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Strong adhesion of wet conducting polymers on diverse substrates

Akihisa Inoue1,2*, Hyunwoo Yuk1‡, Baoyang Lu1,3,4, Xuanhe Zhao1,5†

Conducting polymers such as poly(3,4-ethylenedioxythiophene):poly(styrene sulfonate) (PEDOT:PSS), polypyrrole (PPy), and polyaniline (PAni) have been widely explored as electrodes and coatings for electrodes that interface with biological organisms in bioelectronic devices (1–8), owing to their favorable electrochemical stability (9), electrical and mechanical properties (10–16), and biocompatibility (13, 17, 18). However, weak and unstable adhesion of conducting polymers on substrates and devices can result in interface failures (e.g., debonding of conducting polymers) and subsequent loss of functionality, substantially hampering the reliability and efficacy of bioelectronic devices (5, 6, 19–23). In light of this challenge, a few methods have been reported to provide enhanced adhesion of conducting polymers in wet environment. For example, topological modification of substrates (e.g., nano- and microscale roughness on gold substrates) (24, 25) and electrodeposition of chemically modified EDOT monomers (e.g., vinyl-, carboxylic-, and amine-modified EDOT) (23, 26, 27) have been adopted to improve adhesion of wet PEDOT:PSS. However, the previous approaches require specific types of substrates and/or complicated modification of EDOT monomer, and they are limited to electrodeposited PEDOT:PSS on conductive substrates, substantially limiting the applicability of conducting polymers in bioelectronics devices.

Here, we report a general yet simple method to achieve strong adhesion of various wet conducting polymers including PEDOT:PSS, PPy, and PAni on diverse commonly used insulating and conductive substrates including glass, polymide, polydimethylsiloxane (PDMS), indium tin oxide (ITO), and gold. The method is based on introducing a hydrophilic polymer adhesive layer with a thickness of a few nanometers between the substrate and the conducting polymer. The adhesive layer forms strong adhesion with the substrate and an interpenetrating polymer network with the conducting polymer. To implement the method, we choose hydrophilic polyurethane (PU) to form the nanometer-thick adhesive layer. The adhesive layer can be facilely introduced on a broad range of substrate materials via diverse fabrication approaches including spin coating, spray coating, and dip coating. The resultant interfacial adhesion between wet conducting polymers and various substrates is strong (e.g., lap-shear strength over 120 kPa) and mechanically and electrochemically stable (e.g., no observable interfacial failure after 60 min of ultrasonication and 10,000 charging and discharging cycles). Our method also allows the use of commercially available conducting polymers and hydrophilic PU without the need for complex chemical synthesis or substrate modification, promising ready and broad applicability to bioelectronic devices. Furthermore, the method is compatible with various fabrication approaches for conducting polymers including solvent casting and electrodeposition, without compromising the electrical or mechanical properties of conducting polymers.

INTRODUCTION
Conducting polymers such as poly(3,4-ethylenedioxythiophene):poly(styrene sulfonate) (PEDOT:PSS), polypyrrole (PPy), and polyaniline (PAni) have attracted great attention as promising electrodes that interface with biological organisms. However, weak and unstable adhesion of conducting polymers to substrates and devices in wet physiological environment has greatly limited their utility and reliability. Here, we report a general yet simple method to achieve strong adhesion of various conducting polymers on diverse insulating and conductive substrates in wet physiological environment. The method is based on introducing a hydrophilic polymer adhesive layer with a thickness of a few nanometers, which forms strong adhesion with the substrate and an interpenetrating polymer network with the conducting polymer. The method is compatible with various fabrication approaches for conducting polymers without compromising their electrical or mechanical properties. We further demonstrate adhesion of wet conducting polymers on representative bioelectronic devices with high adhesion strength, conductivity, and mechanical and electrochemical stability.

