A flexible IrO₂ membrane for pH sensing

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An optimized mixture of polydopamine (PDA) and polyvinyl alcohol (PVA) is employed as the surface functionalizing agent and reducing agent to encapsulate individual polypropylene (PP) fibers of polypropylene micromembrane (PPMM). The functionalized PPMM becomes hydrophilic to allow the formation of Au nuclei for subsequent electroless Au deposition. The metalized PPMM is further deposited with IrO₂ nanoparticles, and evaluated as a flexible and porous pH sensor. Images from scanning electron microscope confirms the uniform formation of IrO₂ nanoparticles on Au-coated PP fibers. For pH-sensing performance, the IrO₂-decorated metalized PPMM reveals a super-Nernstian response for a sensing slope of -74.45 mV/pH in aqueous solutions with pH value ranging between 2 and 12. In addition, the pH-sensing performance is properly maintained after 5000 bending cycles and hysteresis is modest in an acidic environment. The cell viability test indicates a negligible bio-toxicity. Our strategy of using a conductive polymeric membrane decorated with IrO₂ nanoparticles enables possible sensing applications in wearable and implantable electronics.

The pH value is a critical indicator for many biological systems¹⁻³. It is known that for patients suffering from cancer, renal failure, multiple sclerosis, and psychiatric illness, severe disorders in acid/base equilibrium might occur⁴⁻⁶. For example, according to Mani et al., cancer cells preferentially convert glucose and other substances into lactic acid, and hence the intracellular fluid becomes slightly acidic⁶⁻⁸. Therefore, an effective pH-sensing technique for tissues is necessary to prevent further physiological damages. So far, a variety of metal oxides including WO₃, RuO₂, and IrO₂ have been investigated for pH-sensing purpose⁹⁻¹¹. Among them, the IrO₂ demonstrates a wide working range, an impressive chemical inertness, and a negligible susceptibility to redox interference.

The IrO₂ is a conductive oxide known for excellent biocompatibility¹². In addition to pH-sensing, the IrO₂ has been explored as a pseudocapacitor for energy storage, as a neuron-stimulating electrode for implantable electronics, and as an oxygen-evolving electrocatalyst in water electrolysis¹³⁻¹⁵. However, the natural reserve for IrO₂ is limited so its expensive cost prohibits its widespread use in commercial products. Therefore, it is necessary to develop a suitable processing technique to produce IrO₂ nanoparticles or thin films in a more effective way. Previously, we have deposited IrO₂ nanoparticles on an ITO substrate, and reported their impressive super-Nernstian pH-sensing behaviors¹⁶. In addition, we have prepared IrO₂ thin films on ITO and FTO substrates for bio-stimulating studies¹⁷,¹⁸. However, those substrates are rigid and thus are not applicable in wearable and implantable electronics, for which a flexible platform is necessary.

For pH-sensing in both in-vivo and in-vitro cases, a desirable flexible substrate also requires sufficient porosity to allow intimate contact of analyte. We recognize that a polymer membrane exhibited ideal attributes since its chemical properties (nature of polymer and hydrophilicity) and physical properties (porosity, pore size, permeability, thickness, mechanical strength, etc.) could be tailored-made to meet the specifications of end applications. In addition, many polymer membranes are mature industrial products with reasonable costs. Unfortunately, conventional polymer membranes are electrically insulating, and a metallization pretreatment is required. To date, the metallization of polymer membranes has been conducted through chemical vapor deposition, physical vapor deposition, and electroless deposition¹⁹⁻²¹. It is noted that both physical and chemical vapor depositions entail vacuum chambers with sophisticated equipment, and the conformal deposition of metals within the polymer membrane is rather difficult. In contrast, the electroless route is relatively simple in terms of processing tools, and is more likely to achieve homogeneous coating throughout the entire polymer structure.

In this work, we demonstrated the fabrication of a flexible pH sensor using polypropylene micromembrane (PPMM) as a porous substrate. Our process started with a mixture of polydopamine (PDA) and polyvinyl alcohol...
(PVA) as the hydrophilic agent to functionalize individual polypropylene (PP) fibers of PPMM. The PDA also served as the reducing agent to prepare nucleation sites for subsequent electroless Au deposition, for which a conformal formation of Au overcoat encapsulating every PP fiber was achieved. The metalized PPMM was further electrodeposited with IrO₂ nanoparticles, and then evaluated as a flexible pH sensor.

