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Functionalized two-dimensional nanochannel membranes to distinguish methylated/unmethylated peptides for sensing cellular G9a protein

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Histone methyltransferase G9a-catalyzed histone modifications serve as critical epigenetic marks that regulate diverse cellular functions. Aberrant enzymatic activities of G9a are closely associated with various human pathologies, making them promising biomarkers for disease diagnosis and therapeutic targeting. However, the lack of efficient assays for accurate and sensitive detection of G9a, especially in complex crude cellular extracts, limits elucidation of cancer-related mechanisms and advancements of therapeutic innovations. Here, a G9a detection system based on functionalized 2D solid-state nanochannel membranes through the specific recognition between small molecule Crown probes and dissociative peptide probes was developed. The Crown probe exhibited a strong binding force with the unmethylated peptide through hydrogen bonding, thereby altering the ion transport characteristics within the nanochannels and enabling the differentiation of methylated and unmethylated peptides followed by the detection of histone methyltransferase G9a. The excellent sensing performance enables diverse detection scenarios, including differentiation of different subtypes of breast cancer cells, screening of enzyme inhibitors and sensing of hypoxic cellular environments. This research proposed that the GO-Crown membrane could be an effective detection system for G9a, which provides a powerful tool for cancer research and clinical application.

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Introduction

As one of the most important post-translational modifications, protein methylation has attracted much attention because of its important role in many biological processes such as DNA repair, RNA processing, transcriptional regulation, and protein phase separation.^{1–3} Methylation mostly occurs on transcription factors and histones.^{4,5} With the assistance of histone methyltransferases such as G9a (also known as EHMT2), histone methylation primarily takes place on the N-terminal tail regions of histone H3 and H4.^{6,7} Utilizing *S*-adenosyl-*L*-methionine (AdoMet) as a methyl donor, G9a catalyzes the addition of one or multiple methyl groups to lysine residues (*e.g.*, H3K9), forming mono-, di-, or trimethylation modifications, which are closely related to the formation and maintenance of chromatin structure, silencing or activation of gene expression, *etc.*^{8–11} The aberrant expression of G9a is closely related to the occurrence

and development of a variety of diseases. In breast cancer cells, aberrantly expressed G9a leads to gene silencing through hypermethylation of the tumor suppressor gene promoter region H3K9, thereby promoting tumorigenesis and progression.^{12,13} In addition, G9a is also highly expressed in many other cancers compared to expression in normal tissues, including human bladder cancer, lung cancer, colon cancer, *etc.*^{14–16} From a diagnostic point of view, changes in the content of G9a can serve as a potential biomarker for the diagnosis of cancer described above. In terms of treatment, G9a can also be an important therapeutic target. For cancer cells with high expression of G9a, the development of specific inhibitors can effectively block their abnormal methylation regulation, thereby inhibiting the growth and spread of cancer cells, which can provide new ideas and methods for precision treatment of cancer.^{17,18} Therefore, the development of G9a detection tools is crucial for elucidating cancer-related mechanisms and advancing therapeutic innovations.

Current methods for detecting histone methyltransferases predominantly rely on traditional spectrophotometry,¹⁹ mass spectrometry methods,²⁰ *etc.*, most of which involve radio-labeling, indirect detection of by-product (*S*)-adenosyl-*L*-homocysteine (AdoHcy),²¹ the use of specific antibodies,¹⁹ and, more

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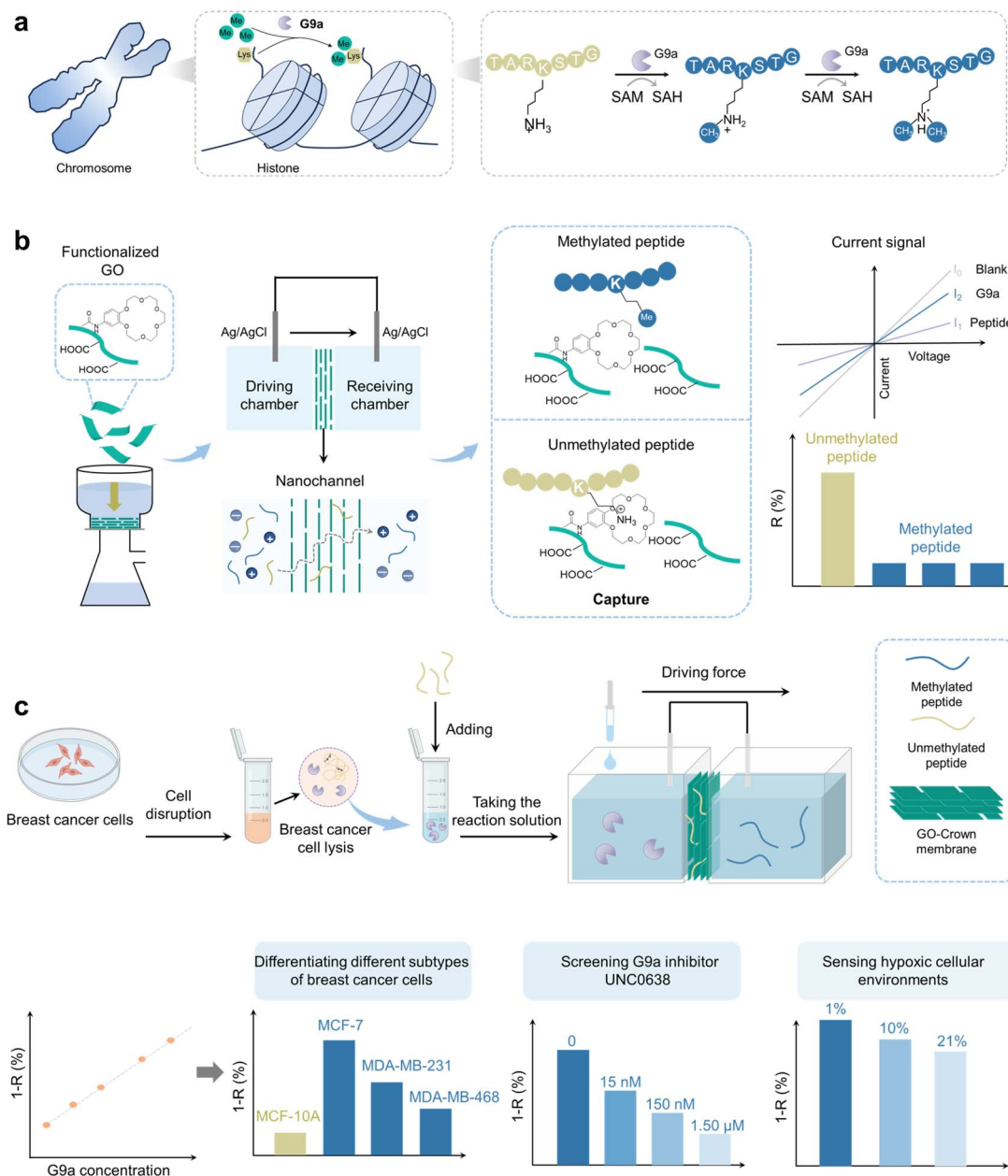


Fig. 1 Functionalized 2D nanochannel membranes to distinguish methylated/unmethylated peptides for sensing cellular G9a protein. (a) Schematic illustration of histone lysine methylation. (b) The functionalized 2D solid-state nanochannels based on the GO-Crown membrane could distinguish methylated/unmethylated peptides through the strong binding force between the Crown probe and unmethylated peptide, which induces a transmembrane current change. (c) The detection procedure of a G9a detection system. The detection is based on the specific recognition between small molecule Crown probes that anchored on the nanochannels and dissociative unmethylated peptide probes, which could be methylated by G9a. This system can be used in diverse scenarios, including differentiation of different subtypes of breast cancer cells, screening enzyme inhibitors and sensing hypoxic cellular environments.

recently, supramolecular replacement methods.²² While some existing approaches exhibit high sensitivity, they suffer from limitations including multistep immobilization or separation procedures, necessary post-reaction treatments, safety concerns about the use of radiotracers, interference from background fluorescence, and high costs.^{23,24} What's more, given G9a's potential as a tumor biomarker, few methods are capable of directly analyzing G9a in complex crude cellular extracts

without prior purification or separation. There remains an urgent need for a label-free, highly selective, sensitive, and simple tool for G9a detection in native biological matrices.

