstly, it was found that the dispersibility state of the PN4P aqueous solutions at pH = 2 (inset of Fig. 3) changed from the transparent state (Fig. 3a, 20 °C) to an emulsion (Fig. 3b, 34 °C), and then to a turbid state with perceptible precipitation (Fig. 3c, 50 °C), which corresponds to the worsening solubility of the copolymer as temperature increased. In other words, the clustering degree of copolymer was enhanced with increasing temperature, as reflected in the micelle morphology shown in the TEM images (Fig. 3). Although the structures of the copolymer cluster exhibited core-shell-like spherical composite micelles (Fig. 3b and c), more NMR data are still needed to verify which block would be the core or shell-forming segment. Fig. 5 shows the ¹H NMR spectra of PN4P in D₂O at different temperatures and pH values. The characteristic peaks of the methine and methyl protons of the PNIPAM block are supposed to be at about 3.8–4.0 (peak a) and 1.0–1.4 ppm (peak b), and the chemical shifts at about 8.4–8.6 (peak c) and 7.3–7.5 ppm (peak d) can be assigned to the 2,2' and 3,3' protons of the pyridine ring of the P4VP

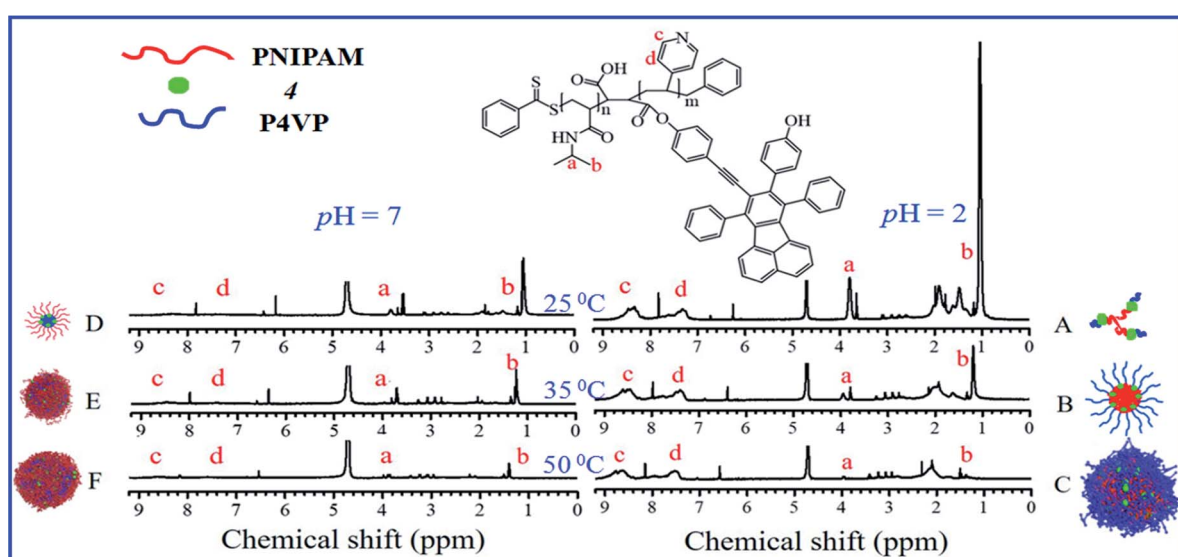


Fig. 5 ¹H NMR spectra obtained for PN4P in D₂O at (A) 25 °C and pH = 2, (B) 35 °C and pH = 2, (C) 50 °C and pH = 2, (D) 25 °C and pH = 7, (E) 35 °C and pH = 7, (F) 50 °C and pH = 7.



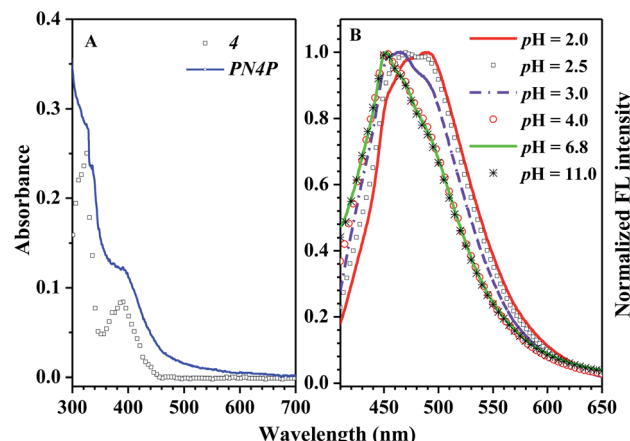


Fig. 6 (A) Absorbance spectra of PN4P and **4** at 25 °C. **4** in ethanol (squares) at the concentration of 1×10^{-5} mol L $^{-1}$, and PN4P in water at pH = 2 (solid line) with concentration of 1.0 mg mL $^{-1}$. (B) Normalized FL spectra against pH value at 25 °C, PN4P in water with concentration of 0.5 mg mL $^{-1}$ and $\lambda_{\text{ex}} = 390$ nm.