Experimental section

Chemicals and reagents. The PPMM (Rone Scientific; catalog number: 201PP-47-045-50; pore size of 450 nm) was in a circular shape with diameter of 47 nm and thickness of 200 µm. Tris(hydroxymethyl)aminomethane hydrochloride (Tris–HCl), sodium l-ascorbate, acetic acid (CH₃COOH), and boric acid (H₃BO₃) were purchased from Sigma-Aldrich. Dopamine hydrochloride (DA-HCl; the monomer for PDA) was purchased from Acros Organic. PVA (MW: 70 kDa) was purchased from Merck. Potassium hydroxide (KOH), sodium hydroxide (NaOH), sodium thiosulfate (Na₂S₂O₃), sodium sulfite (Na₂SO₃), and hydrochloric acid (HCl) were purchased from SHOWA. Anhydrous citric acid (C₆H₈O₇) was purchased from J.T. Baker. Phosphoric acid (H₃PO₄) was purchased from Honeywell. Hydrogen tetrachloroiridate (IV) trihydrate (HAuCl₄·3H₂O) and potassium hexachloroiridate (IV) (K₂IrCl₆) were purchased from Alfa Aesar. All these chemicals were obtained in analytical grade and were used without further purification.

Functionalization of PPMM. The functionalization of PPMM was carried out by a conformal deposition of PDA and PVA on individual PP fibers of PPMM. The PPMM was cleaned by ethanol (99.9 vol%) before its use. First, a buffer solution for Tris–HCl was prepared by mixing deionized water and anhydrous ethanol in 7:3 volume ratio. For the codeposition of PDA and PVA, 40 mg DA-HCl (effective DA amount was 32 mg) was dissolved in 20 mL 10 mM Tris–HCl buffer solution, followed by adding different amounts of PVA (64, 128, and 256 mg). At this stage, the pH of the mixture was 6. Subsequently, minute amount of aqueous 1 M KOH solution was added to adjust the pH to 8.5. Next, the PPMM was submerged in the mixture for 20 h at 25 °C. The immersion step was under constant stirring in ambient atmosphere so the dissolved oxygen was able to initiate the oxidation (polymerization) of DA to PDA. Lastly, the sample was retrieved and washed by deionized water. The PPMM after functionalization of PDA and PVA was denoted as the PPMM@PDA/PVA(1/2), PPMM@PDA/PVA(1/4), and PPMM@PDA/PVA(1/8), respectively. The ratio of 1/2, 1/4, and 1/8 represented the weight ratio of PDA and PVA on individual PP fibers of PPMM. For control experiments, we also prepared PDA-coated and PVA-coated PPMM by conducting the synthetic process in identical conditions (using 40 mg DA-HCl or 256 mg PV A), and the resulting samples were denoted as the PPMM@PDA and PPMM@PVA, respectively.

Deposition of Au and IrO₂ on functionalized PPMM. The functionalized PPMM (PPMM@PDA/PVA(1/8)) underwent a seeding step by submerging the sample in 0.1 M HAuCl₄ aqueous solution (pH 5.2) at 25 °C for 10 min. In this step, the catechol group of PDA was able to reduce the adsorbed Au⁺ ions to form Au nuclei for following electroless Au deposition. Next, the sample was immersed in an electroless Au deposition bath containing 8 mL 1 M citrate acid aqueous solution and 9.8 mL deionized water (the pH for the mixture was adjusted to 6.5 by adding 1 M KOH aqueous solution), as well as 0.4 mL 1 M Na₂SO₃ aqueous solution (serving as the reducing agent). Next, 1 mL 0.1 M H₃AuCl₄ aqueous solution was added under constant stirring, along with the addition of 0.8 mL 1 M Na₂SO₃ aqueous solution (serving as the complexing agent). At this stage, the Na₂SO₃ reduced the Au³⁺ ions for the formation of Au⁺ ions that were complexed by Na₂S₂O₃. Afterward, the reduction of Au⁺ ions was completed by the addition of 0.792 mg sodium l-ascorbate, and the bath temperature was raised to 30 °C. The formation of conformal Au deposit on individual PP fibers lasted for 2 h to fabricate the metalized PPMM, which was denoted as the PPMM@PDA/PVA(1/8)@Au.

