A stretchable epidermal sweat sensing platform with an integrated printed battery and electrochromic display

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Wearable electronic devices are of potential use in various health monitoring applications including non-invasive chemical sensing. However, such platforms are typically limited by the need to connect to external devices for power and data visualization. Here we report a stretchable epidermal sweat sensing platform that integrates a stretchable battery and a low-power digital electrochromic display. The patch can operate as a standalone device to directly display the concentration of various electrolytes or metabolites in sweat, such as glucose and lactate, without any wired or wireless connection to external devices. It consists of electrochemical sensors, a stretchable Ag2O–Zn battery, ten individually addressable electrochromic pixels and a small microcontroller unit. All the components and interconnections, except the microcontroller, are fabricated via the high-throughput screen printing of customized elastomeric or silver inks. The integrated system is robust to mechanical deformation and is unaffected by 1,500 stretching cycles at 20% strain. The electrochromic display exhibits stability for 10,000 on/off cycles, and the battery can power 14,000 sensing sessions over a week-long use.

Soft electronics are of use in on-body sensing and human–machine interfacing9–11. Integrated epidermal sensing systems have, for instance, been developed as ‘labs on the skin’ for recording various mechanical, electrical, physiological and electrochemical signals9–11. In particular, epidermal sweat sensing with electrochemical sensors can be used to detect electrolytes, metabolites, nutrients and drugs, providing non-invasive health monitoring for precision medicine, personalized nutrition and sports performance9–11. The development of wearable sensors for such applications is increasingly moving away from separate physical and chemical sensors, and towards the integration of sensors with electronics for energy management, signal acquisition and data interfacing9–11. However, most wearable electronics currently operate with commercial lithium polymer pouches or coin cells, which are rigid, unsafe and add bottlenecks to the product design. Conformal epidermal sensors with wired connections or short-range wireless power delivery have been designed to avoid battery-related limitations, but these can compromise the autonomy of the system and limit user mobility9–11.

Integrated sensors also typically rely on wireless data transmissions, which require external devices (such as computers, smartphones and receivers) for users to obtain the sensing results9,11,14–16. A lack of direct access to sensing results means that many existing wearable sensors can be inconvenient in practical applications. Skin-conformal flexible electronic systems that include visual data presentation could...
address these limitations, simplify designs and improve user experience. There are, however, major barriers to integrate accurate on-body sensing and real-time display capabilities within a compact, flexible and wearable monitoring platform.

Conventional display technologies can be modified to have flexible form factors for wearable electronics, but their high power consumption limits application in compact low-energy electronic systems. Alternatively, non-emissive displays, such as liquid crystal and electronic ink displays controlled by a microcontroller, consume low power and can effectively present data. However, their lack of flexibility and stretchability limits integration with wearable devices. Nevertheless, integrated sensing systems with miniaturized display elements have been developed for visualizing data in real time for direct readout. These include colorimetric assays that directly react with the target analytes (such as glucose or chloride ions) and qualitatively report the results via the naked eye. Electrochromic materials that change their colour or opacity by applying an electrical potential have also been integrated with sensing systems for data visualization. However, colorimetric display technologies have long reaction times, and naked-eye readout lacks spatial and colorimetric resolution, resulting in inaccurate and unclear data presentation. Further analysis is usually required to convert qualitative colorimetric information into numerical data using smartphones, spectral analysers or cameras.

In this Article, we report an epidermal sensing platform with an integrated electrochromic display (ECD) for the real-time visualization of analytical data and a high-performance stretchable battery for power supply. The epidermal patch consists of stretchable enzymatic and potentiometric electrochemical sensors for sensing various metabolites and electrolytes in sweat, a stretchable AgₐO–Zn battery, ten fast responding and reversible poly(3,4-ethylene dioxythiophene) polystyrene sulfonate (PEDOT:PSS)-based electrochromic pixels and a low-power microcontroller unit (MCU) chip (Supplementary Video 1). All the sensors, batteries and ECDs are fabricated via the high-throughput screen printing of customized stretchable composite inks on a heat-sealable thermoelectric styrene–ethylene–butylene–styrene (SEBS) block co-polymer substrate (Fig. 1a and Supplementary Fig. 1). Electrical connections between the components of the system and with the MCU are printed using stretchable silver ink layers treated with a chloride/lactate acid treatment, which exhibit stable electrical connections under repeated stretching (Supplementary Fig. 2).