RESULTS
Method for strong adhesion of wet conducting polymers on substrates
Instead of modifying a specific conducting polymer or substrate, we propose a general method to achieve strong adhesion of various wet conducting polymers on diverse substrates by using a hydrophilic polymer adhesive layer with a thickness of a few nanometers, as illustrated in Fig. 1A. To avoid effects from substrates’ surface roughness, we use substrates with smooth surfaces (i.e., mean roughness below 5 nm) in this study (fig. S1). A substrate is first functionalized with primary amine groups (fig. S2), which can provide an enhanced interfacial adhesion between the substrate and the adhesive layer via covalent bonds and/or electrostatic interactions (28). Thereafter, the adhesive layer can be applied to a wide range of substrates by various coating approaches including spin coating, spray coating, and dip coating. Conducting polymers can then be prepared on the substrate with the adhesive layer from conducting polymer

1Department of Mechanical Engineering, Massachusetts Institute of Technology, Cambridge, MA 02139, USA. 2JSR Corporation, Tokyo 1050021, Japan. 3Flexible Electronics Innovation Institute, Jiangxi Science and Technology Normal University, Nanchang 330013, China. 4School of Pharmacy, Jiangxi Science and Technology Normal University, Nanchang 330013, China. 5Department of Civil and Environmental Engineering, Massachusetts Institute of Technology, Cambridge, MA 02139, USA.
*These authors contributed equally to this work.
†Corresponding author. Email: zhaox@mit.edu
precursors via various approaches such as solvent casting of aqueous conducting polymer solutions (e.g., PEDOT:PSS, PPy, and PAni) and electrodeposition of aqueous monomer solutions (e.g., EDOT:PSS). The hydrophilicity and subsequent swelling of the adhesive layer allow diffusion of the conducting polymer precursors throughout the adhesive layer (29), subsequently forming an interpenetrating polymer network between the infiltrated conducting polymer and the adhesive layer (30). The strong adhesion between the adhesive

Fig. 1. Strong adhesion of wet conducting polymers on diverse substrates. (A) Strong adhesion of a wet conducting polymer on an amine-functionalized substrate with a hydrophilic polymer adhesive layer. (B) Strong adhesion of a conducting polymer on a substrate with an adhesive layer in wet environment. (C) Solvent-casted wet PEDOT:PSS (10 μm thickness) on a polyimide substrate with a PU adhesive layer (60 nm thickness) can withstand mechanical deformations including twisting, bending, and even folding (the radius of curvature ~ 250 μm) without interfacial failure. Photo credit: Hyunwoo Yuk, MIT. (D) Weak and unstable adhesion of a conducting polymer on a substrate without the adhesive layer in wet environment. (E) Solvent-casted wet PEDOT:PSS (10 μm thickness) on a polyimide substrate without the adhesive layer undergoes interfacial failure upon mechanical deformation of the substrate. Photo credit: Hyunwoo Yuk, MIT. (F) Distribution of carbon and sulfur atoms in a PU adhesive interface between an adhered PEDOT:PSS (5 μm thickness) and a PU-coated silicon substrate (1500 nm PU thickness).
layer and the substrate, and the interpenetration between the adhesive layer and the conducting polymer synergistically provide mechanically robust adhesion against interfacial failure in wet physiological environment (Fig. 1B). Furthermore, the conducting polymer interpenetrates throughout the adhesive layer and directly contacts the underlying substrate, providing electrical conductivity through the adhesive layer.

The proposed method can achieve strong adhesion of various wet conducting polymers on diverse substrates that are inaccessible to previous bonding methods. For example, a solvent-casted wet PEDOT:PSS on an insulating and flexible polyimide substrate with the PU adhesive layer exhibits robust adhesion without failure under twisting, bending, and even folding (with the radius of curvature ~ 250 μm) (Fig. 1C). In contrast, a wet PEDOT:PSS on substrates (e.g., glass and polyimide) without the PU adhesive layer exhibits interfacial failure upon swelling in phosphate-buffered saline (PBS; Fig. 1, D and E, and fig. S3B) due to weak and unstable adhesion formed by physical interactions of wet conducting polymers with the substrate surfaces (Fig. 1D) (23). To validate the formation of an interpenetrating layer between a conducting polymer and a hydrophilic PU adhesive layer, we investigate an elemental dispersion of carbon and sulfur atoms in a cross section of the PU adhesive layer of a cryo-fractured sample (solvent-casted PEDOT:PSS on Si wafer with the PU adhesive layer) by scanning electron microscopy–energy-dispersive x-ray spectroscopy (SEM-EDS). Because both PEDOT and PSS contain sulfur, while the hydrophilic PU is absent of sulfur (fig. S4), the sulfur signal can serve as an indicator for the interpenetration of the PEDOT:PSS into the PU adhesive layer. It is evident that the sulfur signal can be detected in the PU adhesive layer, indicating the interpenetration of the PEDOT:PSS throughout the PU adhesive layer (Fig. 1F). We further investigate the interpenetration between the PEDOT:PSS and the PU adhesive layer by comparing x-ray photoelectron spectroscopy (XPS) spectra of the surface of a wet PEDOT:PSS and the surfaces of 60-nm and 1500-nm PU adhesive layers with PEDOT:PSS interpenetrated from the other side (fig. S5; see the Supplementary Materials for preparation and testing of the samples). We find that the thin PU adhesive layer (60 nm) exhibits almost the same sulfur S(2p) spectral intensity as the PEDOT:PSS layer (fig. S5, D and E), whereas the thick PU adhesive layer (1500 nm) shows substantially lower sulfur S(2p) spectral intensity compared to the PEDOT:PSS layer (fig. S5F). These results indicate that the conducting polymer can interpenetrate throughout the thin adhesive layer (60 nm) but only partially through the thick adhesive layer (1500 nm).