Inspired by biological protein channels, the construction of artificial solid-state nanopores capable of smart responses to various targets has attracted widespread interest. Compared to biological nanopores, artificial solid-state nanopores offer distinct advantages, such as mechanical stability, controllable pore size



and geometry, and ease of functional modification.^{25–27} By immobilizing recognition probes on nanopore surfaces, target-induced changes in transmembrane ionic currents enable the detection of analytes. Leveraging the inherent size-exclusion effect of nanopores, such biomimetic systems can filter out large-sized biological interferences such as cellular debris and organelles, simplifying sample pretreatment and realizing real-time and convenient analysis of complex samples.^{28–31} In previous work, many groups have achieved sensitive detection of various proteins using solid-state nanochannels.^{32–34} Compared with the horizontal transport path, two-dimensional (2D) nanochannel membranes can carry mass transport in the vertical direction due to the different structures of 2D layered materials, which holds promise for high-throughput membrane-based applications.^{35–40} In addition, the unique layered structure of 2D nanomaterials provides abundant binding sites for probes, enhancing analyte capture efficiency, and its unique mass transfer path can prolong analyte residence time within the nanochannels, thereby improving sensing performance.^{41,42} Among them, the graphene oxide (GO)-based 2D nanochannel stands out for biosensing due to its high sensitivity, excellent biocompatibility, facile functionalization, *etc.* Its surface is rich in oxygen-containing groups (*e.g.*, hydroxyl and carboxyl), enabling covalent or non-covalent conjugation with biomolecules or functional moieties for target-specific recognition.^{43,44} Based on this, we propose that 2D nanochannel-based platforms may offer a groundbreaking solution to the challenges of G9a detection in complex cellular environments.

In this work, we developed a G9a detection system based on functionalized 2D solid-state nanochannels through the specific recognition between small molecule probes that anchored on the surface and dissociative peptide probes (Fig. 1). The small molecule probe 4'-aminobenzo-18-Crown-6 (briefly referred to as the Crown probe) was modified on the GO sheet, and the GO-Crown membrane was then obtained by vacuum filtration technology. The Crown probe exhibited a strong binding force with the unmethylated peptide through hydrogen bonding, thereby altering the ion transport characteristics within the nanochannels and enabling the differentiation of methylated and unmethylated peptides. Furthermore, the GO-Crown system was applied to the detection of histone methyltransferase G9a. The sensing performance in application scenarios involving different subtypes of breast cancer cells, the use of inhibitors, and simulated hypoxic environments was discussed as well. This research proposed that the GO-Crown membrane could be an effective detection system for G9a, which provides a powerful tool for cancer research and clinical application.

Results and discussion

Construction of the GO-Crown membrane

To construct the functionalized 2D nanochannel membrane, namely the GO-Crown membrane, GO was first dispersed in ultra-pure water and modified with 4'-aminobenzo-18-crown-6 (abbreviated to Crown), serving as the capture probe through an amidation reaction between the carboxyl groups on GO and the amine groups on Crown (Fig. 2a). As shown in Fig. 2b, because of the abundant negatively charged groups on the

surface of GO, the zeta potential of GO dispersion liquid was -27.87 mV. After the Crown probe was successfully modified, the amounts of carboxyl groups on the GO surface decreased and the zeta potential of GO-Crown increased to -4.25 mV. The particle size increased from 1400 nm to 2200 nm (Fig. S1). Fourier transform infrared spectroscopy (FTIR) was also used to monitor the functionalization process (Fig. 2c). Compared to the absorption of GO, the absorption of functionalized GO-Crown at 1718 cm^{-1} from carboxyl groups declined, while the absorption at 2920 cm^{-1} , ascribed to the asymmetric stretching of CH_2 from Crown, increased. In addition, the absorption at 1515 cm^{-1} from stretching vibration of the CO-NH bond increased, demonstrating the formation of the amide bonds and the successful covalent conjugation of the Crown probe.^{45,46}

Subsequently, the GO-Crown membrane was obtained by pumping the functional dispersion liquid into a film by vacuum filtration technology and peeling off after drying (Fig. 2d). A scanning electron microscope (SEM) was utilized to observe the microstructure of GO and GO-Crown membranes. As shown in Fig. 2e, both membranes exhibited distinct lamellar structures; however, after modification with the probes, the GO-Crown membrane was more incompact and its surface was rougher. Consistent with the SEM images, atomic force microscopy (AFM) also showed that the roughness of the GO-Crown membrane was greater than the GO membrane (Fig. S2). X-ray photoelectron spectroscopy (XPS) illustrated that, except for the two peaks of C 1s and O 1s at 287 eV and 533 eV, respectively, in the GO membrane, a new peak corresponding to N 1s at 399.5 eV emerged for the GO-Crown membrane (Fig. 2f, Tables S1 and S2).⁴⁷ Energy dispersive spectroscopy (EDS) also showed that the nitrogen content of the functionalized GO-Crown membrane was higher than that of the GO membrane, verifying the successful functionalization of the Crown probe again (Fig. S3). Moreover, the GO membrane and functionalized GO-Crown membrane were characterized by surface contact angle (CA) tests. As shown in Fig. 2g, after modification with the hydrophilic Crown probes, the CA decreased from 63.8° to 46.2° . From Raman spectra, the D peak was correlated with the vibration of carbon atoms of the sp^3 bond in GO, and the G peak was correlated with the vibration of carbon atoms of the sp^2 bond in GO. I_G/I_D (strength ratio) of GO was 1.01 and that of GO-Crown was 1.2, suggesting a reduction in structural defects⁴⁸ (Fig. S4). Then the layer distance of the 2D nanochannel was analyzed by X-ray diffraction (XRD) (Fig. 2h). The GO membrane and functionalized GO-Crown membrane exhibited diffraction peaks at 11.3° and 8.8° , respectively. According to the Bragg formula $d = \lambda/2 \sin \theta$, it could be calculated that the inter-layer distance of the GO membrane was 0.78 nm and that of the GO-Crown membrane was 1.01 nm, indicating that functionalization increased the layer spacing of the 2D nanochannel. Together, the above results proved that the GO-Crown membrane with 2D nanochannels was successfully constructed.

