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Nano-visualization of Hydrogel Dynamics via Surface Plasmon-Enhanced Aggregation-Induced Emission

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Understanding conformational transitions in hydrogels is essential for deciphering their stimulus-responsive behavior, yet real-time, label-free monitoring at the nanoscale has remained a fundamental barrier. Here, we introduce surface plasmon coupling aggregation-induced emission (SPCAIE) as a new observational paradigm that enables in situ visualization of hydrogel dynamics with sub-nanometer axial displacement sensitivity. Our approach establishes a synergistic, label-free feedback loop between molecular conformation and optical output by merging the interfacial sensitivity of plasmonics with the microenvironmental responsiveness of non-covalently embedded AIE luminogens. This new transduction mechanism directly converts intrinsic nanoscale motions into quantifiable, amplified optical signals, overcoming the limitations of conventional invasive or ensemble-averaging techniques. The SPCAIE framework not only reveals previously inaccessible stimulus-induced rearrangements in hydrated environments, but also establishes a generalizable principle for probing soft matter interfaces. Our work provides a transformative conceptual and methodological platform for analyzing the dynamics of hydrated soft materials, with broad applicability to diverse stimulus-responsive polymer systems, opening new pathways in soft matter research and advanced material design.

Introduction

In recent decades, polymer and gel materials have experienced rapid development and widespread adoption, with advanced hydrogels emerging as particularly promising candidates. These materials exhibit exceptional tensile properties, including remarkable strain capacity, attributable to their distinctive hybrid network architectures. Furthermore, their mechanical performance can be precisely tailored by modulating the composition, concentration, and relative ratios of constituent components. Hydrogels represent a class of cross-linked polymeric materials featuring three-dimensional network structures that demonstrate exceptional water absorption capacity while preserving structural integrity without dissolution. Through sophisticated molecular design, researchers have developed stimuli-responsive hydrogels capable of sensing and responding to various environmental cues, including temperature fluctuations, pH variations, mechanical pressure, light exposure, electric fields, and magnetic fields.¹⁻⁷ These unique characteristics have propelled

their extensive utilization across diverse applications, ranging from controlled drug delivery systems and tissue engineering scaffolds to advanced biosensing platforms.⁸⁻¹³

The swelling behavior of hydrogels—characterized by their reversible transition between collapsed and extended states and quantified through both equilibrium swelling ratios and kinetic swelling rates—represents a fundamental property governing hydrogel performance.¹⁴ Precise characterization of these swelling dynamics is critical for the rational design of hydrogels tailored to specific applications.¹⁵ Since swelling dynamics are intrinsically linked to dimensional changes, various analytical techniques have been developed to capture these variations, particularly in film thickness. Current techniques for probing hydrogel swelling, a key conformational transition, are largely limited to ex-situ methods. Conventional approaches include spectroscopic ellipsometry,¹⁶ scanning electron microscopy (SEM),¹⁷ atomic force microscopy (AFM),¹⁸ nuclear magnetic resonance,¹⁹ and neutron reflectometry.²⁰ However, accurate nanoscale characterization of hydrogel thickness presents significant experimental challenges.²¹ These ex-situ techniques often require sample dehydration or complex modeling, which can alter or infer rather than directly capture hydrated, dynamic states. For instance, SEM requires freeze-drying or critical point drying to prevent the structural collapse of the three-dimensional hydrogel network, while AFM imaging of hydrated soft materials demands exceptional tip performance to minimize imaging artifacts, and neutron scattering requires extensive data modeling and either prior structural knowledge or explicit assumptions about the hydrogel architecture.

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Beyond those ex-situ methods, several research efforts, including our own, have employed fluorescence-based techniques for in situ investigation of nanoscale environments within polymer materials.²²⁻²⁵ Förster Resonance Energy Transfer (FRET)-based polymer brushes can monitor conformational transitions through specifically integrated donor-acceptor pairs, but they report only on intra- or inter-molecular distances within a narrow range (1–10 nm) and require covalent labeling, which limits material generality.²² Autofluorescent polyelectrolytes avoid exogenous probes yet are restricted to specific polymer chemistries that exhibit native fluorescence, severely narrowing their applicability.²³ Our group previously introduced the angle scanning-surface plasmon coupled emission (AS-SPCE) technique for real-time monitoring of the growth process of fluorophore-labeled polymers;²⁴ however, SPCE alone still requires covalent labeling and, without an environmentally sensitive luminogen, cannot report on local chain packing or hydration independently of the emitter-interface distance. These techniques thus face a fundamental dilemma: they rely on specific, covalently attached fluorescent reporters that constrain material scope and introduce an unavoidable compromise between signal generation and measurement fidelity, as the label may perturb the native properties and interactions of the polymer. Thus, a transformative platform that combines non-invasive operation, nanoscale spatial resolution, and universal material compatibility has remained an outstanding challenge.