The electrodeposition of IrO₂ nanoparticles on the metalized PPMM started with the preparation of IrO₂ colloidal suspension according to what was reported by Zhao et al. The deposition of IrO₂ nanoparticles was performed in a three-electrode cell using a Pt foil (2 × 2 cm²) and a saturated calomel electrode (SCE) as the counter and reference electrodes, respectively. The metalized PPMM (1 × 1 cm²), serving as the working electrode, was subjected to a potentiostatic mode at 0.4 V (vs. SCE) for 30 min. The resulting sample was denoted as the PPMM@PDA/PVA(1/8)@Au@IrO₂.

Materials characterization and pH-sensing analysis. The morphologies and microstructures for samples including PPMM, PPMM@PDA, and PPMM@PVA, as well as PPMM@PDA/PVA(1/2, 1/4, 1/8), PPMM@PDA/PVA(1/8)@Au, and PPMM@PDA/PVA(1/8)@Au@IrO₂, were observed by a scanning electron microscope (SEM, JEOL JSM6700F). Measurements for water contact angles were performed by a SURFTENS-Basic (OEG Co.) to determine the hydrophilicity of PPMM, PPMM@PDA, PPMM@PVA, and PPMM@PDA/PVA(1/2, 1/4, 1/8) in size of 1 × 1 cm². X-ray photoelectron spectroscopy (XPS; Thermo Fisher Scientific, ESCALAB XI⁺) was employed to determine the nature of oxygenated functional groups, and their relative amounts for PPMM and PPMM@PDA/PVA(1/8). Bending test was conducted in a custom-made mechanical device by which the PPMM@PDA/PVA(1/8)@Au@IrO₂ was bended to 60° repeatedly for 5000 cycles.

To determine its pH-sensing responses, the PPMM@PDA/PVA(1/8)@Au@IrO₂ in size of 1 × 1 cm² was immersed in Britton-Robinson buffer solutions (0.04 M H₂BO₃, 0.04 M CH₃COOH, and 0.04 M H₃PO₄) with pH values adjusted to 2, 3, 4, 6, 7, 8, 10, 11, and 12, respectively. The open-circuit potential (OCP) for PPMM@PDA/PVA(1/8)@Au@IrO₂ in Britton-Robinson buffer solutions with different pH values was recorded for 200 s to obtain its average and standard deviation. Identical experiments were carried out for samples after bending 1000, 3000, and 5000 cycles, respectively. The hysteresis behavior was explored by recording the OCP in Britton-Robinson buffer solutions with pH value adjusted in the order of 2–3–4–6–7–8–10–11–12–10–8–7–6–4–3–
In each measurement, the OCP was recorded for 200 s and the sample was rinsed with deionized water afterwards. All those electrochemical experiments were conducted by a potentiostat (VersaSTAT4) at 25 °C. The SCE and Pt foil (3 × 3 cm²) were used as the reference and counter electrodes, respectively.

**Cell viability test.** To evaluate any possible bio-toxicity from PPMM@PDA/PVA(1/8)@Au or PPMM@PDA/PVA(1/8)@Au/IrO₂, a cell viability test was conducted in which L929 cells (mouse lung fibroblasts, ATCC, USA) were cultured within our samples for 1 and 4 days. The cell viability test entailed the CellTiter 96° AO™ One Solution Cell Proliferation Assay Kit (Promega Co.) and 3-(4,5-dimethylthiazol-2-yl)-5(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium (MTS). The experiments were carried out in three different groups; 1) 48-well plate only (serving as the control group), 2) PPMM@PDA/PVA(1/8)@Au, and 3) PPMM@PDA/PVA(1/8)@Au/IrO₂. First, the samples in 0.5 × 0.5 cm² were dipped into ethanol and rinsed with phosphate-buffered saline. Afterwards, they were dried in an oven at 50 °C for 48 h. Next, the samples were positioned in 48-well plates for cell viability tests. Before the tests, all samples were sterilized under UV illumination (Germicidal lamp GL15, Sankyo Denki) for 24 h at 25 °C. In each group, every individual well was seeded with 400 μL cell suspension which contained L929 cells, Dulbecco's Modified Eagle Medium (DMEM; Sigma-Aldrich), and 1 vol% antibiotics (Antibiotic–Antimycotic (100X), Gibco®). The effective cell concentration in suspension was 1 × 10⁴/mL. Subsequently, the 48-well plates were maintained in an incubator (5% CO₂, 37 °C) for 96 h. The control sample was maintained in DMEM without the cell suspension, antibiotics, and cell culture medium. The determination of cell viability was undertaken by following the absorbance measurements at 450 nm (Epoch™ 2, BioTek Instruments Inc.) and the cell viability was determined by following formula.