block. At pH 2 and 25 °C, it is apparent that the copolymer was molecularly dissolved since all the featured signals belonging to both blocks are visible (Fig. 5A). When temperature increased to 35 °C at pH 2 (Fig. 5B), the signals of the methyl and methine protons of **PNIPAM** block at about 1.2 and 3.9 ppm were substantially suppressed, indicating that the **PNIPAM** blocks may form the less mobile and sparsely aggregated micellar cores. However, as temperature was further increased to 50 °C at pH 2 (Fig. 5C), the signals of the characteristic peaks of the **PNIPAM** block at about 1.4 and 3.9 ppm were greatly restrained, signifying the formation of the least mobile and densely aggregated **PNIPAM** micellar cores. In contrast, as the pH value is increased to 7 at varied temperatures (Fig. 5D–F), the signals attributed to the 2,2' and 3,3' protons of the pyridine ring of the **P4VP** block at about 8.5 and 7.4 ppm almost disappeared, suggesting the poorer mobility and worse solvation of **P4VP** blocks, and therefore the **P4VP** blocks preferred to form

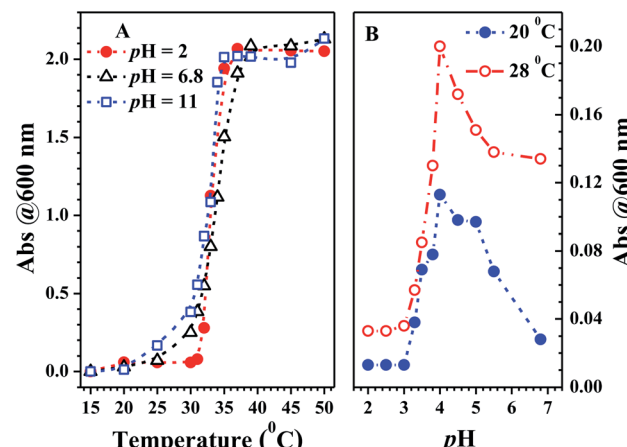


Fig. 8 Solution turbidity of PN4P in water. (A) Temperature responses at pH = 2, 6.8 and 11; (B) pH responses at 20 and 28 °C. PN4P in water at the concentration of 0.5 mg mL $^{-1}$, and turbidity data measured as a function of absorbance (Abs) at 600 nm.

micellar cores as the pH value increased. Meanwhile at pH = 7, we also found that the signals of the featured peaks of **PNIPAM** block at about 1.0–1.4 and 3.8 ppm were becoming more and more weak with temperature increase, which may suggest that the shell-forming **PNIPAM** blocks were getting more and more densely aggregated as temperature increased at pH = 7. We can infer from the combined NMR and TEM data that the copolymer could form core–shell micelles/clusters with **P4VP** cores and **PNIPAM** shells at pH = 7 (Fig. 5D–F), or micelles with **PNIPAM** cores and **P4VP** shells (Fig. 5B and C) of larger size (Fig. 3c) at 50 °C and pH = 2, or smaller-sized aggregates (Fig. 3a) at pH = 2 and 20 °C. Regardless of the pH value, the micelles/clusters became more and more aggregated and the size of the **PN4P** cluster became larger with increasing temperature, which will be proved later by the hydrated diameter data from the DLS in Section 3.2.3.