To overcome these challenges, we present surface plasmon coupling aggregation induced emission (SPCAIE), a strategy that integrates the axial ranging capability of SPCE with the microenvironmental sensitivity of AIE into a single measurement. The SPCE phenomenon arises from the interaction between fluorescent molecules and surface plasmon polaritons at the metal-dielectric interface,²⁶⁻²⁹ providing exceptional spatial sensitivity through the exponential decay of plasmonic coupling intensity.³⁰⁻³⁴ This SPCE platform can precisely quantify fluorophore displacement distance normal to the interface by analyzing the angular distribution and polarization state of the SPCE signal. Meanwhile, AIE luminogens (AIEgens) exhibit a photophysical behavior where molecular aggregation under a microenvironment restricts intermolecular motions (e.g., rotation and vibration), resulting in greatly enhanced fluorescence.³⁵⁻⁴⁰ The synergistic combination of these two effects in our work establishes the first non-invasive approach for monitoring polymer conformational transitions with simultaneous real-time temporal resolution and sub-nanometer axial displacement sensitivity. This work establishes SPCAIE not merely as a new technique, but as a conceptual and methodological framework that redefines the observation of soft matter interfaces. SPCAIE induces a label-free feedback loop: it leverages the material's intrinsic conformational change to modulate an AIE process, the emission of which is then dramatically amplified and spatially confined by surface plasmons. This direct, simultaneous delivery of both spatial and environmental information from an unlabeled sample represents a unique conceptual advance not achieved by

previous techniques. This synergy directly transduces nanoscale mechanical motions into a quantitative, macroscopic optical readout with sub-nanometer axial displacement sensitivity, circumventing the traditional trade-off between non-invasiveness and sensitivity.

Results and Discussion

As a proof-of-concept demonstration, we selected pH-responsive poly(acrylic acid)/poly(vinyl alcohol) (PAA/PVA) hydrogel (pKa \approx 4.5)^{41, 42} as a model system. The AIEgens triphenylaminothiophene pyridine (TTPy) was facilely incorporated into the hydrogel matrix through simple mixing without covalent modification. Employing our AS-SPCE microscopy platform in the Kretschmann configuration, we successfully monitored hydrogel thickness variations in real time during pH stimulation by characterizing the SPCAIE response (Figure 1). Below the pKa, the collapsed hydrogel state restricts the intramolecular motion of TTPy, while above the pKa, swelling leads to the restoration of intramolecular motion of TTPy within the polymer network. These structural transitions occur within the decay length of the interfacial evanescent field in SPCE. Remarkably, the system exhibits dual signal amplification mechanisms: first through distance-dependent SPCE effect that enables nanoscale thickness sensitivity at the interface, and second via the AIE effect that responds to microenvironmental changes during TTPy aggregation driven by polymer conformational transitions. The SPCAIE approach overcomes fundamental limitations of conventional methods by synergistically combining the nanoscale spatial precision of SPCE with the exceptional microenvironmental responsiveness of AIEgens, establishing a label-free, real-time paradigm for studying soft matter dynamics. The non-covalent incorporation of AIEgens offers material versatility beyond the PAA/PVA hydrogel system demonstrated here. We anticipate broad applicability of this platform across diverse stimuli-responsive polymers, enabling new opportunities in biological sensing, smart material development, and fundamental studies of polymer dynamics at the nanoscale.

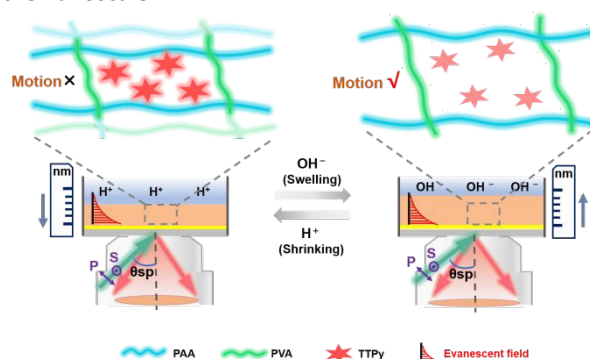


Figure 1. Schematic illustration of real-time nanoscale monitoring of hydrogel conformational dynamics via surface plasmon-coupled aggregation-induced emission. A pH-responsive poly(acrylic acid)/poly(vinyl alcohol) (PAA/PVA)



hydrogel incorporated with a triphenylaminothiophene pyridine (TTPy) molecular probe serves as the model system.