\[
\text{Cell viability} = \frac{A_{\text{exp}}}{A_{\text{control}}} \times 100\%
\]

where the \( A_{\text{exp}} \) is the absorbance for our sample, and the \( A_{\text{control}} \) is the absorbance for the control group.

**Results and discussion**

**Material characterization.** Figure 1 displays the SEM images for PPMM, PPMM@PVA, PPMM@PDA, and PPMM@PDA/PVA(1/2, 1/4, 1/8), respectively. In Fig. 1a, the PPMM demonstrated a non-woven microstructure in which individual PP fibers revealed a smooth surface with fiber diameters between 0.9 and 6.5 μm. In Fig. 1b, the PPMM@PVA demonstrated a similar morphology as the deposition of PVA occurred exclusively on the external surface of PPMM. This was anticipated due to the hydrophilic nature of PVA that limited its infiltration and formation of intimate bonding with the hydrophobic PPMM. For PPMM@PDA, shown in Fig. 1c, there appeared many agglomerates in sizes of 60 ~ 300 nm on the surface of PP fibers. These agglomerates were PDA deposited from the solution via the self-polymerization of DA. Since the DA contained a phenol group, the PDA was likely to form CH₃-π bonding with the PP fibers, resulting in a more pronounced deposition. However, the formation of PDA on PP fibers involved an oxidation reaction from the participation of dissolved oxygen. As a result, the PDA deposition was also observed on the external surface of PPMM. In Fig. 1d–f, the images of PPMM@PDA/PVA(1/2, 1/4, 1/8) demonstrated significantly improved surface morphologies, and the PDA agglomerates were formed uniformly on individual PP fibers. Since there was strong hydrogen bonding between the PVA and DA, the PDA was able to facilitate homogeneous formation of PDA with smaller agglomerate sizes. This allowed the PDA/PVA to infiltrate to the internal space of entire PPMM and produced a conformal coating on every PP fibers. Interestingly, with a smaller PDA/PVA ratio, the surface morphology became more refined with smaller PDA agglomerates. In our experiments, the smallest PDA/PVA mass ratio was 1/8 as additional PVA amounts caused preferential PDA deposition on the external surface of PPMM again.

The hydrophilicity of functionalized PPMM is critical for successful deposition of Au and IrO₂. Figure 2 displays the dynamic water contact angles for PPMM, PPMM@PDA, PPMM@PVA, and PPMM@PDA/PVA(1/2, 1/4, 1/8), respectively. The insets are the photographs for water droplets at 12 min. As shown, the PPMM exhibited a superhydrophobic behavior with a contact angle greater than 120°. This is anticipated as the PPMM consists of nonpolar PP fibers, and is designed for filtering of nonaqueous solution. After coating with PVA, the contact angle for PPMM@PVA was 117.4° initially, and became slightly reduced to 105.9° after 12 min. The hydrophobic behavior of PPMM@PVA suggested that the PVA was not forming a conformal coating on individual PP fibers of PPMM. Instead, only the PP fibers on the external surface of PPMM were partially covered, as evidenced by Fig. 1b. For PPMM@PDA, a similar pattern was recorded with an initial contact angle of 109° that decreased to 105° after 12 min. Again, the hydrophilic PDA agglomerates were predominately deposited on the external surface of PPMM, as evidenced by Fig. 1c, so the PPMM@PDA still revealed a hydrophobic behavior. In contrast, the PPMM@PDA/PVA(1/2, 1/4, 1/8) exhibited substantial improvements in hydrophilicity. For example, their starting contact angles were consistently smaller than 90°, and became even smaller as time progressed, suggesting a steady infiltration of water. Notably, with a smaller PDA/PVA ratio, the hydrophilicity became even more pronounced. For instance, the PPMM@PDA/PVA(1/8) revealed a notable hydrophilic behavior, and the water droplet became completely flat after 12 min. We recognized that the hydrophilic nature of PPMM@PDA/PVA(1/2, 1/4, 1/8) was associated with the conformal coating of PDA/PVA on the PP fibers. In particular, the PPMM@PDA/PVA(1/8) demonstrated the most desirable hydrophilic behavior, and therefore was selected as the flexible substrate for following deposition of Au and IrO₂ nanoparticles.
The XPS was employed to elucidate the chemical nature of functionalized PPMM. Figure 3 displays the XPS profiles and fitting curves for C1s and N1s from PPMM and PPMM@PDA/PVA(1/8). As shown in Fig. 3a, the C1s signal from PPMM revealed the characteristic C–C bonding from the primary molecular structure of PP. However, the curve fitting result indicated the presence of minor signals which were associated with the alcohol/amine (C–O/C–N) and ketone (C=O) bonds. Those peaks were presumably caused by the additives in the PPMM fabrication process.