Evaluating the peptide responsiveness of 2D nanochannels

To verify the feasibility of the Crown probe for recognizing and screening unmethylated peptides and methylated peptides, a set of model peptides ($\text{NH}_2\text{-MARTKQTA}$, termed MA8) was



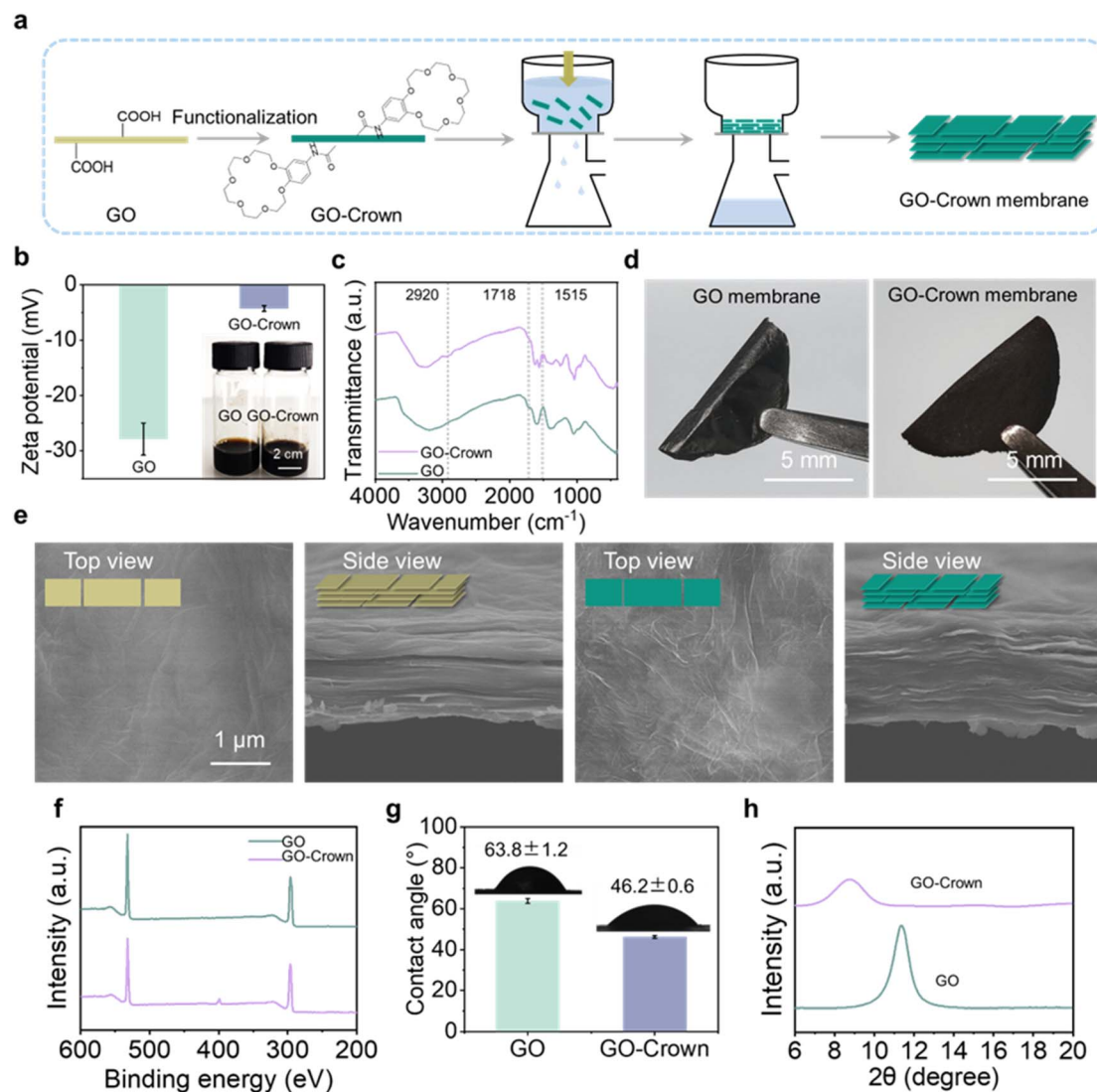


Fig. 2 Construction of the GO-Crown membrane. (a) Construction diagram of the functionalized GO-Crown membrane. (b) The zeta potential of GO and GO-Crown dispersion liquids. (c) FTIR spectra of GO and GO-Crown. (d) Photographs of GO and GO-Crown membranes. (e) SEM, (f) XPS, (g) CA and (h) XRD of GO and GO-Crown membranes.

used, in which K could undergo methylation modification at different levels (mono-, di-, and tri-methylated, termed MA8me1, MA8me2 and MA8me3, respectively) (Fig. 3a and S5–S9).⁴⁹ Then, the binding force between the Crown probe and the peptides was measured by isothermal titration calorimetry (ITC) tests. As shown in Fig. 3b, c and S10, the dissociation constant K_d of unmethylated peptides was 6.9 μM , while the K_d of methylated peptides was higher, demonstrating the higher binding force of the Crown probe toward unmethylated peptides and its strong potential to differentiate unmethylated and methylated peptides.

After verifying the specific recognition ability of the Crown probe toward unmethylated peptides, the response performance of the GO-Crown system was investigated (Fig. 3d). Under incubation conditions with a peptide concentration of 1 μM , the R value of the GO-Crown membrane for MA8 peptides was $54.7\% \pm 4.0\%$, and the R values for MA8me1, MA8me2 and

MA8me3 peptides were $4.2\% \pm 2.7\%$, $6.8\% \pm 3.9\%$, and $4.7\% \pm 1.1\%$, respectively. By means of incubation, the response of the GO-Crown membrane to MA8 peptides and methylated peptides showed significant differences. In contrast, for GO membranes, the responses of the R values were all less than 6% for MA8 peptides and methylated peptides (Fig. 3e–g and S11). To explore the response performance of the GO-Crown membrane toward MA8 peptides, the concentrations of MA8 peptides were set at 10 fM, 1 pM, 100 pM, 10 nM, and 1 μM . It could be seen from Fig. 3h and S12 that with the increase of peptide concentration, the R value gradually increased from $24.8\% \pm 2.7\%$ to $54.7\% \pm 4.0\%$, which proved that, in the incubation form, the GO-Crown membrane could achieve sensitive response performance to changes in peptide concentration.

Next, the response performance of the GO-Crown system under electric drive was investigated (Fig. 4a). To obtain the best