The architectural configuration of the TTPy-incorporated hydrogel and the experimental setup of the AS-SPCE microscopy are illustrated in Figure 2a, which enables reliable modulation of both polarization and excitation angle (see Figure S1 for instrument details). TTPy was synthesized according to established methods⁴³ and characterized as shown in Figure S2. Fluorescence measurements confirmed its AIE characteristics (Figure S3). Efficient SPCE coupling is facilitated by the spectral overlap between the emission band of TTPy and the surface plasmon resonance (SPR) of the underlying gold substrate.^{31, 44} Structural analyses via FT-IR spectroscopy confirmed the successful formation of a cross-linked hydrogel network within the PAA/PVA matrix and importantly indicated that the incorporation of TTPy did not perturb the intrinsic chemical properties of the polymer blend (Figure 2b). Furthermore, SEM imaging revealed a smooth and homogeneous morphology of the hydrogel (Figure 2c), supporting reliable thickness characterization. The SPCAIE technique allows precise characterization of the fluorescent hydrogel film thickness through analysis of excitation angle distribution patterns, particularly leveraging the directional and polarization-dependent nature of SPCE. Validation studies demonstrated excellent agreement between SEM cross-sectional

measurements (311 nm dry-state thickness) and SPCAIE-derived thickness values (see Figure S4 for details), strongly underscoring the reliability and precision of this optical methodology. This integrated platform is particularly powerful for monitoring hydrogel swelling dynamics with nanoscale resolution, owing to its intrinsic compatibility with liquid-phase environments, thereby enabling real-time observation of polymer conformational changes under physiologically relevant hydrated conditions.

As demonstrated in Figure S5a, incubation of the hydrogel in buffer solution (pH 6.0) yielded distinct optimal excitation angles—66.0° for p-polarization and 68.9° for s-polarization under laser excitation—corresponding to a swollen thickness of 498.5 nm. Variations in the solution pH (6.0, 6.5, and 6.8) induced measurable and systematic shifts in the angular distributions: the p-polarized angle shifted from 64.7° to 65.5°, whereas the s-polarized angle remained constant at 68.9° (Figures 2d-e). These angular shifts correspond to a progressive increase in film thickness from 498.5 nm to 503.3 nm (Figure S6). A pivotal advantage of polarization-resolved SPCAIE is its capacity to provide deep molecular-level insight into the hydrogel's structural reorganization during pH-induced conformational transitions. By simultaneously capturing emissions from perpendicular (p-polarized) and parallel (s-polarized) molecular dipoles relative to the metal interface,