In contrast, the curve fitting of C1s profile from PPMM@PDA/PVA(1/8), shown in Fig. 3b, exhibited stronger intensities for both C–O/C–N and C = O bonds, as well as moderate presence of carboxyl (O–C = O) bond. Since the PDA contains phenol, amine, and ketone groups in its chemical structure, and the PVA is synthesized by partial or fully hydrolysis of polyvinyl acetate, the coexistence of O–C = O, C–O, C = O, and C–N was reasonably expected. Figure 3c,d display the N1s profiles for PPMM and PPMM@PDA/PVA(1/8), respectively. The curve fitting for PPMM, shown in Fig. 3c, exhibited two distinct peaks associated with –N = and –NH–, which were additives in the PPMM fabrication process. In Fig. 3d of PPMM@PDA/PVA(1/8), the predominant signal consisted of –N= and –NH–, but a minor one associated with –NH2 was also recorded. According to Liebscher et al., three distinct amine groups (primary, secondary, tertiary) might be present in PDA, contingent on its...

**Figure 1.** The SEM images for (a) PPMM, (b) PPMM@PVA, and (c) PPMM@PDA, as well as (d) PPMM@PDA/PVA(1/2), (e) PPMM@PDA/PVA(1/4), and (f) PPMM@PDA/PVA(1/8), respectively. The insets are their respective high magnification images.
degree of oxidation. Therefore, the presence of amine (–NH$_2$) group and an increasing ratio of N/C provided the solid evidence validating the existence of PDA. Table 1 lists the XPS fitting results of C1s and N1s signals from PPMM and PPMM@PDA/PVA(1/8), respectively.

Figure 4 displays the SEM images for PPMM@PDA/PVA(1/8)@Au and PPMM@PDA/PVA(1/8)@Au@IrO$_2$ in low and high magnification views. As shown in Fig. 4a, the PPMM@PDA/PVA(1/8)@Au in low magnification view demonstrated a uniform coating of Au. In high magnification view, shown in Fig. 4b, there appeared a notable coalescence of Au grains on individual PP fibers. The thickness of conformal Au overcoat was around 150 nm. In Fig. 4c, the PPMM@PDA/PVA(1/8)@Au@IrO$_2$ in low magnification view exhibited a rough surface, a phenomenon attributed to the deposition of IrO$_2$ nanoparticles. The high magnification view, shown in Fig. 4d, confirmed the presence of IrO$_2$ nanoparticles on the surface of Au grains. In addition, the size of IrO$_2$ nanoparticles was around 50 nm.

Analysis of pH-sensing performance. Figure 5a displays the OCP of PPMM@PDA/PVA(1/8)@Au and PPMM@PDA/PVA(1/8)@Au@IrO$_2$ in Britton-Robinson buffer solutions with pH values adjusted from 2 to 12. The corresponding pH-sensing slopes for both samples were −49.35 and −74.45 mV/pH, respectively. For pH-sensing, the Au was primarily used as a current collector instead of functioning as an active pH-sensing element. In addition, the Au was reported to reveal a pH sensing slope of −53.4 mV/pH, a value that was rather close to what we observed. On the other hand, the IrO$_2$ has been established as an impressive pH-sensing material as it is known to show a super-Nernstian response. In the literature, the pH sensitivity of IrO$_2$ was found to be affected by the preparation methods involved. For example, from solution-based processes, the surface of IrO$_2$ was mostly occupied by hydrate IrO$_2$, and therefore a super-Nernstian behavior with a pH-sensing slope between −60 and −80 mV/pH was anticipated. The redox potential behavior for super-Nernstian response is listed below.