The system is solely powered by the integrated battery, which is activated by a switch that connects the batteries to the MCU. The signal input from the electrochemical sensor is then sampled by the MCU via an analogue-to-digital converter (ADC), which converts the analytical data into digital outputs that control the on and off states of the ten ECD pixels (Fig. 1b). As the sensor relies on potentiometric reading when in contact with the analyte and independently operates from the MCU, the sensing and data visualization occur in under one second once the switch is pressed. We illustrate the capabilities of the integrated system using two potentiometric sweat sensors for monitoring sodium ions and pH, and two enzymatic amperometric sweat sensors for monitoring glucose and lactate, with the analyte concentrations displayed on the ECD (Fig. 1c). The epidermal patch exhibits excellent durability and mechanical robustness (Fig. 1d(i), (ii)), demonstrating 10,000 sensing sessions over five days and 1,500 stretching cycles at 20% strain without affecting its sensing or display capability. A rapid room-temperature ‘solvent-welding’ process was developed to bond the rigid chip and soft printed circuit, which have very different elastic moduli; consequently, the MCU can be securely connected without using conductive adhesives (Fig. 1d(iii) and Supplementary Fig. 3). The use of maleic-anhydride-grafted SEBS (SEBS-MA) further strengthens the bonding of the MCU metal contact pads to the substrate, allowing up to 300% strain without detachment (Supplementary Fig. 4).

**Stretchable PEDOT:PSS ECDs**

Integrating ECDs into skin-interfaced electronic systems provides timely data visualization, such as the level of a target biomarker in the case of on-body sensing platforms. Our skin-interfaced integrated sensing platform features a dedicated and highly responsive low-power ECD for the direct visualization of analytical data. The ECD was chosen for its low power consumption compared with light-emitting display technologies, as it features low self-discharge and only requires energy as the displaying content is refreshed. The electrochromic PEDOT:PSS electrode material was selected for its stretchability and compatibility with screen-printing processes. Fabricated via layer-by-layer screen printing, the ECD was composed of a top PEDOT:PSS panel and ten separately addressable pixels on the bottom panel, with a highly viscous polystyrene sulfonate (PSS)-based electrolyte used to create ionic pathways between the two panels as the opposing electrodes are physically separated; stretchable and printed silver traces were used to connect the ECD to the MCU and were covered with an insulating elastomer layer to avoid short-circuiting and corrosion (Fig. 2a and Supplementary Fig. 1). The colour of the PEDOT:PSS conducting polymer is dependent on its redox state, as described in the half-reaction below.

\[
\text{(PEDOT)}^+ (\text{PSS})^- (\text{light blue}) + e^- \rightarrow \text{PEDOT} (\text{dark blue}) + \text{PSS}^- 
\]

The colour change, thus, occurs on the front panel during the reductive PSS-undoping process on applying a potential above 1 V between the two electrodes, during which the reverse oxidative doping process takes place on the back-panel PEDOT:PSS electrodes. Thus, to induce a colour change, a positive potential can be applied to the bottom pixels, which results in the colour-changing reduction of PEDOT:PSS within the corresponding region on the front panel covered by the block of PSS electrolyte (Fig. 2b and Supplementary Fig. 5). The printable PEDOT:PSS ink was optimized for stretchability and compatibility with the elastic SEBS substrate and shows excellent mechanical behaviour and electrochemical stability on repeated 20% stretching deformation (Fig. 2c and Supplementary Fig. 6). Cyclic voltammetry of the electrochromic pixel was carried out with scan rates of 0.1, 0.5 and 1.0 V s⁻¹ over the 0–1.5 V range (Fig. 2d). The on/off switching of the ECD was characterized using potential stepping between 0 V and a set switching voltage as the current was monitored. As shown in Fig. 2e, Supplementary Figs. 6 and 7 and Supplementary Video 3, a switching voltage of 1.5 V was determined to be optimal as it results in the fastest turn-on response with a turn-off response below 500 ms. In comparison, the 1 V switching voltage resulted in a slower turn-on and turn-off speed, whereas switching voltages of 2 and 3 V, as a fast turn-on rate is given, led to a slower turn-off speed due to excessive charge accumulation on the electrodes. Such a high voltage can also lead to electrolyte breakdown as well as irreversible electrode overoxidation that limits the cycle life of the electrochromic cells and should be avoided. The average power consumption of turning on each pixel, on applying the optimal 1.5 V for 1 s, was determined to be 0.08 mW (Supplementary Fig. 8). This behaviour was examined again after 1,500 cycles of repeated 20% uniaxial stretching along two orthogonal directions. The results (Fig. 2f, Supplementary Fig. 9 and Supplementary Video 2) demonstrate a slightly reduced turn-on and turn-off current but a negligible change in switching speed, indicating the excellent stretchability of the PEDOT:PSS electrodes. As characterized via electrochemical impedance spectroscopy (Fig. 2g), the reduction in switching current can be attributed to a slight increase (≈200 Ω) in the electrode serial resistance, whereas the charge transfer and diffusion impedance remained mostly constant. The electrochemical stability of the ECD was tested via repeated cycling of the PEDOT:PSS electrodes between on and off (Fig. 2h(i)) over 10,000 cycles (Fig. 2h(ii)) via potential steps between 0 and 1.5 V. The electrical current response of the pixel shows high reproducibility with only 17% decrease in the peak current after...
about 10,000 on/off cycles (Fig. 2h(iii),(iv) and Supplementary Fig. 10), which can be caused by the increased cell impedance due to electrode overoxidation from repeated cycling (Supplementary Fig. 11). Such high mechanical and electrochemical stabilities ensure the continued and reliable operation of the ECD within the designed lifetime of the system.