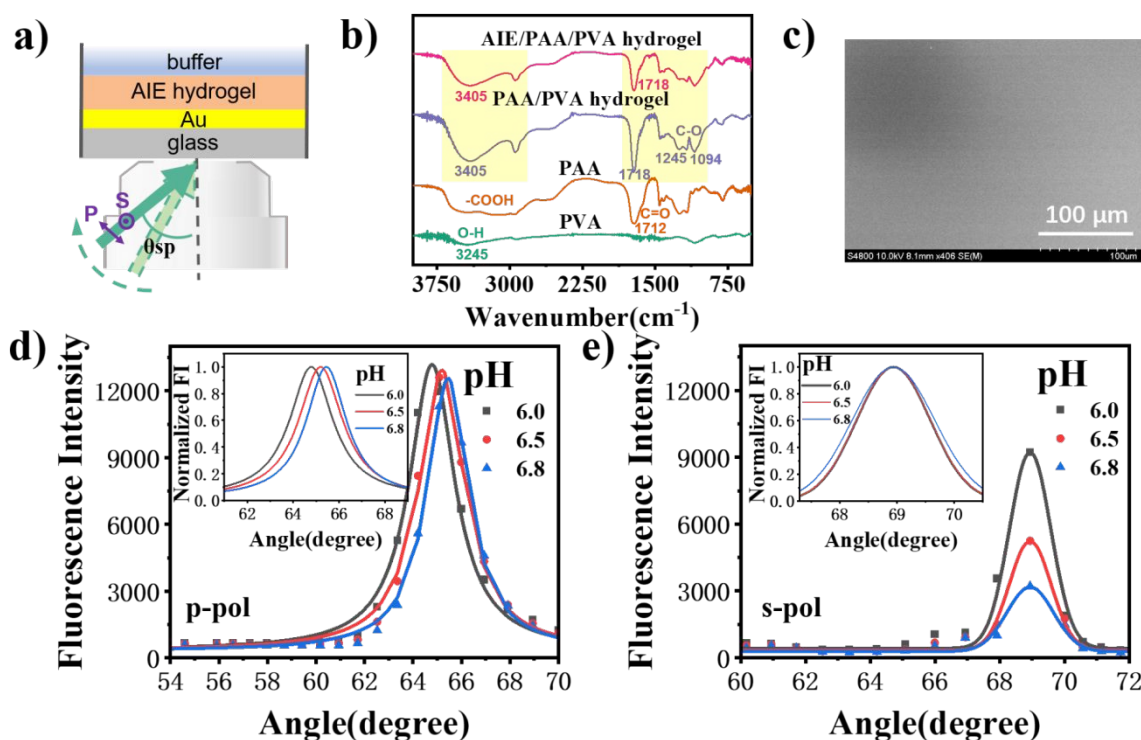


Figure 2. Characterization of the AIE-incorporated hydrogel chip and its SPCE sensing performance. (a) Schematic of the SPCE experimental setup used for real-time hydrogel analysis. (b) FT-IR spectra of pure PAA, pure PVA, PAA/PVA hydrogel, and TTPy/PAA/PVA hydrogel, showing characteristic peaks at 1094 cm^{-1} , 1245 cm^{-1} , 1718 cm^{-1} , and 3405 cm^{-1} , confirming successful network formation. (c) SEM image of the TTPy/PAA/PVA hydrogel film spin-coated on a gold-coated coverslip (30 nm Au). (d) p-Polarized and (e) s-polarized excitation angular distributions measured by the AS-SPCE system at pH 6.0, 6.5, and 6.8, with incident angle scanned from 0° to 80°. Symbols represent experimental data; solid lines denote fitting results; insets display normalized fluorescence intensity as a function of incident angle.



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SPCE affords unique three-dimensional structural resolution,⁴⁵ which extends far beyond the thickness-only information accessible via conventional techniques such as ellipsometry.

The polarization-dependent responses offer particularly valuable insights into molecular orientation and dipole alignment within the swelling hydrogel. The more pronounced decrease in *s*-polarized SPCE intensity suggests significantly greater mobility and increased spatial separation among TTPy dipoles oriented parallel to the substrate surface during swelling. This anisotropic emission response reflects constrained swelling perpendicular to the film plane, with dimensional changes occurring predominantly along the thickness direction in spatially confined hydrogel films.^{46, 47} The persistence of *s*-polarized angle, coupled with intensity reduction, indicates that in-plane dipoles aligned parallel to the interface experience fewer restrictions and undergo more substantial rearrangement, whereas vertically aligned (*p*-polarized) out-of-plane dipoles are relatively constrained and

more strongly influenced by the film's vertical expansion. Such discrimination between in-plane and out-of-plane molecular motions is not accessible by conventional thickness measurements (e.g., ellipsometry or profilometry), highlighting the unique capability of SPCAIE to probe anisotropic chain dynamics within the hydrogel network. Given the enhanced sensitivity of *s*-polarized excitation to in-plane dipole rearrangements and its pronounced intensity response during swelling, it was selected as the standard mode for all subsequent experiments, providing optimal resolution for tracking molecular-scale reorganization during hydrogel transitions.

We propose that the SPCAIE signal arises from a synergistic combination of the aggregation-induced emission (AIE) effect and surface plasmon-coupled emission (SPCE). To decouple these contributions, we designed two control systems: a TTPy-incorporated hydrogel on a non-plasmonic glass substrate (exhibiting AIE only; Figure 3d) and a conventional Rhodamine