\[
E = E_0 - \frac{mRT}{nF}pH
\]  

(2)

where the $E$ is the redox potential, the $E_0$ is the standard redox potential, the $n$ and $m$ are the numbers of electrons and protons involved in the redox reaction, the $R$ is the ideal gas constant, the $T$ is the absolute temperature, and the $F$ is the Faraday constant. In the case of hydrated IrO$_2$, the redox reaction is listed below.

\[
\text{Ir}_2\text{O}((\text{OH})_3\text{O}_3^{3-} + 3\text{H}^+ + 2\text{e}^- \leftrightarrow 2\text{Ir(OH)}_2\text{O}^- + \text{H}_2\text{O}
\]  

(3)

We substituted those known constants into the equation, and the theoretical pH-sensing slope could be expressed as:
In our pH-sensing results, the PPMM@PDA/PVA(1/8)@Au@IrO2 exhibited a super-Nernstian response with a pH-sensing slope of −74.45 mV/pH. This value was slightly smaller than the theoretical value of −88.7 mV/pH because according to our earlier XPS analysis, our sample was consisted of both hydrated and anhydrous IrO2. It is known that the anhydrous IrO2 reveals an inferior sensing response to that of hydrated IrO2. The redox reaction of anhydrous IrO2 is listed in Eq. (5), and the sensing slope is expressed in Eq. (6) as −59.1 mV/pH. Therefore, the expected overall sensitivity for our sample is between −59.1 and −88.65 mV/pH.

![Figure 3](image)

**Table 1.** The XPS fitting results of C1s and N1s profiles from PPMM and PPMM@PDA/PVA(1/8), respectively.

| Compound type | C–C sp²/sp³ | C–O/C–N | C=O | O–C=O |
|---------------|-------------|---------|-----|-------|
| PPMM          | 83.5 at%    | 13.1 at%| 3.4 at%| 0     |
| PPMM@PDA/PVA(1/8) | 63.8 at% | 29.1 at%| 4.5 at%| 2.6 at%|

| Compound type | Atomic ratio |
|---------------|--------------|
| −N=           | 47.2 at%     |
| −NH–          | 52.8 at%     |
| −NH₂          | 0            |
| N/C           | 0.057        |

\[
(-m/n) \left( \frac{RT}{F} \right) = (-3/2)(0.05916) = -88.7 \text{ mV/pH}, \quad \text{at } 25^\circ \text{C}
\]  

\[
(\text{5})
\]

The redox reaction of anhydrous IrO₂ is listed in Eq. (5), and the sensing slope is expressed in Eq. (6) as −59.1 mV/pH. Therefore, the expected overall sensitivity for our sample is between −59.1 and −88.65 mV/pH.

\[
2\text{IrO}_2 + 2\text{H}^+ + 2\text{e}^- \leftrightarrow \text{Ir}_2\text{O}_3 + \text{H}_2\text{O}
\]  

\[
(\text{5})
\]
We also obtained the pH-sensing slopes in different pH ranges to further explore the pH sensitivity of PPMM@PDA/PVA(1/8)@Au@IrO2. Figure 5b displays the corresponding bar charts of pH-sensing slope in aqueous solutions with different pH values. Apparently, the PPMM@PDA/PVA(1/8)@Au@IrO2 revealed a higher slope of −78.63 mV/pH in solutions with the pH range of 4 to 10. It is noted that the pH values for human's...

\[
(-\frac{m}{n}) \left( \frac{RT}{F} \right) = (-2/2)(0.05916) = -59.1 \frac{mV}{pH}, \text{ at } 25^\circ C
\]  

Figure 4. The SEM images for PPMM@PDA/PVA(1/8)@Au in (a) low and (b) high magnification views, as well as PPMM@PDA/PVA(1/8)@Au@IrO2 in (c) low and (d) high magnification views.