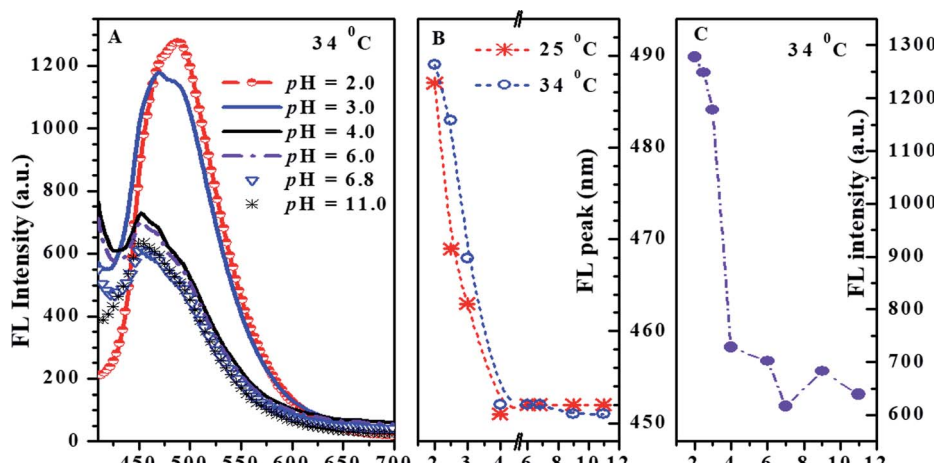


Fig. 7 (A) Emission spectra of PN4P at varied pH and 34 °C; the effect of pH on the emission peak wavelength (B) and maximum FL intensity (C) of PN4P at constant temperature. PN4P in water at the concentration of 0.5 mg mL $^{-1}$ and $\lambda_{\text{ex}} = 390$ nm.



3.2 The pH- and thermal-responsive properties and plausible emission mechanism of PN4P in water

The absorption and fluorescence of **PN4P** in water within a wide pH range (pH = 2–11) were measured, and the results are displayed in Fig. 6–8.

The absorbance spectra of **PN4P** and **4** at 25 °C can be found in Fig. 6A. The normalized pH-responsive emission spectra of **PN4P** in water at 25 °C are shown in Fig. 6B. A dramatic redshift of the emission band from 451 (pH > 4) to 487 nm (pH = 2) at 25 °C was observed in aqueous solution of the probe (Fig. 6B). In contrast, Fig. 7 displays the emission spectra at varied pH at 34 °C, the pH-dependent emission intensity and peak wavelength of **PN4P** in water. It is interesting that the pH-responsive emission peaks at 34 °C (Fig. 7B) actually tell a similar story to that for 25 °C, with a peak redshift from 451 (pH > 4) to 489 nm (at pH = 2) being recorded.

3.2.1 The pH-responsive emission mechanism of PN4P in water. As shown in Fig. 7A and B, **PN4P** exhibits a maximum emission at about 451 nm in water within the pH range of 4–11, and there was almost no peak-shift observed with the decrease in pH until pH = 4. Then, the addition of more acid into the aqueous solution resulted in enhanced emission, along with the respective new emission maxima at 468 (pH = 3) and 489 nm (pH = 2). Simultaneously, we also found that the emission intensity of **PN4P** at 34 °C decreased monotonically with the increase in pH until pH = 6.8, along with a sharp FL attenuation within the pH range of 2–4, and the maximum FL intensity at pH = 2 was enhanced by about 2.1-fold as compared with that at pH = 6.8 (Fig. 7C). One possible reason for the pH-dependent emission could be because it is reasonable to think about a deprotonated state being involved since the dye unit **4** of **PN4P** has a free phenolic OH group and water is able to support the charges. The emission enhancement along with the peak-shift could be attributed to a pH-driven switching of the deprotonation equilibrium as follows. If **PN4P**-n represents the neutral form of **PN4P** (the chromophore group **4** with the phenol unit), and **PN4P**-d denotes the deprotonated form (fluorophore group **4** with the phenoxide ion unit), the phenol unit of group **4** might offer its hydrogen ion and hydrogen-bond with water at conditions of pH > 4, and the emission peak at 451 nm could be proof of the formation of **PN4P**-d, whereas the peak at around 489 nm is allocated to the neutral form **PN4P**-n. Obviously, the formation of **PN4P**-d, the weaker fluorescent form, would induce an emission decay accompanied by a blue peak-shift (Fig. 7). The deprotonation equilibrium of **PN4P** was introduced to explain the spectra change, and the pH-responsive deprotonation equilibrium of the phenolic OH groups was anticipated to influence the absorption and emission spectra of **PN4P**. As expected, the data in Fig. 6B and 7B (25 °C) help to support the above assumption of deprotonation equilibrium. At conditions of pH < 4, mainly **PN4P**-n, the stronger fluorescent form exists, with an emission peak at about 488 nm, whereas at conditions of pH > 4, **PN4P**-d, the weaker fluorescent form occurs with emission peak at about 451 nm; *i.e.*, both **PN4P**-n and **PN4P**-d will coexist within the pH range of 4–11. In other words, the increase in solution basicity might shift the deprotonation

equilibrium towards a greater amount of **PN4P**-d, together with a decay in emission intensity within the pH range of 2–7 (Fig. 7C), which is in favor of the deprotonation-driven emission decay mechanism.