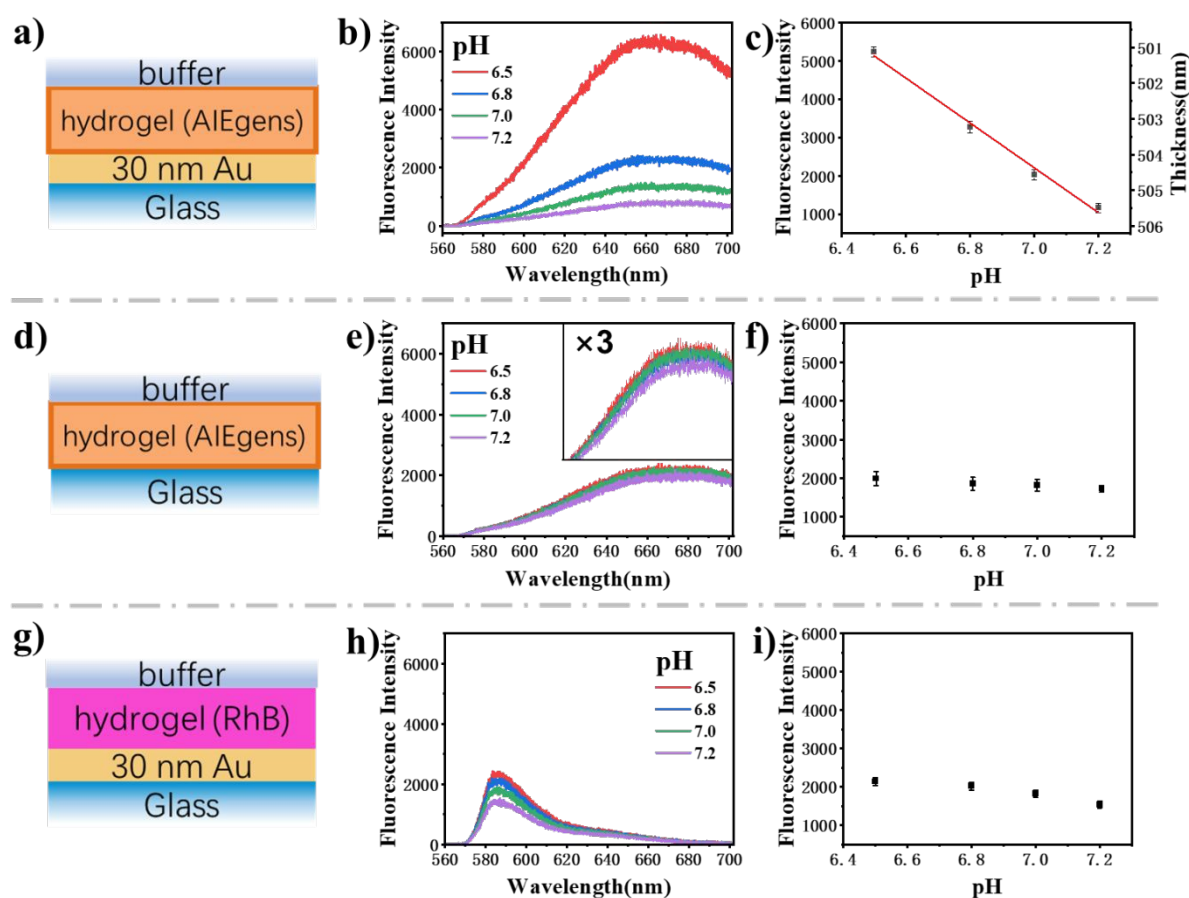


Figure 3. Comparative analysis of the response characteristics for three systems. pH values of 6.5, 6.8, 7.0, and 7.2 were used for demonstration. (a–c) AIEgen-doped hydrogel on a gold substrate (combining both SPCE and AIE effects); (d–f) AIEgen-doped hydrogel on a non-plasmonic glass



substrate (AIE effect only); and (g–i) RhB-doped hydrogel on a gold substrate (SPCE effect only). Each row presents schematic illustrations (a, d, g), corresponding emission spectra (b, e, h), and fluorescence intensity as a function of pH and film thickness ($N=4$) (c, f, i). DOI: 10.1039/D6SC00713A

B (RhB)-doped hydrogel on a gold film (exhibiting SPCE only; Figure 3g). Using our custom-built dual-mode AS-SPCE platform, which supports simultaneous spectral acquisition and spatial imaging, we systematically evaluated the fluorescence responses of all three systems across a range of physiologically relevant pH conditions (Figures 3b, e, h). The PAA-based hydrogel exhibits pH-responsive swelling behavior governed primarily by the deprotonation of carboxyl groups along the polymer backbone. This process disrupts interchain hydrogen bonding and introduces electrostatic repulsion among ionized carboxylate groups, leading to network expansion.^{48,49} Although these mechanisms collectively promote progressive swelling across the physiological pH range (6.5–7.4), the resultant volumetric changes remain constrained by the cross-linked network structure. Crucially, the limited swelling magnitude proves insufficient to significantly alter either the spatial distribution of TTPy molecules within the evanescent field region (which critically governs SPCE coupling efficiency) or the molecular packing state of the AIEgen (which fundamentally dictates AIE intensity). Consequently, both control systems exhibited only modest fluorescence variations under identical pH stimuli (Figures 3d, g).