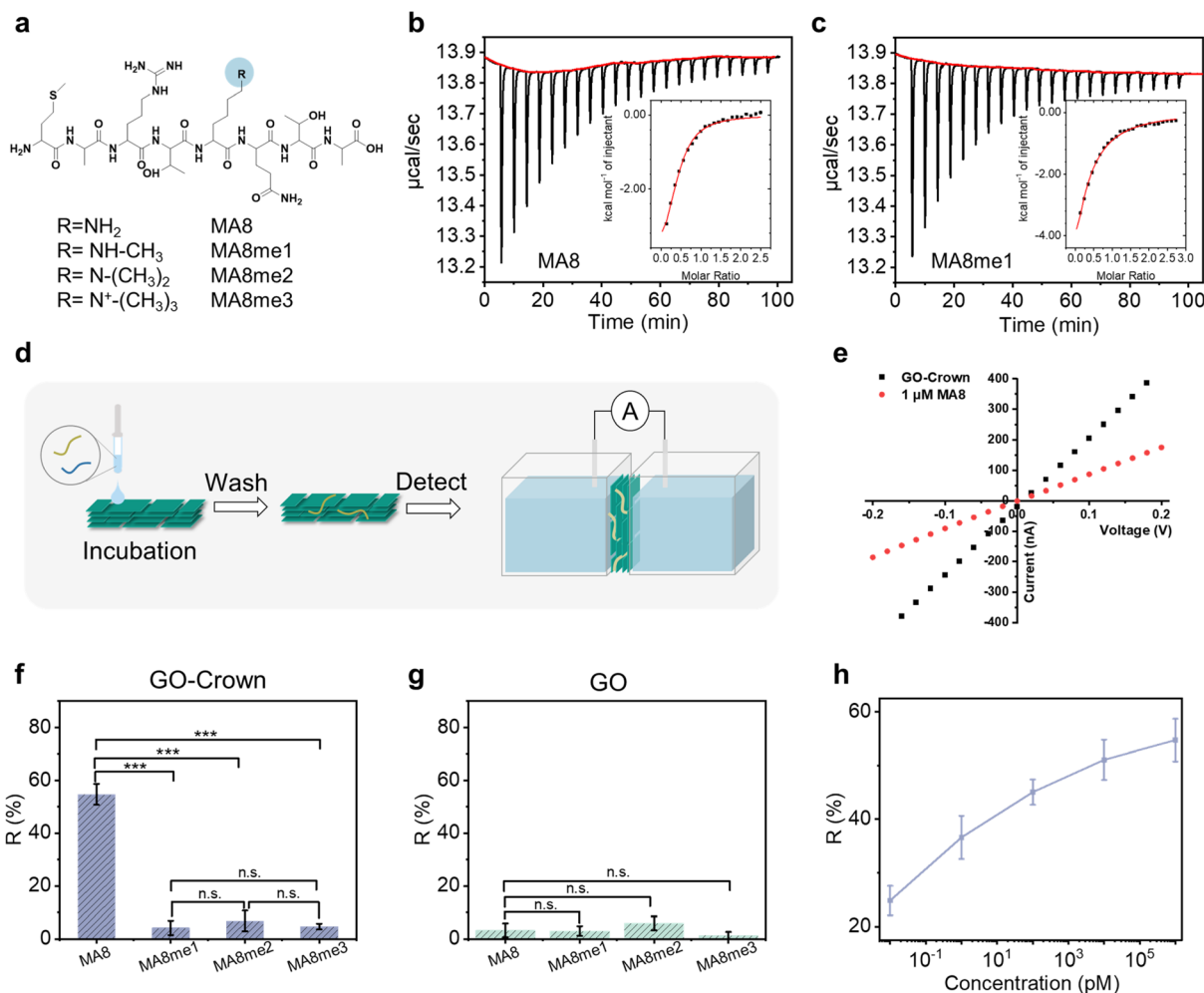


Fig. 3 Evaluating the peptide responsiveness of 2D nanochannels. (a) Peptide structures of MA8, MA8me1, MA8me2, MA8me3. ITC tests investigated the interaction of the Crown probe with (b) MA8 peptides and (c) MA8me1 peptides. (d) Schematic diagram of detection procedures of the GO-Crown membrane by incubation. (e) I - V curves obtained at the GO-Crown membrane before (black dashed line) and after (red dashed line) incubated with 1 μM MA8. The current change R value of (f) the GO-Crown membrane and (g) the GO membrane after incubated with 1 μM peptides with different methylation degrees. (h) The response of the GO-Crown membrane to different concentrations of MA8 peptides by incubation. The current change ratio (R) was calculated as $R = (I_0 - I)/I_0$, where I_0 and I are the current intensities at +0.2 V before (blank control) and after peptide treatments, respectively. Data are expressed as mean \pm SD ($n = 3$). * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, and n.s. not significant, as determined by a two-sided Student's t -test or one way ANOVA.

electric driving time, the ion current variation of the GO-Crown membrane was measured after different driving times ranging from 15 min to 10 h. As shown in Fig. 4a and S13, when unmethylated peptides passed through the membrane under electric drive from the driving chamber to the receiving chamber, they could be captured by the Crown probes on the membrane, thus inducing a decrease in current. When the electric drive time was 15 min, the R value (defined as $(I_0 - I)/I_0$, where I_0 and I are the current intensities at +0.2 V before and after driving, respectively) was about $18.0\% \pm 3.9\%$. Increasing the time to 1 h increased the R value to $44.6\% \pm 2.8\%$. With further increases in time, the R value gradually increased and then reached a plateau at about 50%. Therefore, 1 h was selected as the optimal driving time. The I - T result also showed that the current gradually decreased with the increase of time and the current value stabilized around -900 nA at about 1 h (Fig. S14).

Next, the screening ability of 2D nanochannels between methylated and unmethylated peptides under electric drive for 1 h was investigated. As shown in Fig. 4b and S15, the GO membrane without the capture probe was used to detect the peptides. As expected, there was no significant difference between methylated and unmethylated peptides ($p > 0.05$), and the current signals of both methylated and unmethylated peptides were less than 10%, demonstrating that GO itself could not recognize and distinguish the two kinds of peptides. Meanwhile, the GO-Crown membrane efficiently captured MA8, thus leading to a remarkable decrease in ion current and the R value was $44.6\% \pm 2.8\%$. In contrast, the R values for MA8me1, MA8me2, MA8me3 were $6.2\% \pm 1.3\%$, $6.0\% \pm 1.1\%$ and $3.1\% \pm 2.4\%$, respectively, which were much lower than that of MA8 ($p < 0.001$), indicating that the GO-Crown membrane could only capture the unmethylated peptides, instead of methylated