3.2.2 The critical pH and critical solution temperature (CST) of the micellization of PN4P in water. The pH responses of emission peak wavelength and solution turbidity (absorbance at 600 nm) of the copolymer **PN4P** in water at constant temperature are shown in Fig. 7B and 8B, respectively. The transition of the solution turbidity against pH (Fig. 8B) suggests a critical aggregation pH value at about 4.0, highly likely due to the deprotonation of the **P4VP** block as mentioned by Xu *et al.*²³ The effect of pH on the emission peak-shift at constant temperature (Fig. 7B) also supports a critical clustering pH value of around 4.0. The probe could form core–shell micelles in water at a certain critical temperature with a core–shell inversion at pH = 4. In other words, at the critical temperature, **PN4P** in water could self-assemble into core–shell micelles, together with a micellar structure inversion from **PNIPAM** cores and **P4VP** shells at pH < 4 to **PNIPAM** shells at pH > 4. As such, the next important task was to find the critical temperature of micellization.

From Fig. 8A, the critical solution temperature (CST) of **PN4P** was determined to be at about 32, 30 and 27 °C as the pH value switched from 2, 7 to 11, owing to the worsening solubility of the shell-forming block with increasing pH and temperature. Obviously, the solubility improvement in **PN4P** would lead to a rise in the CST.

In brief, the above data obtained from solution turbidity (Fig. 8A) and emission peak shifts (Fig. 7B) help suggest a CST around 32 °C in acid water and a critical pH value around 4.0, corresponding to a micellar core–shell inversion for **PN4P**. Therefore, from the combined NMR and TEM data, we can speculate that the copolymer could form core–shell micelles with a **P4VP** core at pH > 4.0 (Fig. 5D–F), or growing-sized micelles with a **PNIPAM** core (Fig. 5B and C) at conditions of

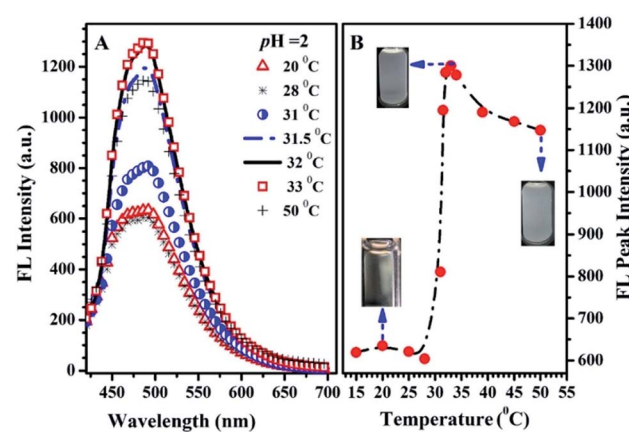


Fig. 9 The emission spectra (A) at various temperatures; the thermo-responsive emission peak intensity and solution dispersibility (B) of **PN4P** in water at pH = 2 and concentration of 0.5 mg mL⁻¹ and $\lambda_{ex} = 390$ nm.



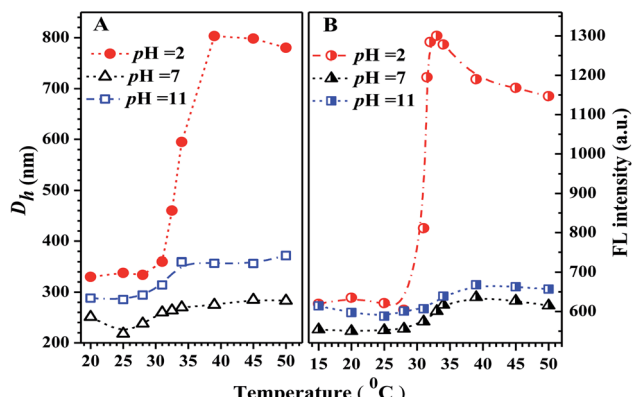


Fig. 10 Temperature responses of (A) hydration diameter (D_h) and (B) the maximum emission intensity of PN4P in water at different pH values. PN4P at the concentration of 0.5 mg mL^{-1} , and $\lambda_{\text{ex}} = 390 \text{ nm}$.