In contrast, the SPCAIE system demonstrated dramatically enhanced signal modulation, exceeding the combined contributions of either AIE or SPCE operating independently (Figure 3c). Despite the relatively limited absolute swelling degree, the system exhibited a responsivity slope more than 50-fold steeper than that of the control system on a glass substrate, endowing it with exceptional detection sensitivity. SPCAIE imaging further confirmed highly uniform signal distribution throughout the hydrogel matrix (Figure S7), indicating that the AIEgens maintain stable integration within the polymer network without leaching during swelling. The absence of observable spectral shifts across all conditions (Figures 3b, e) suggests that fluorescence intensity changes primarily stem from restricted intramolecular motion of the AIEgens, rather than from alterations in their aggregation state. This is supported by the control experiment (Figure 3e), in which an AIE-active film (prepared under AIE-inducing conditions) exhibited only a weak pH response, confirming that the strong signal modulation observed in SPCAIE originates from the synergistic coupling effect rather than from the AIEgen's inherent sensitivity alone.

The exceptional sensitivity of SPCAIE is rooted in a synergistic dual-amplification mechanism, which constitutes the core conceptual advance of this work. This mechanism operates through a cascading process: first, the nanoscale swelling or contraction of the hydrogel network directly modulates the intramolecular motion (rotation/vibration) of the non-covalently embedded AIEgens. Second, this AIE emission is not passively collected but is actively and selectively amplified by SPCE effect. Specifically, the s-polarized SPCE component enhances emission from dipole moments oriented parallel to

the metal interface, providing exquisite interfacial specificity and distance-dependent sensitivity. Critically, these two processes are not sequential but are intrinsically coupled, creating a coherent, label-free feedback loop. This loop directly and dynamically transduces minute conformational changes into a strongly amplified, quantifiable optical signal with exceptional signal-to-noise ratio and spatial precision, overcoming the historic trade-off between sensitivity and non-invasiveness in interfacial analysis. Owing to this foundational mechanism, the SPCAIE platform exhibits broad material compatibility and robust operability in liquid environments, establishing a generalizable paradigm for real-time interfacial science. Its utility extends far beyond the model hydrogel system, positioning it as a promising tool for real-time monitoring within complex chemical and biological settings, including biomarker detection, cellular microenvironment mapping, and dynamic analysis of soft matter transitions.

To demonstrate its capability in resolving nanoscale dynamics, we characterized the hydrogel's real-time switching behavior through alternating exposure to acidic (pH 1.0) and neutral (pH 7.0) solutions. Transition from pH 1.0 to 7.0 was initiated by applying a microdroplet of acidic solution to the hydrogel surface, followed by rapid infusion of neutral buffer. The reverse transition (pH 7.0 to 1.0) was performed using an analogous exchange protocol. Real-time conformational dynamics were monitored via SPCAIE signal acquisition, with simultaneous recording of excitation angular distributions for comprehensive analysis (Figure S9). As quantitatively summarized in Figure 4a, the hydrogel exhibited rapid and reversible pH-responsive behavior. Both switching directions achieved detectable thickness changes within 1 minute and reached equilibrium within 3 minutes after pH alteration. Complementary SPCAIE imaging (Figure 4b) provided spatial insight into swelling/deswelling dynamics, enabling direct correlation among fluorescence intensity, angular distribution profiles, and temporal evolution.

Notably, shifts in angular distributions—reflecting thickness variations on the tens-of-nanometer scale—were accompanied by synchronous fluorescence intensity changes sensitive to even subtler structural rearrangements. These dual optical signatures yielded distinct kinetic profiles for swelling and deswelling processes. Together, the results demonstrate the hydrogel's rapid response kinetics and high structural stability over multiple pH cycles. We attribute the fast-switching behavior to the highly hydrated nature of the nanoscale hydrogel film, which enables rapid hydration triggered by deprotonation (swelling) and efficient water expulsion during protonation (collapse), facilitating efficient conformational transitions between collapsed and swollen states.