Figure 5. (a) The potentiometric response of PPMM@PDA/PVA(1/8)@Au and PPMM@PDA/PVA(1/8)@Au@IrO2 in aqueous solutions with pH values adjusted from 2 to 12. (b) The corresponding pH-sensing sensitivities of PPMM@PDA/PVA(1/8)@Au@IrO2 in aqueous solutions of different pH ranges.
blood and urine are 7.3 to 7.4 and 4.5 to 7.8, respectively. Thus, our PPMM@PDA/PVA(1/8)@Au@IrO2 not only is suitable for wearable electronics but also is expected to be potentially useful as a pH sensor in in vivo environment.

To validate the flexibility of PPMM@PDA/PVA(1/8)@Au@IrO2 and its pH-sensing performance after multiple bending cycles, we tested the sample under two different bending directions, as shown in Fig. 6a. Figure 6b displays the photographs for bending tests in which the sample was bended along the X axis and Y axis to 60° and 90°, respectively. We recorded the pH-sensing slopes at the 0th, 1000th, 3000th, and 5000th cycles, respectively. Figure 6c displays the corresponding pH-sensing slopes and standard deviations for PPMM@PDA/PVA(1/8)@Au@IrO2 undergoing bending action along the X axis. The sensitivity over repeated bending was relatively steady without noticeable deterioration, an expected behavior as any physical deformation was confined on the metalized PPMM exclusively. In contrast, for the sample undergoing bending along the Y axis, the area undergoing deformation covered both metalized PPMM and IrO2. The corresponding pH-sensing behavior, as shown in Fig. 6d, revealed minor deterioration at the 1000th cycle, followed by stable sensing responses afterwards. Nonetheless, the pH-sensing slope still retained the desirable super-Nernstian response within 5000 bending cycles, which validated the flexible and durable nature of PPMM@PDA/PVA(1/8)@Au@IrO2.

Figure 7a displays the OCP of PPMM@PDA/PVA(1/8)@Au@IrO2 in Britton-Robinson buffer solutions with different pH values. The sample was immersed in the solution in which the pH value was adjusted in the order of pH 2–3–4–6–7–8–10–11–12–11–10–8–7–6–4–3–2. The hysteresis behavior, defined as the hysteresis potential (dV), is determined by the difference of the OCPs when the sample is immersed at different times in the solution with identical pH values. Figure 7b displays the hysteresis potentials at respective pH values. Apparently, under an
acidic solution, the sample revealed a relatively smaller hysteresis potential, which was likely caused by the faster ionic exchange of protons in an acidic solution. In a neutral solution, the concentration of proton or hydroxyl ion was relatively low which led to a slower ionic exchange rate on the electrode surface. The greater hysteresis phenomena in an alkaline solution was attributed to the strong interaction between the hydroxyl groups and oxide surface as the IrO$_2$ was unstable in an alkaline solution.

To investigate the stability of our sample, we measured the variation of OCP in acidic, neutral, and alkaline solutions. Figure 8 displays the drift effect at pH values of 2, 7, and 12 for 7200 s. The drift potential was obtained by comparing the average potential for the final 200 s and the first 200 s. The drift rate at pH values of 2, 7, and 12 was 0.14, −0.76, and 57.75 mV/h, respectively. Apparently, our sample revealed an impressive stability in both acidic and neutral solutions. However, at pH 12, the hydrated IrO$_2$ became unstable so the drift behavior was rather poor. Since in the in-vivo condition, the body fluid is either neutral or weakly acidic, the drift performance for our sample is proven to be useful for implantable electronics.

We have compared our sample with what have been published in the literature. The comparison is listed in Table 2 below. As listed, among these flexible pH sensors, our sample demonstrates a superior sensitivity which is attributed to the presence of hydrated IrO$_2$ from our wet chemical synthetic route. It is known that hydrated IrO$_2$ exhibits a better pH sensing behavior than that of anhydrous IrO$_2$.

Figure 9 displays the bar chart for cell viability experiments. After 1 day of cell culture, the cell viability for PPMM@PDA/PVA(1/8)@Au and PPMM@PDA/PVA(1/8)@Au@IrO$_2$ was 91.2% and 90.6% of that of blank sample, which served as the control group. Interestingly, after 4 days, their cell viability was slightly increased to 96.1% and 93.5%, respectively. This moderate improvement in cell viability was possibly attributed to the presence...
of IrO2 because the IrO2 was reported to promote cell growth in the literature42. From our results, both PPMM@PDA/PVA(1/8)@Au and PPMM@PDA/PVA(1/8)@Au@IrO2 revealed impressive biocompatibility, which was defined with a cell viability above 80%.