$\text{pH} < 4.0$ and $T > 32 \text{ }^\circ\text{C}$ (Fig. 3b and c) whereas smaller-sized aggregates formed at $\text{pH} < 4.0$ and $T < 32 \text{ }^\circ\text{C}$ (Fig. 3a).

3.2.3 The dual-responsive emission mechanisms of PN4P in water. The emission spectra at various temperatures and the thermal-responsive emission peak intensity of PN4P in water at $\text{pH} = 2$ are shown in Fig. 9. The temperature responses of hydration diameter (D_h) and emission peak intensity of PN4P in water at different pH are displayed in Fig. 10. As shown in Fig. 9, PN4P in $\text{pH} = 2$ water displays relatively weak emission at $<31 \text{ }^\circ\text{C}$ with two emission peaks at 488 and 451 nm being assigned to PN4P-n and PN4P-d, respectively. With the increase in temperature above $31 \text{ }^\circ\text{C}$, the emission peak at 451 nm gradually became vague while the peak at 488 nm became more dominant (Fig. 9A). As the temperature increased, firstly we saw a slight change in the FL intensity of the PN4P aqueous solution within the temperature range of 15 to $28 \text{ }^\circ\text{C}$, followed by a swift emission enhancement from 29 to $33 \text{ }^\circ\text{C}$ and then a slow FL decay until $50 \text{ }^\circ\text{C}$ (Fig. 9B). Obviously, PN4P exhibited a remarkable emission enhancement between 29 and $33 \text{ }^\circ\text{C}$, and the maximum FL intensity at $33 \text{ }^\circ\text{C}$ was enhanced 2.10-fold as compared with that at $15 \text{ }^\circ\text{C}$ (Fig. 9B). In contrast, although

a similar thermal response of emission behavior to that at $\text{pH} = 2$ was observed for PN4P in aqueous solution at $\text{pH} = 7$ and 11, (Fig. 10B), the corresponding emission enhancement against temperature was much weaker than that at $\text{pH} = 2$, and the maximum FL intensity at $33 \text{ }^\circ\text{C}$ was only enhanced 1.08- ($\text{pH} = 7$) and 1.03-fold ($\text{pH} = 11$) compared with that at $15 \text{ }^\circ\text{C}$, respectively. Furthermore, the hydration diameter (D_h) of PN4P in water at different pH values also exhibited a similar thermal-responsive trend to the corresponding emission intensity (Fig. 10). For example in aqueous solution at $\text{pH} = 2$, a mild increase in the polymer clustering size was recorded within the temperature range of 25 to $30 \text{ }^\circ\text{C}$, then a strikingly drastic size enhancement from 31 to $39 \text{ }^\circ\text{C}$ and a subsequent small size shrinkage until $50 \text{ }^\circ\text{C}$ were observed (Fig. 10A). The respective maximum clustering size D_h at $39 \text{ }^\circ\text{C}$ was about 800, 280 and 360 nm for PN4P at $\text{pH} = 2, 7$ and 11. The maximum D_h at $39 \text{ }^\circ\text{C}$ of PN4P in $\text{pH} = 2$ water was enhanced by 2.43-fold as compared with that at $20 \text{ }^\circ\text{C}$, whereas the counterparts were merely increased by 1.10- ($\text{pH} = 7$) and 1.24-fold ($\text{pH} = 11$) as compared with that at $20 \text{ }^\circ\text{C}$.

Why was it that D_h at $\text{pH} 2 > D_h$ at $\text{pH} 11 > D_h$ at $\text{pH} 7$ at comparable temperature (Fig. 10A)? The mechanism of the pH response of D_h could be as follows. Above the critical solution temperature (CST), the PN4P probe in water could self-assemble into core-shell micelles with P4VP shells at $\text{pH} 2$, but with PNIPAM shells at $\text{pH} = 7$ and 11. In aqueous solution at $\text{pH} = 7$, the minimum D_h could be due to the weakest dehydration of PN4P micelles with PNIPAM shells corresponding to the relatively strong hydrogen bonding between the acylamino N of PNIPAM and the OH group of neutral water. In aqueous solution at $\text{pH} 11$, basic water could not supply enough protons for the hydrogen bonding between the acylamino N of PNIPAM and water, thus the dehydration degree of PN4P micelles with PNIPAM shells became stronger, or the hydrogen bonding between PNIPAM and basic water became weaker as compared with that in neutral water, so we observed D_h at $\text{pH} 11 > D_h$ at $\text{pH} 7$. The maximum D_h in acid water at $\text{pH} = 2$ might be due to the strongest dehydration of PN4P micelles with P4VP shells and PNIPAM cores, where the coupling between the pyridine N of