To elucidate the conformational changes in detail, we systematically investigated the equilibrium swelling behavior of the hydrogel across a range of pH conditions. Quantitative analysis of both angular distribution and fluorescence intensity



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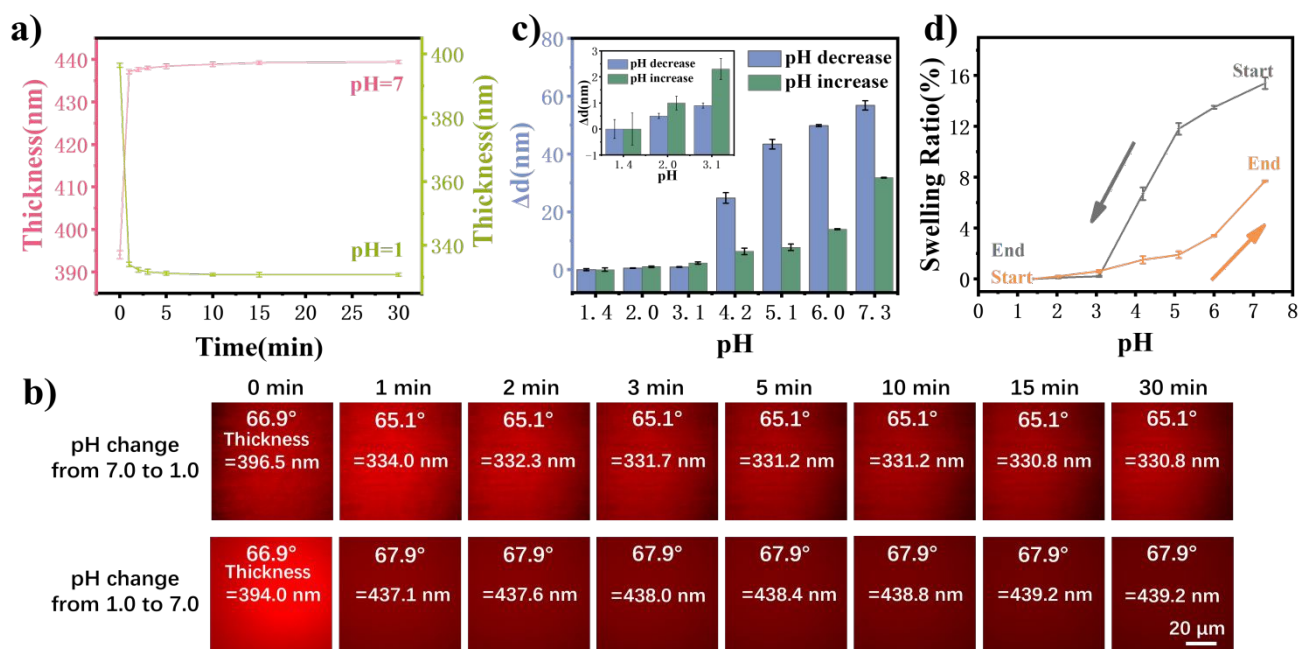


Figure 4. pH-dependent thickness variations of PAA/PVA hydrogel characterized by SPCAIE. (a) Real-time monitoring of hydrogel transient response during pH transitions between acidic (pH 1.0) and neutral (pH 7.0) conditions. Pink squares and olive circles separately denote transitions starting at pH 1.0 and at pH 7.0. (b) Corresponding fluorescence micrographs documenting the dynamic swelling/deswelling process. (c) Quantified thickness dynamics during pH cycling (the magnified inset revealing sub-nanometer resolution). pH-dependent transitions are represented by blue columns (pH 7.3 to 1.4, deswelling) and green columns (pH 1.4 to 7.3, swelling). (d) pH-cycling behavior showing swelling ratio hysteresis: gray curve represents neutral-to-acidic transition (pH 7.3 to 1.4); orange curve represents the reverse transition. All error bars represent mean \pm standard deviation (n=3). SPCE angles scanned from 0° to 80°; film thickness was obtained by fitting the angular reflectivity curves using a four-layer Fresnel model. pH was adjusted using buffer solutions.

variations was performed during reversible transitions between acidic and neutral environments (Figure S10). The results revealed asymmetric dimensional changes: swelling from pH 1.4 to 7.3 induced a 32 nm increase in thickness, while the reverse transition (pH 7.3 to 1.4) resulted in a more pronounced contraction of 57 nm (Figure 4c). Since the hydrogel film is confined on a planar substrate, dimensional changes are restricted to the thickness direction (z-axis), enabling precise determination of the swelling ratio (SR) according to:

$$SR(\%) = \left(\frac{d_t - d_0}{d_0} \right) \times 100$$

where d_0 and d_t denote the hydrogel film thickness at reference (pH 1.4) and test conditions during either forward or reverse pH transitions, respectively.