**Adhesion strength of wet conducting polymers on diverse substrates**

To measure adhesion strength of wet conducting polymers on diverse substrates, we perform lap-shear tests to measure interfacial shear strength (ASTM D3163) between the adhered wet conducting polymer and the substrate (Fig. 2). Figure 2A shows a typical lap-shear test sample and a setup in which a wet conducting polymer is sandwiched between a hydrophilic microporous nylon filter (pore size, 1 μm) and a substrate with the PU adhesive layer. Note that we choose hydrophilic microporous nylon filters as the backing for the laboratory-shear tests due to two reasons: (i) The nylon filters’ high mechanical stiffness (Young’s modulus, >1 GPa) can prevent undesirable deformation of wet conducting polymers during lap-shear tests. (ii) Hydrophilicity and microporosity of the nylon filters allow infiltration of conducting polymers and subsequent reliable mechanical integration during lap-shear tests. Wet PEDOT:PSS on glass substrates with the PU adhesive layers exhibits a high shear strength (54 kPa), while wet PEDOT:PSS on pristine glass substrates without the PU adhesive layer shows a very low shear strength (0.08 kPa) and easily detaches from the substrate (Fig. 2, B and C).

We further investigate the effect of surface functionalization of substrates with primary amine groups on the adhesion performance. We compare shear strength and modes of failure in lap-shear tests for wet PEDOT:PSS on PU-coated glass substrates with and without primary amine functionalization. The wet PEDOT:PSS on the amine-functionalized glass substrate with the PU adhesive layer exhibits cohesive failure (i.e., failure occurs within wet PEDOT:PSS) (Fig. 2C), while the wet PEDOT:PSS on the pristine glass substrate with the PU adhesive layer demonstrates adhesive failure (i.e., failure occurs between the PU adhesive layer and the substrate) (Fig. 2B). Moreover, the amine-functionalized glass substrate with the PU adhesive layer provides a much higher shear strength than the pristine glass substrate with the PU adhesive layer (160 kPa versus 54 kPa) (Fig. 2D and fig. S6A). Hence, the amine functionalization is a critical step to achieve strong adhesion between the PU adhesive layer and the substrate due to enhanced adhesion of hydrophilic PU on amine-functionalized substrates (30). We also find that the PU adhesive layer can provide strong adhesion of wet PEDOT:PSS (shear strength over 120 kPa) consistently over a wide range of the PU layer’s thickness (6 to 1500 nm) (fig. S7).

The proposed method is also applicable to various wet conducting polymers as well as a wide range of insulating and conductive substrates (Fig. 2, E and F). Solvent-casted wet PEDOT:PSS, PPy, and PANi on amine-functionalized glass substrates with the PU adhesive layers provide substantially higher shear strengths than the counterparts on pristine glass substrates without the PU adhesive layer (160 kPa versus 0.1 kPa for PEDOT:PSS; 39 kPa versus 0.4 kPa for PPy; 37 kPa versus 1.9 kPa for PANi) (Fig. 2E). Note that all conducting polymers undergo cohesive failure in the laboratory-shear tests, indicating that the adhesive interface with the PU adhesive layer is stronger than the bulk wet conducting polymer and that the measured adhesion strength is determined by the shear strength of the bulk wet conducting polymer. The proposed method also enables strong adhesion of wet PEDOT:PSS on diverse amine-functionalized insulating and conductive substrates including glass (shear strength, 160 kPa), polyimide (shear strength, 116 kPa), PDMS (shear strength, 111 kPa), ITO-glass (shear strength, 149 kPa), and gold (shear strength, 146 kPa) (Fig. 2F and figs. S6 and S8). We find that wet PEDOT:PSS on polyimide substrates with the PU adhesive layers exhibits negligible enhancement in the shear strength by amine functionalization (fig. S6C), which might result from good inherent adhesion of the hydrophilic PU on polyimide (fig. S9).