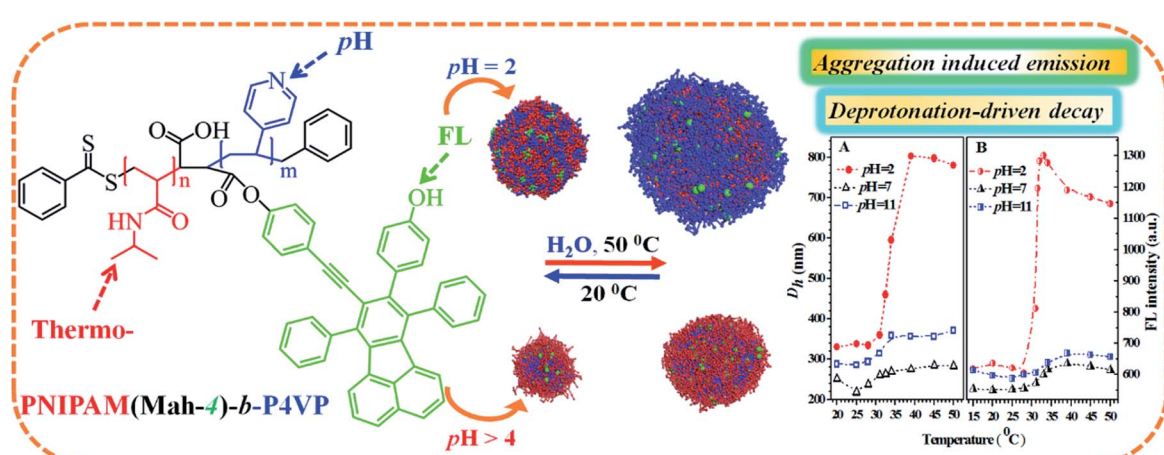


Fig. 11 The proposed pH- and thermal-responsive micellar/clustering structures and emission mechanisms of the nanoprobe PN4P in water.



the **P4VP** blocks and acid water would lead to the most intense degree of aggregation of the probe.

The possible reasons for the dual-responsive emission variation might be as follows: **PN4P** is a **PNIPAM-*b*-P4VP**-based polymer labeled with a dye unit **4** possessing AIE and deprotonation-driven emission decay (DDED) features under suitable conditions. Besides, **PN4P** can self-assemble into micellar structures in water to form core-shell-like micelles with **4** units anchored in the middle layer between the micellar cores and shells. We prefer to think that the AIE effect and the deprotonation equilibrium may be the two major factors responsible for the thermal- and pH- responsive FL changes of **PN4P** in water. Firstly, as shown in Table 2, the molar ratio of **PNIPAM** to unit **4** in **PN4P** used in this work is 15/1, implying that the **4** groups are likely separated from each other if polymer chains are in the hydrated coil state at lower temperatures. However, high temperature generally supports the polymer chains changing from coils to globules, which favors the aggregation of **4** and activates the restricted intra-molecular rotation (RIR) of the propeller moiety of fluorogen **4** and the resultant AIE phenomena. The thermally driven intra-polymer clustering of **4** units would possibly restrict the intra-molecular rotation in the fluorogen, which helps to enhance the emission of the **4** units. Secondly, in aqueous solution at pH = 2, **P4VP** block was demonstrated to be the shell-forming chain of the polymer aggregates as temperature increased. The **P4VP**-shell started to dehydrate when **PN4P** was heated to 25–30 °C, along with a gradual transition of **P4VP** chains from coil to globule and a cluster size (D_h) evolution from 338 to 350 nm (Fig. 10A). The more folded shell-forming chains may lead to the geometric confinement of **4** and may favor its RIR effect, making it more emissive (Fig. 10B). Remarkably intensive dehydration in the temperature range of 31 to 39 °C would result in the formation of more and more dense aggregates, as proven by the transition of the polymer cluster size from 360 to 803 nm from DLS data (Fig. 10A), which greatly triggered the RIR process of the **4** units and thus led to a 2.10-fold more striking emission increase of **PN4P**, revealing its AIE feature from 25 to 33 °C (Fig. 10B). At higher temperatures above the CST, the molecular motions including the vibration and intra-molecular rotation of the fluorescent group **4** would be spurred, which may gradually and mildly counterbalance the AIE effect and help to explain the observed slow emission decay above 33 °C (Fig. 9B and 10B). Meanwhile, similar thermal-responsive emission trends in water at pH = 7 and 11 also contributed to supporting the AIE feature of the probe. Thirdly, as illustrated in Section 3.2.2, **PN4P** could form core-shell micelles with a **PNIPAM** shell at pH > 4.0 or micelles with a **P4VP** shell at pH = 2, and the clustering degree of **PN4P** would become greater as temperature increases, as reflected in the DLS data shown in Fig. 10A. It seems that the **P4VP** shell is more easily dehydrated subject to temperature rise than **PNIPAM** as the shell due to the 2.43-fold enhancement in the D_h of **PN4P** clusters in water at pH = 2, whereas the respective counterpart was only increased by 1.10- (pH = 7) and 1.24-fold (pH = 11) as compared with that at 20 °C. The remarkable most intensive dehydration of the **P4VP** shell at pH = 2 as compared to that of