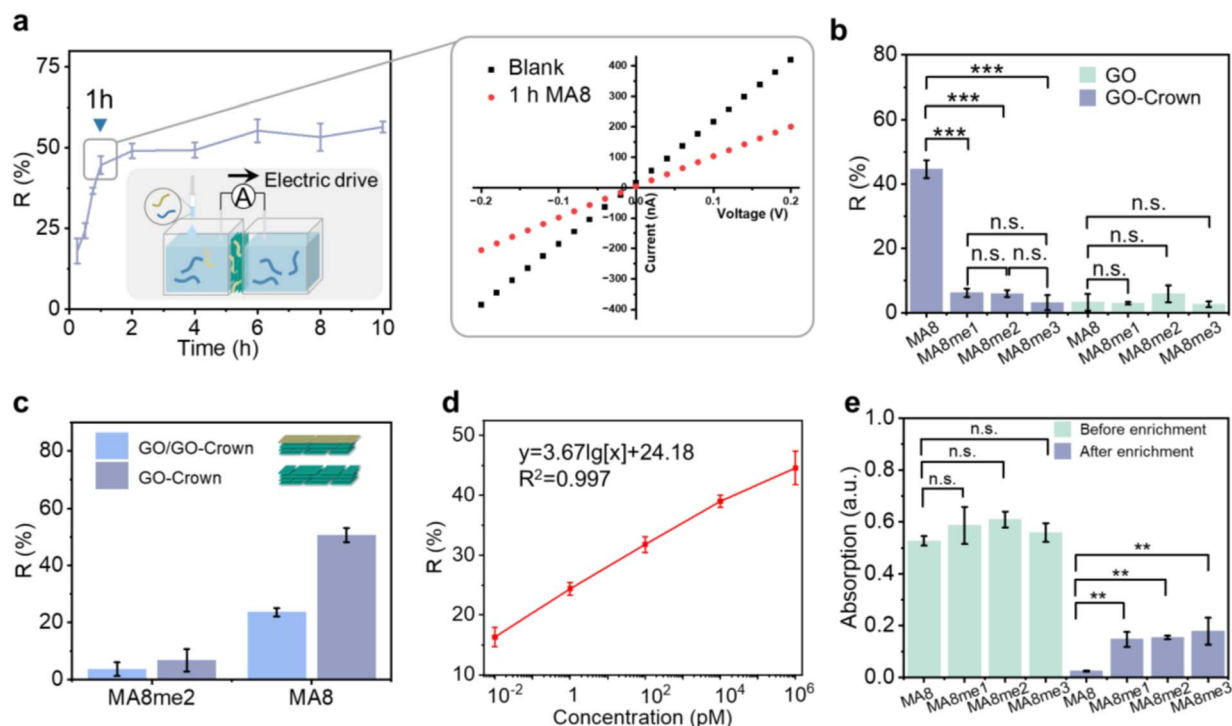


Fig. 4 Evaluating the peptide responsiveness of 2D nanochannels by electric drive. (a) The current change R value of the GO-Crown membrane after different electric driving times with MA8 peptides. The inset shows the schematic diagram of detection procedures of the GO-Crown membrane by electric drive. (b) The current change R value of GO-Crown or GO membranes after electric driving the peptides with different methylation degrees. (c) The current change R value of GO/GO-Crown/GO, GO/GO-Crown or GO-Crown membranes after electric driving the peptides with different methylation degrees. (d) The response of the GO-Crown membrane to different concentrations (10 fM, 1 pM, 100 pM, 10 nM, and 1 μ M) of MA8 peptides. (e) The UV absorption value of the unmodified peptide and methylated peptides before and after the enrichment. The current change ratio (R) was calculated as $R = (I_0 - I)/I_0$, where I_0 and I are the current intensities at +0.2 V before (blank control) and after peptide addition, respectively. Data are expressed as mean \pm SD ($n = 3$). * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, and n.s. not significant, as determined by a two-sided Student's t -test or one way ANOVA.

peptides including mono-, di- and tri-methylated, and cause the ion current change.

To further confirm that the variation in current was induced by the binding between the Crown probe and unmethylated peptides, composite membrane with different contents of Crown, namely "GO/GO-Crown," were constructed by layer-by-layer pumping. The CA variation proved their successful construction (Fig. S16). Then, the current signal response performance of the composite membrane was tested. As shown in Fig. 4c and S17, GO/GO-Crown could not bind MA8me2 peptides and did not induce current changes. Compared to the pure GO-Crown membrane, the R value of GO/GO-Crown for MA8 peptides was about 24%, demonstrating that as the proportion of the Crown probe decreased, the capture capability of the membrane for unmethylated peptides was distinctly weakened.

Subsequently, the MA8 peptide concentration–response curve of the GO-Crown membrane system was examined, as shown in Fig. 4d and S18. When the MA8 peptide was only 10 fM, the R value was $16.3\% \pm 1.6\%$. As the peptide concentration increased to 1 μ M, R value gradually increased and reached $\sim 45\%$, indicating that, in the form of electric drive, the GO-Crown membrane was sensitive to unmethylated peptides, and the current decrease ratio exhibited a positive relationship to the concentration of

unmethylated peptides. The plot showed good linearity at concentrations of 10 fM–1 μ M, and the linear regression equation is $R = 3.67 \lg X$ [with the concentration in pM of MA8] + 24.18, with a correlation coefficient of 0.997.

To explore the saturation concentration of the GO-Crown membrane, the concentration of MA8 peptides was further enhanced. As shown in Fig. S19, when the concentration was increased from 1 μ M to 25 μ M, the R value did not increase obviously; the R values of 5 μ M and 25 μ M MA8 were about 48%, respectively. After statistical analysis, there was no significant difference in response between 1 μ M, 5 μ M and 25 μ M MA8 peptide ($p > 0.05$), confirming that the saturation concentration of the peptide was 1 μ M. In addition, to realistically simulate the working conditions of peptide fragments from histone degradation, a mixture of methylated and unmethylated peptides, exemplified by dimethylated peptides, was detected by the GO-Crown membrane. The R value of the mixed peptide (1 μ M MA8 + 0.1 μ M MA8me2) was $49.5\% \pm 2.0\%$, which was nearly the same as that of 1 μ M MA8 without the addition of MA8me2, and it was found that no significant difference existed. The situation of the mixed peptide with a high concentration (25 μ M MA8 + 10 μ M MA8me2) was similar, illustrating that the existence of methylated peptides in the mixed solution would not obviously affect the recognition of unmethylated peptides.



In histone lysates, methylated peptides generally occupy a small proportion, leading to difficulty in their detection; in this case, the enrichment of methylated peptides plays a crucial role.^{50,51} Thanks to the screening ability of methylated and unmethylated peptides, the GO-Crown membrane could also be utilized for enrichment of methylated peptides. When mixed peptides pass through the membrane under electric drive, unmethylated peptides bind to the Crown probe on the membrane, while the methylated peptides do not bind to the probe and are driven to the receiving chamber, achieving the enrichment of the methylated peptides in the mixed sample. UV-vis spectrophotometry and mass spectrometry were used to verify this process. As shown in Fig. 4f and S20, four peptides passed through the GO-Crown membrane under electric drive and their concentration variation was monitored by UV-vis absorbance, respectively. Compared to the approximately same absorbance ($p > 0.05$) before driving, the absorption value of methylated peptides was in the range of 0.1–0.2, which is about 5–10 times of unmethylated peptides (about 0.01) and showed a significant difference ($p < 0.01$). After enrichment, the absorption value of the unmethylated peptide was too low to obtain the converted concentration, while the concentrations of MA8me1, MA8me2 and MA8me3 were 0.0915 μM , 0.103 μM and 0.145 μM , respectively, suggesting the enrichment ability of the GO-Crown membrane for methylated peptides. Then, the solution in the receiving chamber was collected after driving a mixed solution (1 μM MA8 + 0.1 μM MA8me2) and characterized by mass spectrometry. Only the peaks from MA8me2 (m/z value of $[\text{M} + \text{H}]^+$ 934.51 and $[\text{M} + 2\text{H}]^{2+}$ 467.76) could be found, rather than MA8 (m/z value of $[\text{M} + \text{H}]^+$ 906.48 and $[\text{M} + 2\text{H}]^{2+}$ 453.75) (Fig. S21). These results further proved that the GO-Crown membrane system could be used for the separation of unmethylated and methylated peptides and had the enrichment ability for methylated peptides.

In the GO-Crown membrane system, the electric driving form played a crucial role in the process of peptides passing through the nanochannel membrane. In this complex process, both the electric field force and the concentration difference had significant effects on the transmembrane behavior of peptides. The electric field force exerted a specific driving force when the peptide molecules passed through the nanochannel membrane by its directional effect on the charged particles. The difference in concentration was generated due to the difference in the concentration of peptides between the two sides of the electrolytic cell. The imbalance of chemical potential promoted the diffusion of peptides from the region of high concentration to the region of low concentration, and then pushed the peptides through the nanochannel membrane. Three models were designed to explore the influence of F_C (concentration difference) and F_E (electric field force) in the electric drive model (Fig. S22 and S23). First, to investigate the influence of F_E , the concentration was the same on both sides of the cell ($F_C = 0$). The results showed that there was no significant difference between the two solutions, with the UV absorption values was around 0.7, so it was speculated that the influence of F_E was tiny. Second, to investigate the influence of F_C , the electric field force was set to 0 ($F_E = 0$). The result showed that the UV

absorption value of the solution on the other side was about 0.06, which proved that there were peptides driven to the other side. Third, there were two cases in which F_E and F_C were in the same direction or in opposite directions.

The results showed that when F_E and F_C were in the same direction, the absorption value of the solution collected on the other side was about 0.13, while when F_E and F_C were in opposite directions, the absorption value was about 0.08. In summary, it could be concluded that both F_E and F_C had a certain effect on the driving force of peptides under electric drive, and it was mainly driven by F_C , with the driving effect being optimal when F_E and F_C were combined in the same direction.