**Electrical and mechanical properties of adhered wet conducting polymers**

While the hydrophilic PU adhesive layers can provide strong adhesion between wet conducting polymers and substrates, they should not affect the electrical or mechanical properties of the conducting polymers in applications. To investigate this requirement, we systematically characterize the electrical and mechanical properties of PEDOT:PSS adhered by the PU adhesive layer as a representative conducting polymer (Fig. 3 and fig. S10). We measure the electrical conductivity of wet PEDOT:PSS without and with the PU adhesive
layer of varying thickness by a standard four-point probe method in wet physiological environment (hydrated in PBS) (Fig. 3A). The electrical conductivity of the wet PEDOT:PSS exhibits no statistically significant differences ($P > 0.5$) over a wide range of the PU adhesive layer thickness (6 to 1500 nm), indicating insignificant effect of the PU adhesive layer on the electrical properties of wet PEDOT:PSS (Fig. 3B). To probe the effect of the PU adhesive layer on electrical conductivity between wet PEDOT:PSS and underlying electrodes, we measure the sheet resistance of the wet PEDOT:PSS on the amine-functionalized glass substrates without the PU adhesive layer and amine-functionalized glass substrates with the PU adhesive layer. (D) Lap-shear strengths of a wet PEDOT:PSS on various amine-functionalized insulating and conductive substrates with the PU adhesive layers. Conducting polymers with 50 $\mu$m thickness and PU adhesive layers with 60 nm thickness were used for all experiments. Values in (C) to (E) represent the mean, and the error bars represent 95% CI of the measured values ($n = 5$). $P$ values are determined by Student’s $t$ test. **$P \leq 0.01$; ***$P \leq 0.001$.

To further investigate the effect of the PU adhesive layer’s thickness on electrical property of the adhesive interface, we perform electrochemical impedance spectroscopy (EIS) analysis of wet PEDOT:PSS on amine-functionalized ITO-glass electrodes without and with the PU adhesive layer (60 or 1500 nm thickness) (Fig. 3E and fig. S11) (31). It can be seen that the PEDOT:PSS adhered on the ITO-glass electrode via the thin PU adhesive layer (60 nm) can provide favorable electrical property (i.e., lower impedance than the bare ITO-glass electrode in the low frequency region) due to enhanced capacitance of the electrode from the adhered wet polymer and the underlying electrode, whereas the sufficiently thin (i.e., nanometers thick) PU adhesive layer can provide low interfacial resistance.
that the nanometer-thick PU adhesive layer is essential to provide highly conductive interface between the adhered wet conducting polymers and the underlying electrodes.

To quantitatively evaluate the effect of the PU adhesive layer on the mechanical properties of wet PEDOT:PSS, we perform tensile tests of solvent-casted wet PEDOT:PSS in PBS without and with the PU adhesive layer of varying thickness (fig. S11A). The Young’s moduli and the ultimate tensile strain of the wet PEDOT:PSS (100 µm thickness) with the PU adhesive layer of varying thickness (6 to 1500 nm) exhibit no statistically significant difference compared to the wet PEDOT:PSS without the PU adhesive layer (P > 0.5) (fig. S10, B and C). Hence, the PU adhesive layer can provide strong adhesion of wet PEDOT:PSS (fig. S12) without compromising the mechanical properties of wet PEDOT:PSS.