Battery performance

The all-printed fabrication of the flexible Ag$_2$O–Zn battery was designed based on the selection of substrate and ink formulation that ensures mechanical stretchability, electrochemical stability, chemical resilience, operational safety, light weight and comfort. SEBS was selected as the binder for electrodes, separator and substrate due to its chemical stability in the alkaline electrolyte used in the Ag$_2$O–Zn battery. Furthermore, thermoelectric seric SEBS is compatible with both heat and vacuum sealing, which grants reliable packaging to protect and contain the electrolyte. Polyvinyl alcohol (PVA) hydrogel electrolyte infused with 9 M potassium hydroxide was also formulated to physically contain the electrolyte and avoid leakage. A stacking configuration was designed to minimize the device footprint as the ion diffusion distance between the battery electrodes is reduced, being minimized the internal resistance. The batteries, with the customized silver current collector, zinc anode, Ag$_2$O cathode and titanium dioxide (TiO$_2$) separator inks, were printed in a layer-by-layer manner on the top and bottom panels and assembled via heat and vacuum sealing on placing the electrolyte hydrogel (Fig. 3a). The redox reaction of the resulting Ag$_2$O–Zn battery relies on the dissolution of zinc ions (Zn$^{2+}$) and silver ions (Ag$^+$) in the alkaline electrolyte and their supersaturation-induced precipitation, which takes place rapidly as a stable voltage is maintained at 1.56 V (Fig. 3b). The reactions are described below ($E$, electrode potential; SHE, standard hydrogen electrode).

Anode : $\text{Zn} + 2\text{OH}^- \leftrightarrow \text{ZnO} + \text{H}_2\text{O} + 2e^- E = -1.22 \text{ V versus SHE}$ (2)

Cathode : $\text{Ag}_2\text{O} + \text{H}_2\text{O} + 2e^- = 2\text{Ag} + 2\text{OH}^- E = +0.34 \text{ V versus SHE}$ (3)
The capacity of the printed stretchable battery can be adjusted by the loading of cathode material, as controlled by the number of layers of printing. As shown in Fig. 3c, batteries with different (1–4) layers of printed cathode were prepared and tested at the discharge rate of 1 mA cm\(^{-2}\). Such a thickness change results in a proportional increase in capacity with the number of layers, reaching as high as 11.5 mAh cm\(^{-2}\) with four layers of cathode loading. The power consumption of the MCU was separately characterized at various operation potentials, and the current draw at the voltage of 1.5 V was determined to be at roughly 1 mA (Supplementary Fig. 13). The operation of the system was tested with 1 s fluctuations for the 14–22 cycles (iii) and 9,986–9,994 cycles (iv).

### Stretchable display system with potentiometric sensors

Two types of potentiometric sensors, namely, for monitoring pH and sodium, were employed for demonstrating the applicability of the all-in-one sensing device. Both sensors rely on measuring the potential difference between the modified working electrode and silver/silver chloride (Ag/AgCl) reference electrode when in contact with electrolyte and packaging materials. By printing the stretchable silver ink and carbon ink on the SEBS substrate, the two-electrode sensor shows satisfactory resiliency to 20% stretching (Fig. 4b(i)(ii)). The electrode–electrolyte interface of the sodium-ion-selective-membrane-modified sodium sensor results in a logarithmic dependence on the sodium concentration (Fig. 4c(i)). Figure 4c(ii) depicts the calibration of the fabricated sodium sensor, demonstrating a slope of 62.4 mV per decade of sodium concentration, with a high correlation coefficient (\(R^2 = 0.9996\)), over the 0.1–100.0 mM range. The high selectivity of the sodium sensor is demonstrated by the negligible change in sensor potential response to a wide range of clinically or environmentally important electrolytes.