Evaluating G9a responsiveness of the system

Encouraged by the satisfactory separating ability of the GO-Crown membrane toward methylated and unmethylated peptides, its application in detecting G9a was considered, since G9a could catalyze the methylation of specific positions in histone. According to literature reports,^{52–54} G9a interacts specifically with 6–11 amino acid residues of H3, and the smallest *in vitro* substrate recognition unit is heptapeptide with the TARKSTG (TG7) sequence. In this case, the TG7 peptide was used here as a detecting probe, combined with the GO-Crown membrane to construct a G9a detection system. In the presence of G9a, the substrate TG7 peptide could be methylated to mono-, di-, and tri-methylated peptides (termed TG7me1, TG7me2 and TG7me3) (Fig. S24). Then, the reaction liquid was added to the electrolytic cell for electric drive, and the current response of the GO-Crown membrane was recorded, which was related to the residual TG7 peptide content and also G9a content.

First, to confirm the methylation ability and process of G9a toward the TG7 peptide, the products of the enzymatic reaction after different reaction times were characterized by mass spectrometry (Fig. 5a and S25–S27). The theoretical m/z value $[\text{M} + \text{H}]^+$ of TG7 and TG7me1 was calculated to be 720.40 and 735.04, respectively and the peaks at 720.40 and 735.04 could be found when the reaction time was 1 h, proving the co-existence of TG7 and TG7me1 and the occurrence of methylation at this time. When the reaction time reached 2 h and 3 h, except for the peak of TG7 and TG7me1, the peak at 748.43, which was from TG7me2 (theoretical m/z value $[\text{M} + \text{H}]^+$ of 748.43) could also be found, and the ratio of peak intensity for TG7me2/TG7me1 was increased from 2 h to 3 h. At the 4-h time point, the TG7me3 peptide emerged. The above results demonstrated that the methylation process of G9a toward the TG7 peptide occurred as expected, and with extended reaction time, the methylation degree of TG7 could be gradually strengthened.

Then, the feasibility of this combined system to detect G9a was investigated. The reaction solution of G9a (20 nM) and TG7 peptide (10 μM) after different reaction times was added to the electrolytic cell for testing. As shown in Fig. 5b, S28 and S29, when the reaction time was 0 h, the current signal response R of the GO-Crown membrane was $42.6\% \pm 2.0\%$, confirming that unmethylated TG7 could be captured by GO-Crown and lead to the ion current decrease, which was the same as that of the MA8 peptide. Meanwhile, with the increase of reaction time to 7 h,



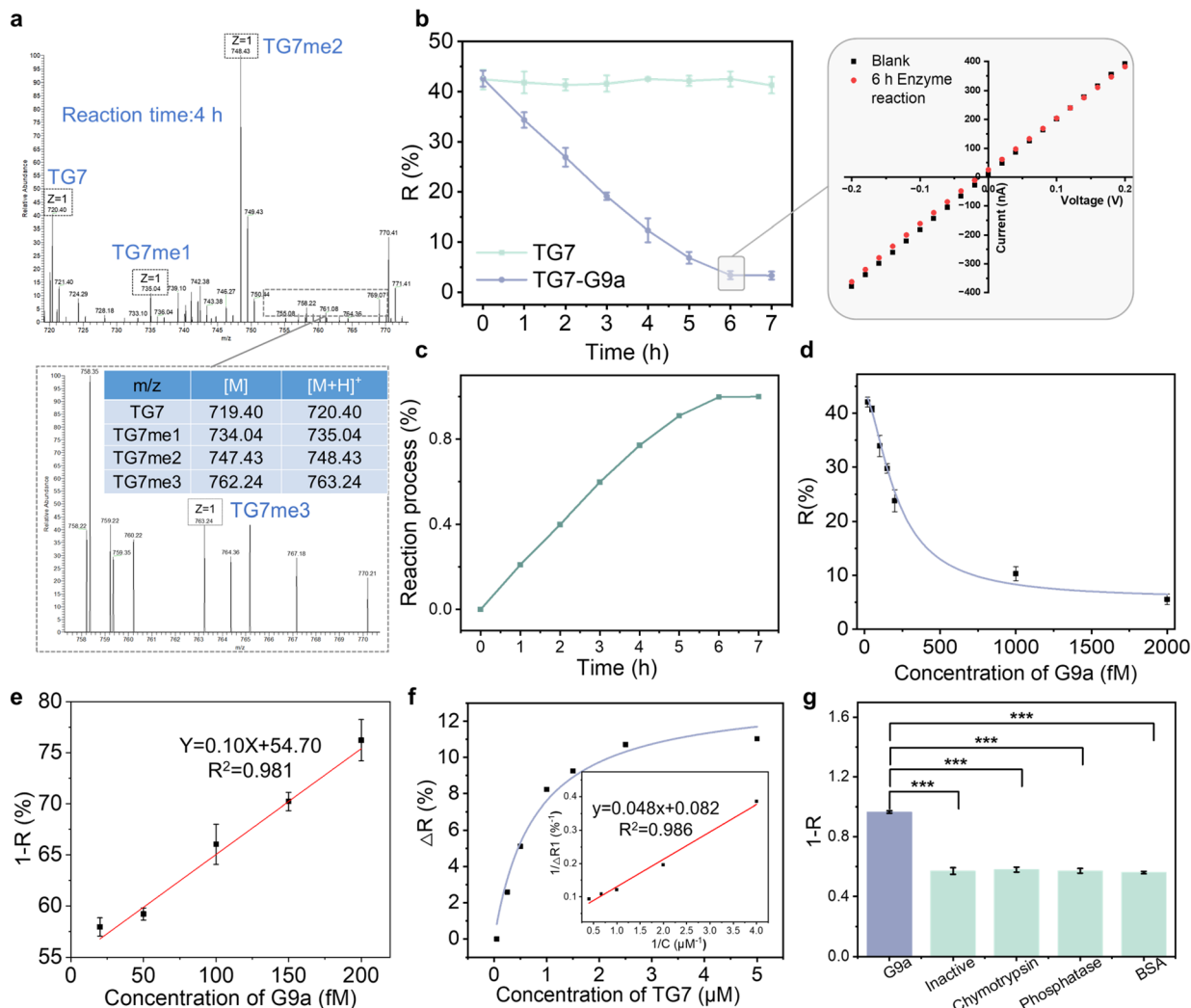


Fig. 5 Evaluating G9a responsiveness of the system. (a) The mass spectrum of the products of enzymatic reaction after 4 h reaction times. (b) The current signal response of the GO-Crown membrane system with G9a or without G9a at different reaction times. (c) The corresponding digestion curve of the GO-Crown membrane system. (d) The current response of the GO-Crown membrane system toward G9a in the concentration range of 20 fM–2 pM. (e) The fitting curve of the G9a concentration range of 20–200 fM. (f) Under certain conditions, the enzymatic kinetics of G9a was studied based on current signals. (g) The GO-Crown membrane system in different media with G9a or other enzyme interferences to study specificity. The current change ratio (R) was calculated as $R = (I_0 - I)/I_0$, where I_0 is the blank control current at +0.2 V and I is the G9a current response at +0.2 V. Data are expressed as mean \pm SD ($n = 3$). * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, and n.s. not significant, as determined by a two-sided Student's t -test.

the TG7 peptide could be methylated to different degrees and thus could not bind to GO-Crown to induce current variation. The R value decreased to 34.13% at 1 h, continued to decline during 6 h and reached a plateau (3.45%) at 6 h, proving the above process and the potential of the system for detecting G9a. For comparison, the TG7 peptide solution without G9a was also added to the electrolytic cell after different times under the same conditions. With the increase of reaction time, the R value of the system for the enzyme-free mixture remained at about 42%, and there was no obvious change in current response, which proved that the current signal of the GO-Crown membrane system was caused by enzymatic action of G9a. Since after 6 h, the process of enzyme catalyzed reaction was maximal and approximately 100%, the following detection experiments selected 6 h as the reaction time (Fig. 5c).