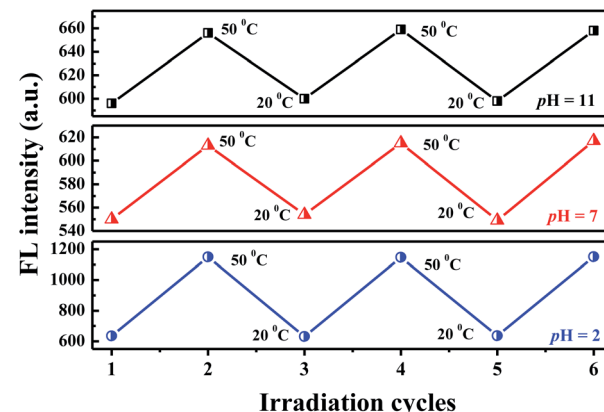


Fig. 12 FL cycling between 20 °C and 50 °C of **PN4P** in water ($\lambda_{\text{ex}} = 390$ nm, 0.5 mg mL⁻¹).

the **PNIPAM** shell at pH = 7 and 11 would make **PN4P** the most strongly aggregated and most AIE-active in acidic water (Fig. 10), which supports the 2.10-fold most distinguished thermal-responsive emission enhancement in water at pH = 2 (Fig. 10B). Furthermore, the question arises as to why the FL at pH = 11 is always greater than the FL at pH = 7 at comparable temperatures ranging from 15 to 50 °C (Fig. 10B). As shown in Fig. 8A, the respective CST of **PN4P** is about 30 and 27 °C for water solutions at pH = 7 and 11, signifying the poorer solubility or stronger dehydratation of the shell-forming **PNIPAM** block in pH = 11 solution, which would result in a denser and larger polymer clustering size at pH = 11 with increasing temperature than that at pH = 7, as proven in Fig. 10A. The denser polymer aggregation at pH = 11 would induce a stronger RIR effect of the fluorogen **4** and make it more emissive than that at pH = 7, which again favors the AIE feature of **PN4P**. As for the deprotonation-driven emission decay mechanism, there primarily exists the strongly emissive **PN4P-n** at pH = 2 while under conditions of pH = 7 and 11, both the strongly emissive **PN4P-n** and less emissive **PN4P-d** will coexist. The shift of the deprotonation equilibrium towards more **PN4P-d** at pH = 7 and 11 would lead to a decrease in the amount of **PN4P-n** and a resultant FL decay, which could clarify why the FL (pH = 2) > FL (pH = 7 and 11) at constant temperature ranging from 15 to 50 °C (Fig. 10B).