The shelf life of the battery was tested over 1,000 h, displaying a negligible change in voltage and internal resistance of \(<40 \Omega\) (Supplementary Fig. 15), reflecting the high chemical stability of the electrodes, electrolyte and packaging materials.
in the presence of potassium and different pH values (Supplementary Fig. 16). As shown in Fig. 4c(iii), the potential response of the sensor in alternating 0.1 and 10.0 mM sodium solutions is highly reversible and exhibits minimal changes after 500, 1,000 and 1,500 stretching cycles, reflecting the excellent stretchability of the sodium sensor.

The pH sensor is fabricated by the electropolymerization of polyaniline onto the working carbon electrode. The operation of the resulting pH sensor is based on the protonation and de-protonation of nitrogen atoms in the polymer chains of polyaniline (Fig. 4d(i)). The sensitivity of the fabricated pH sensor is evaluated by measuring the potential response at pH values in the range of 3–8 (Fig. 4d(i)). The sensors display a linear response of 62.62 mV pH⁻¹ with a high correlation coefficient ($R^2 = 0.9995$) across the entire pH range (Fig. 4d(ii)). As illustrated in Fig. 4d(iii), the modified pH sensor displays excellent stretchability, as evidenced by the highly stable potential signals measured at pH 4 and pH 7, with negligible changes after 500, 1,000 and 1,500 stretching cycles. In addition, the pH sensor demonstrates good selectivity in the presence of Na⁺ and K⁺ (Supplementary Fig. 16). The operating voltage obtained from sodium and pH sensors is converted into logic outputs, programming the MCU and correlating the obtained voltage values to the display content (Fig. 4e and Supplementary Tables 1 and 2). By integrating the pH sensor with the ECD, we, thus, fabricated an all-in-one epidermal patch for rapid pH sensing and display (Fig. 4f(i)(ii)).

**Stretchable display system with enzymatic sensors**

Two enzymatic sensors for glucose and lactate were selected to demonstrate the applicability of the integrated ECD patch in a real-life scenario.
for epidermal sweat sensing of both metabolites. Both enzymatic sensors commonly rely on amperometric measurements, where the electrical current at a set potential can be correlated to the metabolite concentration using a sigmoidal fitting:

$$E = E_0 + \frac{E_0 - a}{1 - \left(\frac{\text{Conc.}}{b}\right)^c}, \quad (4)$$

where $E$ is the sensor voltage, $E_0$ is the open-circuit potential of the sensor without load, Conc. is the analyte concentration and $a$, $b$ and $c$ are constants. To enable rapid and low-power-consumption sensing as the need for potentiostats and amplifiers is obviated and removing them from the circuits, a self-powered sensing mechanism that converts the amperometric signal to a voltage signal was implemented under an optimized load. The sensor electrodes are composed of a printed carbon working electrode along with $\text{Ag}_2\text{O}$ counter electrode, where the carbon electrodes were thereafter functionalized with the corresponding enzymes for their target analytes (Fig. 5a). Figure 5b displays the lengthwise stretching of the enzymatic sensor fabricated using the customized stretchable inks. As shown in Fig. 5c, the epidermal patch perfectly matches the skin curvature of the subject’s forearm. After sweat is induced from the loaded exercise, the subject can mount the integrated system on the forearm for demonstrating its application in real conditions. On pressing the switch for 1 s, the sensor was, thus, able to display the lactate concentration on the wearer’s skin without needing any external devices, allowing the user to directly obtain the sensing results without the need of any stabilization time or delay.