**Mechanical and electrochemical stability of adhered wet conducting polymers**

To ensure long-term functionality and reliability of bioelectronic devices in wet physiological environment, mechanical and electrochemical stability of conducting polymers on substrates is one of the most critical requirements. We evaluate the mechanical stability of adhered wet conducting polymers by characterizing interfacial failures and electrical properties of solvent-casted wet PEDOT:PSS on ITO-glass electrodes with the PU adhesive layers under prolonged ultrasonication in PBS (Fig. 4) (23). The solvent-casted PEDOT:PSS on the ITO-glass electrode without the PU adhesive layer undergoes complete interfacial delamination after ultrasonication for 1 min in PBS (Fig. 4A). In contrast, the solvent-casted PEDOT:PSS on the amine-functionalized ITO-glass electrode with the PU adhesive layer remains intact (Fig. 4B) and shows negligible deterioration in electrical property (Fig. 4C) after ultrasonication for 10 min in PBS, demonstrating superior mechanical stability of the adhesion. Furthermore, a solvent-casted wet PEDOT:PSS on an amine-functionalized PDMS substrate with the PU adhesive layer can withstand and remain intact without interfacial failure after 10,000 cycles of bending (radius of curvature ~ 2 mm) in PBS (movie S1).

We further investigate the electrochemical stability of adhered wet conducting polymers by evaluating the electrochemical properties of solvent-casted PEDOT:PSS on amine-functionalized Pt electrodes with the PU adhesive layers based on cyclic voltammetry (CV) tests in PBS (Fig. 4D) (23, 32). The CV curves show a small change over 10,000 charging and discharging cycles with less than 6% decrease in charge storage capability (CSC) after 10,000 CV cycles, demonstrating superior electrochemical stability of the adhesion (Fig. 4, D and E). Moreover, the wet PEDOT:PSS on the Pt electrode with PU adhesive layer remains intact without any observable interfacial failure after 10,000 charging and discharging cycles (Fig. 4E).

The proposed method is also applicable to conducting polymers prepared by electrodeposition, which has been widely adopted to introduce conducting polymers on existing bioelectronic devices (22). To evaluate mechanical and electrochemical stability of electrodeposited conducting polymers adhered by the PU adhesive layer, we perform the same set of tests (i.e., ultrasonication, EIS, and CV) for electrodeposited PEDOT:PSS (fig. S13). Similar to solvent-casted PEDOT:PSS, the electrodeposited PEDOT:PSS on the ITO-glass electrodes with the PU adhesive layers exhibits superior mechanical stability against prolonged ultrasonication and electrochemical stability in wet physiological environment (fig. S13).
Strong adhesion of wet conducting polymers on bioelectronic devices

While electrodeposited PEDOT:PSS has been intensively studied to improve electrical properties of various bioelectrodes (e.g., low impedance, high charge injection capacity, and low rigidity), poor long-term stability of the electrodeposited PEDOT:PSS on bioelectrodes in wet physiological environment has been one major hurdle that significantly limits their utility in practice (17, 23, 24, 27). The new capability of forming strong, conductive, and mechanically and electrochemically stable adhesion between conducting polymers and diverse substrates allows us to readily electrodeposi conducting polymers on various existing bioelectronic devices, achieving high robustness in wet physiological conditions (Fig. 5).

Existing bioelectronic devices can be approximately classified into two categories based on configurations of their electrodes (33, 34). In the first category, bioelectronic devices such as electrocorticography (ECoG) arrays (35), Michigan probes (36), and multielectrode arrays (MEA) consist of planar microelectrodes on the device substrates to provide bioelectronic sensing or stimulation (Fig. 5A). In the second category, bioelectronic devices such as Utah arrays (37), microwire electrode arrays (38), and fiber probes (39) consist of tip microelectrodes at end tips of the devices (Fig. 5B). To demonstrate flexibility and ready applicability of the proposed method to robustly integrate conducting polymers with various bioelectronic devices in wet physiological environment, we choose commercially available MEA with Au microelectrodes (Fig. 5C) and polytetrafluoroethylene (PTFE)—coated Pt microwire electrodes (Fig. 5F) as representative examples for bioelectronic devices with planar microelectrodes and tip microelectrodes, respectively.