The response performance of the GO-Crown membrane system to G9a concentration in the range of 20 fM–20 nM was investigated (Fig. 5d and S30). When the G9a concentration was in the range of 20 fM–2 pM, the GO-Crown membrane system produced an obvious current signal response with the change of G9a concentration, and the R value gradually changed from $42.1\% \pm 0.9\%$ to $5.6\% \pm 1.0\%$. When the G9a concentration was in the low concentration range of 20 fM–200 fM, the fitted curve had good linearity and the linear regression equation is $Y = 0.10 \times [\text{concentration of G9a}] + 54.70$, with a correlation coefficient of 0.981 (Fig. 5e). The limit of detection ($3\sigma/\text{slope}$, $\sigma = 0.688$) is 21 fM, which was much lower than that of reported methods (Table S3). These results proved the G9a detection ability of the system and also laid a foundation for subsequent detection in complex environments. Moreover, when increasing the G9a



concentration from 2 pM to 20 nM, the current signal was basically unchanged and the R value was about 4–5%, suggesting that the response of the GO-Crown membrane system had reached a saturation state.

By virtue of well G9a responsiveness ability of the system, the enzymatic reaction kinetics of G9a was further discussed (Fig. 5f and S31). To explore the effect of substrate concentration on enzymatic reaction kinetics, the temperature (25 °C), enzyme concentration (20 nM) and reaction time (6 h) were controlled. The concentrations of substrate TG7 peptides were set at 0.05 μ M, 0.25 μ M, 0.5 μ M, 1.5 μ M, 2.5 μ M, and 5 μ M successively. The difference of R at different concentrations was taken as the vertical coordinate. The value of ΔR gradually increased with the increase of concentration. By linear fitting of the obtained current signal data, the K_m value of the enzyme activity fitted under the experimental conditions was 1.71 μ M.

Specificity is an important factor in evaluating the response performance of detection systems. As shown in Fig. 5g and S32, the $1 - R$ of the GO-Crown membrane system towards G9a was 0.97. However, the current response of the system to other enzymes, including α -chymotrypsin, phosphatase and bovine serum albumin (BSA), was lower than 0.6. The significant difference demonstrated reliable specificity for G9a. In addition, G9a was also deactivated by heating, and the $1 - R$ value after the reaction of the GO-Crown membrane system with deactivated G9a was 0.57, proving that the activity of G9a was the key factor inducing the change in ionic current.

Detection of G9a from cells in different application scenarios

With good selectivity and sensitivity, the performance of the GO-Crown membrane system for detecting G9a from MCF-7 cells (breast cancer cell line with high G9a expression) was investigated (Fig. 6a, b, S33 and S34). Along with the MCF-7 cell concentration rising from 50 cells per mL to 5×10^4 cells per mL, the current change ratio R value decreased gradually, proving that there was a positive relationship between concentrations of G9a and the current change ratio. The variation tendency was similar to the G9a concentration–response curves in Fig. 5f. However, when incubated with G9a from the MCF-10A cells (normal mammary epithelial cells with low G9a expression), TG7 peptides cannot be methylated, thus resulting in an unapparent change in ion current, even when the cell concentration was up to 5×10^4 cells per mL.

After successfully applying the system to precisely detect G9a from cells, we wonder if this system could be used in diverse application scenarios, including differentiation of different subtypes of breast cancer cells, evaluation of inhibitor action, and comparison of different oxygen contents, since the secretion of G9a was highly related to these factors. First, three subtypes of breast cancer cells: MCF-7 cells, MDA-231 cells and MDA-468 cells were selected. At the same time, MCF-10A was treated as the control group. The four cell lines were obtained from Wuhan Pricella Biotechnology Co., Ltd. Cell lysates from the four cell lines at a concentration of 5×10^3 cells per mL were incubated with TG7 peptides, respectively, and then added into the GO-Crown membrane system under electrical drive. As

shown in Fig. 6c and S35, the GO-Crown membrane system had obvious differences in electrical signal responses to cancer cells and normal cells. The $1 - R$ value was $58.7\% \pm 0.9\%$ for MCF-10A cells, $79.4\% \pm 1.7\%$ for MCF-7 cells, $74.1\% \pm 0.5\%$ for MDA-231 cells, and $68.2\% \pm 1.0\%$ for MDA-468 cells, and there was a significant difference ($p < 0.001$) between each two groups, indicating that the system could realize the distinction between cancer cells and normal cells relying on the different expression of G9a. For different subtypes of breast cancer cells, the G9a detected by the system from high to low was in the order of MCF-7 cells, MDA-231 cells and MDA-468 cells, and the electrical signal response was significantly different, proving that the system could realize the differentiation of different subtypes of breast cancer cells. To confirm the accuracy of tendency, the enzyme linked immunosorbent assay (ELISA) analysis, as a gold standard, was further utilized to measure the G9a content in cell lysate of the four kinds of cells. The G9a concentration in the four cell lines was significantly different and was calculated to be 0.234 ng mL^{-1} , 0.906 ng mL^{-1} , 0.759 ng mL^{-1} and 0.497 ng mL^{-1} in MCF-10A cells, MCF-7 cells, MDA-231 cells and MDA-468 cells, respectively, which was consistent with the detection results of the functionalized GO-Crown membrane, proving the reliable detection ability of the system for differentiating different subtypes of breast cancer cells.

In another application scenario, an inhibitor of G9a UNC068 with low toxicity was used (Fig. S36), which could interfere with G9a's normal interaction with substrates and cofactors (such as SAM) by binding to the active site of G9a or key regions near it, thereby impeding its ability to catalyze methyl-transfer reactions. The concentration of the inhibitor was set at 5 nM, 150 nM, and 1.5 μ M, and the expression level of G9a in the four cell lysates was detected after co-culture for 72 h (Fig. 6d and S37). With the increase of inhibitor concentration, for MCF-10A cells, the R value did not change significantly and fluctuated within the range of $\sim 43\%$. For MCF-7 cells, the R value changed from $\sim 21\%$ to $\sim 32\%$. For MDA-231 cells, the R value changed from $\sim 26\%$ to $\sim 36\%$. For MDA-468 cells, the R value changed from $\sim 32\%$ to $\sim 40\%$. For convenience of comparison, " $1 - R$ " was selected as the vertical coordinate. It was demonstrated that the GO-Crown membrane system had a reliable ability to detect G9a in an environment where the expression of G9a was down-regulated after inhibitor treatment. ELISA kit was used to detect cell lysate treated with 5 nM, 150 nM, and 1.5 μ M inhibitor, which was consistent with the results determined by the GO-Crown membrane, proving the reliable detection capability of the system.