In fabricating the flexible pH sensor for possible uses in wearable and implantable electronics, the materials we employed (PPMM, PDA, PVA, Au, IrO2) are known for their biocompatibility. In addition, the chemicals involved during the synthetic process were deliberately chosen to minimize any residual toxicity. For example, a typical electroless Au deposition requires a sequential sensitization and activation process using Sn and Pd ions to prepare Pd nuclei for subsequent Ni electroless deposition43. The Ni was then served as the seeds for Au deposition. However, both Pd and Ni exhibit moderate biotoxicity so their presence is not desirable. Therefore, the use of PDA for Au nuclei formation gave us an alternative biocompatible route without involving toxic chemicals for sensitization and activation. Since the PDA was deposited on the surface of PP fibers, the reduction of Au3+ ions for Au nuclei took place exclusively on the PP fibers instead of homogeneously in the solution. In addition, conventional electroless Au depositions entail reducing agents such as N2H4 and NH2OH·HCl, and those chemicals are not biocompatible. On the contrary, in our process the reducing agent was the sodium L-ascorbate because it not only reveals a mild reducing power but also is known for its biocompatibility.

It is further noted that during pH sensing, the IrO2 is not susceptible to interference from monovalent cations (Na+, K+) and divalent cations (Mg2+)28,44,45. In addition, the selection of PPMM over conventional polymeric substrates such as polyimide (PI), polyamide (PA), polytetrafluoroethylene (PTFE), polyethylene terephthalate (PET), and polyvinylchloride (PVC) is based on its relatively low cost, impressive corrosion resistance, and excellent biocompatibility. More importantly, the porous structure of PPMM could be readily tailored to meet specific applications.

**Conclusions**

A flexible pH sensor was fabricated by carrying out deposition of Au and IrO2 on individual PP fibers of functionalized PPMM. Both PDA and PVA were used as the surface agents to improve the hydrophilicity of otherwise hydrophobic PPMM. The SEM images validated the formation of continuous Au overcoat on individual PP fibers, and the deposition of IrO2 nanoparticles with moderate coalescence. Contact angle measurements indicated that a molar ratio of PDA/PVA = 1/8 rendered the most hydrophilic PPMM. The pH-sensing behaviors were explored in aqueous solutions with the pH values adjusted between 2 and 12. The PPMM@PDA/PVA(1/8)@Au@IrO2 revealed a super-Nernstian response for a sensing slope of −74.45 mV/pH. In addition, the PPMM@PDA/PVA(1/8)@Au@IrO2 maintained its sensing ability after repeated bending up to 5000 cycles. The PPMM@

|                      | pH sensitivity (mV/pH) | pH sensing range | References |
|----------------------|------------------------|------------------|------------|
| Our sample           | −74.45                 | 2–12             | This work  |
| Flexible PANI membrane | −58.57               | 5.45–8.62         | 38         |
| Flexible IrO2/AgCl electrode | −51.7               | 1.5–12.1         | 38         |
| Polyaniline nanopillar sensor | −60.3               | 2.09–12.0         | 38         |
| Flexible graphene pH sensor | −62                 | 5–8              | 38         |
| Au composite strip-based sensor | −57                | 3–8              | 38         |

Table 2. The comparison of pH sensing sensitivity among different flexible pH sensors in the literature.

![Figure 9.](image-url)
PDA/PVA(1/8)@Au@IrO₂ revealed a subdued hysteresis behavior in an acidic solution. The raw materials and chemicals used in our sample preparation were deliberately chosen for their compatibility and the resulting PPM@PDA/PVA(1/8)@Au@IrO₂ was validated in cell viability test.

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**Author contributions**
S.C.C. and Y.C.H. designed the idea and developed the method. W.H.C., B.Y.S., and C.Y.C. contributed to the analysis and experiment. The manuscript was written by S.C.C. and further edited by S.Y.C., J.C.C, and P.W.W.

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**Competing interests**
The authors declare no competing interests.

**Additional information**
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