Because of weak and unstable adhesion between electrodeposited PEDOT:PSS and Au electrodes, the electrodeposited PEDOT:PSS on the MEA (electrode diameter, 100 μm) without the PU adhesive layer undergoes deterioration in electrical property (Fig. 5C) and nearly disappears from the electrode surface after ultrasonication for 5 min in PBS (Fig. 5D). In contrast, the electrodeposited PEDOT:PSS on the MEA with the PU adhesive layer exhibits no observable change in electrical property (Fig. 5C) and remains intact after ultrasonication for 60 min in PBS (Fig. 5E), demonstrating strong adhesion and superior stability of the adhesion. Similarly, the electrodeposited PEDOT:PSS on the Pt microwire electrode (wire diameter, 100 μm) without the PU adhesive layer experiences deterioration in electrical property (Fig. 5F) and shows mechanical damage with the bare Pt electrode exposed after ultrasonication for 5 min in PBS (Fig. 5G and fig. S14). In contrast, the electrodeposited PEDOT:PSS on the Pt microwire electrode with the PU adhesive layer (only formed on Pt surface, not PTFE coating) exhibits negligible deterioration in electrical property (Fig. 5F) and mechanical

Fig. 4. Adhesion stability of solvent-casted wet conducting polymers. (A) Images of a solvent-casted wet PEDOT:PSS on a ITO-glass substrate without the PU adhesive layer before and after ultrasonication for 1 min. Photo credit: Hyunwoo Yuk, MIT. (B) Images of a solvent-casted wet PEDOT:PSS on an amine-functionalized ITO-glass substrate with the PU adhesive layer before and after ultrasonication for 10 min. Photo credit: Hyunwoo Yuk, MIT. (C) EIS curves for a solvent-casted wet PEDOT:PSS on amine-functionalized ITO-glass substrates with the PU adhesive layer before and after ultrasonication for 10 min. (D) Long-term CV curves for a solvent-casted wet PEDOT:PSS on an amine-functionalized Pt electrode with the PU adhesive layer in PBS. (E) Measured CSC versus CV cycle number for a solvent-casted wet PEDOT:PSS on an amine-functionalized Pt electrode with the PU adhesive layer in PBS. Photo credit: Hyunwoo Yuk, MIT. Conducting polymers with 10 μm thickness and PU adhesive layers with 60 nm thickness were used for all experiments.

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integrity after ultrasonication for 30 min in PBS (Fig. 5H and fig. S15).

**CONCLUSION AND DISCUSSION**

In this study, we present a general yet simple method to achieve strong adhesion of various wet conducting polymers on diverse substrates through intermediate hydrophilic polymer adhesive layers. While we choose the hydrophilic PU as the adhesive layer in this study, other hydrophilic polymers such as poly(vinyl alcohol) (PVA) may also be adopted as the adhesive layer (fig. S16). The proposed method can provide robust interfacial integration of various wet conducting polymers such as PEDOT:PSS, PPy, and PAni on a wide range of insulating and conductive substrates including glass, PDMS, polyimide, ITO, and gold as well as bioelectronic devices with representative form factors. The resultant conducting polymers adhered on substrates achieve superior adhesion performance (shear strength over 120 kPa) and remarkable mechanical and electrochemical stability without compromising electrical or mechanical properties of the conducting polymers. Our method is compatible with various fabrication approaches for conducting polymers including solvent casting and electrodeposition and allows the use of commercially accessible off-the-shelf materials (e.g., hydrophilic PU and conducting polymers), offering ready and broad applications in bioelectronic devices. The method also gives strong adhesion for very thin spin-coated conducting polymers (100 nm

| Media/wet tissue | Plastic film | Microelectrode | Device | Frequency (Hz) |
|-----------------|--------------|---------------|--------|----------------|
| Media/wet tissue | Plastic film | Microelectrode | Device | Frequency (Hz) |
| Media/wet tissue | Plastic film | Microelectrode | Device | Frequency (Hz) |