In short, although the AIE and the deprotonation-driven emission decay may be the two major factors responsible for the dual responsive emission behavior of **PN4P** in water, there may exist negligible deprotonation and the strongest AIE effect due to the most intensive dehydration of the **P4VP** shell at pH = 2, whereas there is strong deprotonation and moderate AIE at pH = 11, and then weak AIE at pH = 7, owing to the poorer dehydration of the **PNIPAM** shell at pH > 4, which could explain why the FL (pH = 2) > FL (pH = 11) > FL (pH = 7) at constant temperature ranging from 15 to 50 °C (Fig. 10B). Additionally, an increase in the solution basicity would generally shift the deprotonation equilibrium of **PN4P** towards more **PN4P-d**, according to our previous work,²⁶ but we actually observed that FL (pH = 11) > FL (pH = 7) at constant temperatures (Fig. 10B),

which obviously indicates that the AIE effect plays a more important role than the deprotonation-driven decay mechanism.

Based on the above data and discussion, we can propose a possible dual-responsive clustering/micellar structure and emission mechanism of the probe in water as displayed in Fig. 11.

3.3 Sensitivity and reversibility

The variations in the FL peak intensity of **PN4P** in aqueous solutions at constant pH are displayed in Fig. 12, where the temperature was switched repeatedly between 20 and 50 °C. At constant pH values of 2, 7 and 11, the data visibly indicate that our **PN4P** probe can adjust its intensity reversibly by temperature tuning. The reason could be as follows. The shell-forming chains might have a reversible coil-to-globule phase transition against temperature at constant pH in aqueous solution, and therefore the corresponding emission of **PN4P** behaves reversibly with temperature switching. That is, at constant pH in aqueous solution, the phase transition of the copolymer would take place reversibly, despite the cooling/heating process, thus illustrating reversible emission decay/increase along with the temperature change. It was found that the emission peak intensity at 50 °C and pH = 2 was enhanced by about 1.81-fold as compared with that at 20 °C, whereas the counterpart was increased only 1.12- (pH = 7) and 1.10-fold (pH = 11) as compared with that at 20 °C. This may be attributed to the strongest dehydration of **P4VP** as the micellar shell at pH = 2 being subject to an increase in temperature as compared to **PNIPAM** as the shell at pH = 7 and 11. The most intensive dehydration of the **P4VP** shell at pH = 2 would make **PN4P** the most strongly aggregated and the most AIE-active in acid water, favoring the strongest emission enhancement at pH = 2.

Regarding the proper use of the probe, the pH responses based on the absorption data shown in Fig. 8B were employed only to clarify the critical pH value of the probe. We read the pH response *via* the emission data of the probe since the emission data are more sensitive to pH changes as shown in Fig. 7C, which shows a monotonic variation in emission intensity against pH within the pH range of 2 to 7, indicating that **PN4P** is a good pH sensor within the pH range of 2 to 7.

Combined with the data shown in Fig. 7C, Fig. 10B and 12, we are sure that the probe shows the strongest sensitivity at pH = 2, then at pH = 7 and the weakest sensitivity at pH = 11. The probe could, therefore, be a promising sensor that is suitable for working within the pH range of 2 to 7 (Fig. 7C) and the temperature range from 28 to 40 °C (Fig. 10B), simultaneously with very strong pH sensitivity, especially in the pH range of 2–4 (Fig. 7C).

4. Conclusions

We developed a new fluorescent pH- and thermo-meter, **PN4P**, in this work. The micellar/clustering structures of the probe in water were investigated by solution dispersibility, NMR, TEM, DLS, FL and UV-vis measurements, and the emission

performance was systematically examined by temperature and pH tuning.

We illustrated the plausible dual-responsive micellar structures and emission mechanisms of the probe in water. The critical solution temperature of the probe in water is about 32 °C (pH = 2), 30 (pH = 7) and 27 °C (pH = 11). The critical pH value of the probe in aqueous solution was found at about 4, and the micelles showed a core–shell inversion in response to pH and thermal stimuli accompanied by remarkable emission tunability. Acidochromic changes in the emission band from 450 (pH > 4) to 490 nm (pH = 2) was observed in water, and the probe showed the best sensitivity at pH = 2. **PN4P** demonstrates obvious deprotonation-driven emission decay and AIE features in different microenvironments and displays considerable temperature and pH responses, as well as good reversibility. The probe is more sensitive within the pH range of 2 to 7, and the temperature range from 28 to 40 °C. Therefore, we see that our probe is suitable for working in a temperature and pH range that is interesting for biomedical applications, and therefore, we can expect that our **PN4P** probe is suitable for use as a biosensor or in the biomedical field, due to the suitable temperature and pH responsive range.

Conflicts of interest

There are no conflicts of interest to declare.

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