It has been reported that G9a could be up-regulated in hypoxic environments. In brief, under hypoxia conditions, oxygen receptors in cells would sense the reduction of oxygen content, thereby stabilizing and activating hypoxia-inducible factor (HIF-1 α). Activated HIF-1 α could bind to the hypoxia response elements of the G9a gene promoter, recruiting transcription-related complexes, promoting the binding of RNA polymerase to the G9a gene promoter, thus enhancing the transcription of G9a gene and making G9a expression up-regulated.^{55–57} In this case, different levels of oxygen content were constructed for cells, including 21% normoxia, 10% hypoxia and 1% hypoxia. The GO-Crown membrane system had



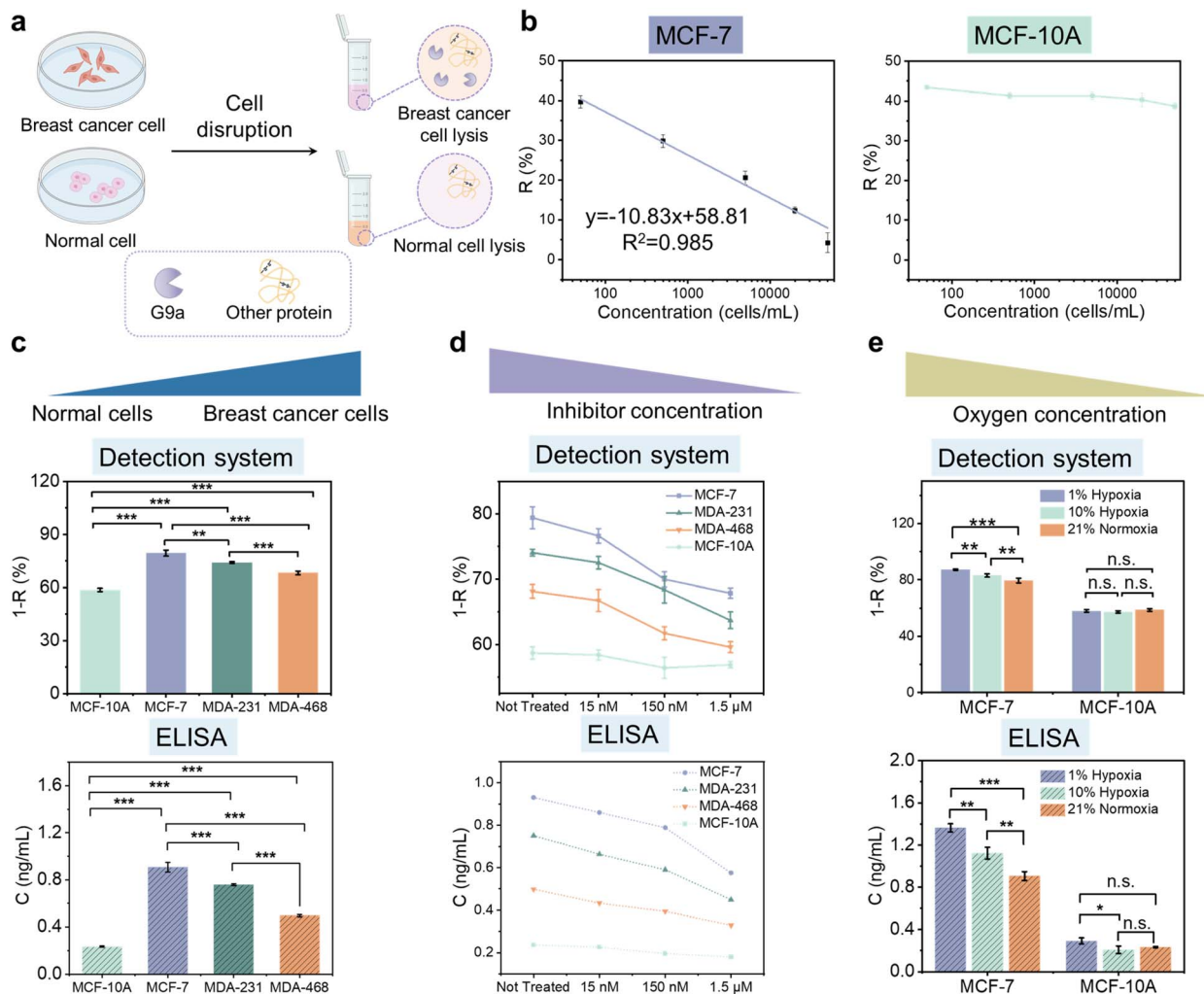


Fig. 6 Detection of G9a from cells in different application scenarios. (a) Acquisition of G9a from complex biological samples. (b) The current signal of MCF-7 cell lysate and MCF-10A cell lysate with the cell number of $50\text{--}5 \times 10^4$ cells per mL. (c) The GO-Crown membrane system current response to MCF-10A, MCF-7, MDA-231, and MDA-468 cell lysates, and the G9a concentration values of MCF-10A, MCF-7, MDA-231, and MDA-468 cell lysates by ELISA kit. (d) The current signal of cell lysate after treatment with different concentrations of inhibitors, and the concentration values of MCF-10A, MCF-7, MDA-231, and MDA-468 cell lysates after inhibitor treatment by ELISA kit. (e) The current signal of cell lysate at different oxygen concentrations, and the concentration values of MCF-10A, MCF-7, MDA-231, and MDA-468 cell lysates after hypoxic culture treatment by ELISA kit. The concentration of TG7 peptides was $10 \mu\text{M}$. The current change ratio (R) was calculated as $R = (I_0 - I)/I_0$, where I_0 is the blank control current at $+0.2 \text{ V}$ and I is the G9a current responses at $+0.2 \text{ V}$. Data are expressed as mean \pm SD ($n = 3$). * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, and n.s. not significant, as determined by a two-sided Student's t -test or one way ANOVA.

a significant response to the expression of G9a in MCF-7 cells at different oxygen contents, with $\sim 13\%$ at 1% oxygen concentration and $\sim 17\%$ at 10% oxygen concentration (Fig. 6e and S38). For MCF-10A cells, the values were $\sim 42\%$ at both 1% oxygen concentration and 10% oxygen concentration, indicating that a hypoxic environment had no significant effect on the expression of G9a in MCF-10A cells. The cell lysates cultured in hypoxic environments were detected by ELISA kit, which was consistent with the results determined by the GO-Crown system. Therefore, the system could be applied in hypoxic environments where G9a expression was upregulated with high sensitivity. The GO-Crown membrane system had the certain ability to distinguish between 1% hypoxia and 10% hypoxia conditions, so it was considered that the system had a potential application as an oxygen indicator.

Conclusions

To construct a label-free, highly selective, sensitive, and simple tool for G9a detection in native biological matrices, a G9a detection system based on functionalized 2D solid-state nanochannel membranes was developed in this study through the specific recognition between small molecule probes that anchored on the surface and dissociative peptide probes. The confined environment in nanochannels enhances interaction times and efficiency between the peptide and the crown ether receptors on the channel walls. This multivalent effect amplifies the subtle chemical difference between methylated and unmethylated peptides into a significant and measurable electrical signal. The excellent sensing performance enables diverse detection scenarios, including differentiation of different



subtypes of breast cancer cells, screening of enzyme inhibitors and sensing of hypoxic cellular environments. This research proposes that the GO-Crown membrane could be an effective detection system for G9a, which provides a powerful tool for cancer research and clinical application.

Author contributions

F. X., X. L. and J. J. H. conceived and supervised the project. J. J. H. and N. L. wrote the manuscript. W. J. and N. L. conducted the experiments. R. L., J. J. H., W. J. and N. L. contributed to data analysis. All authors contributed to manuscript revision.

Conflicts of interest

There are no conflicts to declare.

Data availability

Supplementary information: experimental procedures, characterization data, and *I-V* curves. See DOI: <https://doi.org/10.1039/d5sc09089j>.

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