![Image](image-url)
Formation of strong adhesion of conducting polymers
A hydrophilic PU (HydroMed D3, AdvanSource Biomaterial) and a PVA (molecular weight = 124,000; Sigma-Aldrich) were used as PU adhesive layer and PVA adhesive layer, respectively. To prepare a hydrophilic PU solution, varying concentrations [1 to 5 weight % (wt %)] of PU were dissolved in mixture of ethanol and deionized water (95:5 = ethanol:water, v/v). To prepare a PVA solution, 1 wt % of PVA was dissolved in deionized water at 90°C. For planar substrates and devices in this study, the adhesive layer was introduced in a controlled manner by spin coating of the hydrophilic PU solution or the PVA solution at 2000 rpm for 30 s. For nonplanar substrates and devices in this study, the adhesive layer was introduced in controlled manner by dip-coating of the hydrophilic PU solution or the PVA solution with pulling rate of 1 cm min⁻¹. After spin coating or dip coating, the PU- or PVA-coated substrates were annealed at 80°C for 1 hour. In experiments, PU adhesive layer with 60 nm thickness was used unless otherwise mentioned. For strong adhesion of wet conducting polymers, conducting polymers were prepared by either solvent casting or electrodeposition on the PU- or PVA-coated substrates following the previously reported protocols (10, 23, 40, 41).

For solvent-casted or spin-coated PEDOT:PSS, 15% (v/v) dimethyl sulfoxide was added into a commercially available aqueous PEDOT:PSS dispersion (Clevios PH1000, Heraeus Electric Materials) and vigorously stirred for 6 hours at room temperature. The mixed PEDOT:PSS dispersion was then filtered by PTFE filter (pore size, 5 μm) and solvent-casted or spin-coated on the PU- or PVA-coated substrates followed by air drying for 12 hours at room temperature and subsequent annealing for 30 min at 120°C. To obtain wet PEDOT:PSS adhered on the substrate, the samples were soaked in PBS for 1 hour before use.

For solvent-casted PPy, an aqueous pyrrole solution (0.3 M pyrrole and 0.1 M SDS in deionized water) and an aqueous oxidant solution (0.3 M ammonium persulfate in deionized water) were mixed in 1:1 by volume and vigorously stirred for 3 min at room temperature. The mixed solution was solvent-casted on the PU-coated substrates and incubated in ambient condition for 30 min. To obtain wet PPy adhered on the substrate, the samples were soaked in PBS for 1 hour before use.

For solvent-casted PAni, an aqueous aniline solution (1.42 M aniline and 1.42 M hydrochloric acid in deionized water) and an aqueous oxidant solution (1.42 M ammonium persulfate in deionized water) were mixed in 1:1 by volume and vigorously stirred for 1 min at 4°C. The mixed solution was solvent-casted on the PU-coated substrates and incubated in ambient condition for 30 min. To obtain wet PAni that adhered on the substrate, the samples were soaked in PBS for 1 hour before use.

For electrodeposited PEDOT:PSS, an EDOT:PSS precursor solution (0.02 M EDOT, 0.04 M PSS, and 0.1 M LiClO₄ in deionized water) was electropolymerized by an electrochemical potentiostat/galvanostat (PGSTAT30, Metrohm Autolab). A PU-coated electrode, a Pt sheet, and an Ag/AgCl wire were used as a working electrode, a counter electrode, and a reference electrode, respectively. The electropolimerization of EDOT was performed at 1.0 V versus Ag/AgCl (38 mC cm⁻²) for ITO-glass substrates and at 1.0 V versus Ag/AgCl (43 mC cm⁻²) for MEA with Au electrodes.

Measurement of adhesive layer thickness
The thickness of adhesive layers was measured by using an ellipsometer (Gaertner Scientific) with a wavelength at 633 nm. Changes in polarization were measured by the ellipsometer as a function of the sample thickness, which were then used to determine the thickness of the adhesive layers.

Elemental analysis and SEM imaging
Elemental distribution in adhesive interfaces was characterized by SEM-EDS measurement of a cross section of a dry PEDOT:PSS on a silicon substrate with the PU adhesive layer. The PEDOT:PSS on the amine-functionalized silicon substrate with the PU adhesive
layer was immersed in liquid nitrogen for 3 min and then cryo-fractured to obtain clean cross section. The cleaved sample was observed by using an SEM-EDS facility (6010LA, JEOL) and characterized to monitor the distribution of carbon and sulfur atoms. SEM images of bioelectronic devices with electrodeposited PEDOT:PSS were taken with the same SEM facility after Au sputtering (6 nm Au thickness).

**Atomic force microscopy imaging**

Atomic force microscopy (AFM) images and surface roughness data of diverse substrates were acquired by using an AFM facility (MFP-3D, Asylum Research). The substrates were directly attached onto the sample stage by double-sided carbon tape.