The pH-dependent swelling behavior is summarized in Figure 4d. During the forward transition (pH 1.4 \rightarrow 7.3), the SR increased continuously, with especially pronounced swelling above the pK_a of PAA. This behavior is attributed to carboxyl group ionization, where enhanced electrostatic repulsion promotes hydration, leading to marked hydrogel expansion. In

contrast, the reverse transition (pH 7.3 \rightarrow 1.4) exhibited a more complex SR profile, which can be divided into three distinct regions: (1) Between pH 7.3 and 5.1, the hydrogel retained higher SR values relative to the fully collapsed state, showing gradual deswelling that reflects relaxation of the pre-expanded network; (2) Near the pK_a region (pH 5.1–4.2), a rapid decrease in SR occurred due to reduced electrostatic repulsion and constraints from network elasticity as carboxyl groups became protonated; (3) Below pH 3.1, complete protonation eliminated interchain repulsion, causing the SR to stabilize at a minimum value.

Notably, even in this plateau region, SPCAIE detected sub-nanometer thickness variations, confirming that the fully protonated state exhibits negligible swelling compared to the reverse transition (Figure 4c, inset). The observed swelling–deswelling hysteresis—where forward and reverse pH transitions follow distinct SR pathways—is a characteristic behavior of ionic hydrogels, previously reported only at macroscopic scales.⁵⁰ In contrast to traditional methods that infer such hysteresis from ensemble-averaged bulk responses, SPCAIE directly resolves its nanoscale topography and dynamics

in real time. To our knowledge, this work provides the first quantitative, in situ visualization of asymmetric nanoscale hydrogel dynamics with sub-nanometer axial displacement sensitivity, thereby bridging the critical gap between macroscopic property measurements and molecular-scale interactions. The SPCAIE technique thereby establishes a conceptually straightforward yet methodologically powerful platform for non-invasive, real-time interrogation of soft materials. Its superior z-axis spatial resolution, surpassing the diffraction limit and reaching the nanoscale, places it beyond the reach of conventional optical methods.⁵¹

Conclusions

In summary, this work establishes SPCAIE as a transformative paradigm for probing soft matter interfaces. By synergistically integrating the interfacial sensitivity of SPCE with the microenvironmental responsiveness of non-covalently embedded AIEgens, the SPCAIE platform not only achieves real-time, label-free visualization of nanoscale dynamics with exceptional spatiotemporal resolution, but also ensures non-invasiveness, experimental simplicity, and full compatibility with aqueous environments. The capability of this paradigm is decisively demonstrated through the resolution of a long-standing challenge: the first quantitative, in situ characterization of nanoscale swelling–deswelling hysteresis in ionic hydrogels. Moving beyond macroscopic averaging, SPCAIE captures this hysteresis with high fidelity, revealing the underlying interplay between network elasticity and electrostatic repulsion. It directly visualizes the suppression of swelling under protonated, hydrogen-bond-dominated conditions, providing mechanistic insight previously inaccessible.

Beyond these fundamental advances, the SPCAIE platform offers broad methodological versatility. Its high sensitivity, operational simplicity, and compatibility with hydrated environments position it as a powerful tool for studying hydration-mediated processes in fields such as controlled drug delivery, soft actuators, and bio-integrated sensing. The underlying transduction principle, based on synergistic coupling of evanescent-field sensitivity with microenvironment-responsive emission, is generalizable to systems that satisfy three necessary conditions: optical transparency at the excitation and emission wavelengths, responsive conformational changes occurring within the evanescent-field penetration depth, and compatibility with non-covalent incorporation of AIEgens. The approach is readily adaptable to diverse stimuli-responsive polymers and can be extended to complex biological milieus—from polymer brushes and multilayer films to biomimetic membranes. In practice, successful translation also requires that the AIEgen remains stably entrapped without leaching over repeated stimulation cycles and that its loading does not perturb the intrinsic responsiveness of the matrix. Representative candidate systems include thermo-responsive polymers, polyelectrolyte brushes, and biopolymer networks. Thus, SPCAIE emerges not merely as an analytical technique, but as a generalizable

framework for probing dynamic interfaces, paving the way for next-generation soft-matter design and interfacial science.

Author contributions

Q. B. L., Z. L., S. H. C., and Y. Q. L. designed the study and wrote the manuscript. Q. B. L. and Z. L. performed the majority of the experiments and analyzed the data. Y. H. W., L. C., and J. D. W. assisted with the fabrication and characterization of hydrogels. T. B. W. provided guidance on the preparation of AIE dyes. S. H. C. and Y. Q. L. conceived and supervised the project.

Conflicts of interest

There are no conflicts to declare.

Data availability

The authors declare that all the data are available within the article and its supporting information or from the corresponding author upon reasonable request.

Acknowledgements

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Data availability statement

The authors declare that all the data are available within the article and its supporting information

or from the corresponding author upon reasonable request.