**XPS characterizations**

To prepare XPS samples, PEDOT:PSS was prepared by solvent-casting on a glass substrate, a glass substrate with the 60-nm PU adhesive layer, and a glass substrate with the 1500-nm PU adhesive layer. Before the sample preparation, the glass substrates were functionalized with perfluorosilane (1H, 1H, 2H, 2H-perfluorooctyltriethoxysilane) to aid the peel-off process of the samples from the glass substrates without residues of the PU adhesive layer or PEDOT:PSS. Subsequently, the PEDOT:PSS without and with the PU adhesive layer was peeled off from the glass substrate after swelling in PBS to expose the adhesion interface for XPS characterization. Surface chemical composition of the samples was characterized by using an XPS facility (VersaProbe II, Physical Electronics Inc.) under ultrahigh vacuum of 5 × 10⁻¹⁰ mbar.

**Mechanical characterizations**

Before mechanical characterizations, all samples were immersed in PBS for 1 hour to hydrate the conducting polymers. Interfacial shear strength was measured by lap-shear tests (ASTM D3163) with an overlap area (length, 20 mm; width, 25 mm). The lap-shear test samples were prepared by solvent-casting of conducting polymers. A hydrophilic nylon filter (pore size, 1 μm; Tisch Scientific) was covered on top of a conducting polymer precursor solution on the substrate to provide a robustly integrated backing. The lap-shear tests were performed by using a mechanical testing machine (2-kN load cell, Zwick/Roell Z2.5) at the crosshead speed of 50 mm min⁻¹. The interfacial shear strength was determined by dividing the peak measured force by the overlap area. Tensile tests were performed in PBS bath with dog-bone shape samples by using a mechanical tester (USTretch, CellScale) to avoid dehydration of the wet conducting polymers during tests. Ultrasonication of samples was conducted by using a sonicator (VWR) with PBS bath at 25°C.

**Electrical characterizations**

Electrical conductivity was measured by using a standard four-point probe (SCS-4200, Keithley). Sheet resistance was measured by using a digital multimeter (Fluke) with an applied potential bias of 100 mV. All conducting polymer samples used for electrical conductivity and sheet resistance measurements were fully hydrated by immersing in PBS for 1 hour before the tests. EIS measurements were performed by using a potentiostat/galvanostat (PGSTAT30, Metrohm Autolab) with a potential scan rate of 150 mV s⁻¹ in an electrochemical cell installed with a wet PEDOT:PSS on a Pt electrode with the PU adhesive layer as a working electrode, a Pt sheet as a counter electrode, a Ag/AgCl wire as a reference electrode, and PBS as an electrolyte. The frequency range between 0.1 and 100 kHz was scanned with an applied potential bias of 10 mV versus Ag/AgCl.

EIS measurements were performed by using an electrochemical potentiostat/galvanostat (PGSTAT30, Metrohm Autolab) with a potential scan rate of 150 mV s⁻¹ in an electrochemical cell installed with a wet PEDOT:PSS on a Pt electrode with the PU adhesive layer as a working electrode, a Pt sheet as a counter electrode, a Ag/AgCl wire as a reference electrode, and PBS as an electrolyte. The CSC of the sample was calculated from the measured CV data as

$$\text{CSC} = \int_{E_1}^{E_2} \frac{i(E)}{2na} dE$$

where \(v\) is the scan rate, \(E_2\) and \(E_1\) are the potential window, \(i\) is the current at each potential, and \(A\) is the area of the wet PEDOT:PSS on Pt electrode, respectively.

**Statistical analysis**

MATLAB software was used to assess the statistical significance of all comparison studies in this work. Sample size of 5 was selected otherwise mentioned in all experiments. Data distribution was assumed to be normal for all parametric tests but not formally tested. In the statistical analysis for comparison between multiple samples, one-way ANOVA followed by Tukey’s multiple comparison test were conducted with threshold of \(*P \leq 0.05, **P \leq 0.01, ***P \leq 0.001,\) and \(****P \leq 0.0001\). In the statistical analysis between two data groups, two-sample Student’s \(t\) test was used, and significance threshold was placed at \(*P \leq 0.05, **P \leq 0.01, ***P \leq 0.001,\) and \(****P \leq 0.0001\). All error bars in the graphs represent the 95% confidence interval (CI) values.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/12/eaay5394/DC1

**REFERENCES AND NOTES**

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