The lactate sensor operates based on the selective 1,4-naphthoquinone (NQ)-mediated oxidation of lactate on the working electrode by the lactate oxidase (LOx) enzyme, immobilized on carbon nanotubes. The working electrode is coupled with the $\text{Ag}_2\text{O}$ counter electrode, where the carbon electrodes were thereafter functionalized with the corresponding enzymes for their target analytes (Fig. 5a). Figure 5b displays the lengthwise stretching of the enzymatic sensor fabricated using the customized stretchable inks. As shown in Fig. 5c, the epidermal patch perfectly matches the skin curvature of the subject’s forearm. After sweat is induced from the loaded exercise, the subject can mount the integrated system on the forearm for demonstrating its application in real conditions. On pressing the switch for 1 s, the sensor was, thus, able to display the lactate concentration on the wearer’s skin without needing any external devices, allowing the user to directly obtain the sensing results without the need of any stabilization time or delay.

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counter electrode that undergoes a high reaction-rate reduction of Ag2O to Ag, independent of the analyte concentration. On applying a load, a continuous current flows between the enzyme electrode and Ag2O electrode, which is dominated by the concentration-limited electrocatalytic reaction on the LOx electrode (Fig. 5d(i)). The optimization of the load ensures a maximized signal resolution over the desired concentration region. To this end, linear sweep voltammetry (LSV) was performed at typical sweat lactate concentrations \(^{34-36}\), namely, 0 and 25 mM, and the potential was plotted against the load (Fig. 5d(ii)). As shown, a load of 75 kΩ was selected, where the potential difference between 0 and 25 mM lactate in artificial sweat (AS) was at the maximum. Subsequently, the in vitro voltage response of the lactate sensor at different concentrations was measured under the 75 kΩ load and fitted with a sigmoidal model (Fig. 5d(iii) and Supplementary Fig. 18d)\(^{34-36}\). To further illustrate the importance of optimizing the load to the range and resolution of the lactate sensor, the sensor was also tested under different loads of 10 and 500 kΩ (Supplementary Fig. 17). This test showed a minimal potential change above the lactate concentration of 10 mM under the 500 kΩ load, and poor voltage resolution across 0–30 mM lactate concentrations under the 10 kΩ load. Figure 5c(iv) examines the stretchability of the lactate sensor, demonstrating highly stable and reversible voltage signal after 1,500 cycles of lengthwise uniaxial 20% stretching. Additional characterization shows that the lactate sensor exhibits effective discrimination against common sweat constituents, including urea, ascorbate, acetaminophen and glucose, as well as high electrochemical stability, operating continuously over 9 hours (Supplementary Fig. 18b). Considering the potentially changing pH of human sweat, the fabricated lactate sensor was tested in different AS scenarios with variable pH values of 5.5, 6.5 and 7.4 (Supplementary Fig. 19). No noticable differences in potentiometric response were observed among the tested samples, demonstrating the reliability and practicality of the sensor. Based on the results obtained from the lactate sensors, the MCU can be programmed to convert the voltage response of the sensors to the 10 pixel digital display within the desired range (Fig. 5e(i) and Supplementary Tables 3 and 4). The on-body application of the epidermal device was demonstrated by integrating the all-in-one lactate sensor system. A sweat-compatible adhesive layer with a simple fluidic design was attached to the back panel of the patch for directing the sweat flow through the sensor and to establish temporal resolution (Supplementary Figs. 23 and 24). The experimental sweat flow data is supported by the theoretical simulation (Supplementary Fig. 24c). Validated using commercial lactate meters, the sensor, thus, shows the ability to track the temporal profile of lactate level in sweat over a given period of time. The observed lactate level changed from the initial 6 mM at the beginning of perspiration to 9 mM after 5 min, reflecting changes in the metabolic activity of the sweat gland (Fig. 5e(ii)(iii)).

Using a similar two-electrode construction, the NQ-mediated glucose oxidase was used to obtain a highly selective glucose-sensing working electrode in connection to the Ag2O counter electrode (Fig. 5f(i)). The load was similarly optimized via LSV at 75 kΩ and using 0 and 20 mM glucose solutions (Fig. 5f(ii)). The well-defined potential response of the resulting glucose sensor over the 0–10 mM range is presented in Fig. 5f(iii), and its high selectivity against common sweat constituents, including uric acid, ascorbic acid, acetaminophen and lactate, is displayed in Supplementary Fig. 20. The stretchability of the sensor was assessed by stretching the electrode as its potential response to 2.5 and 10.0 mM glucose solutions is recorded. As illustrated in Fig. 5f(iv), the glucose response remained highly stable after 1,500 cycles of lengthwise uniaxial 20% stretching. Similar to the lactate sensor, the glucose sensor was tested in AS under variable pH values, demonstrating a stable voltage signal regardless of the environment, and confirming its practicality under real-life operating conditions (Supplementary Fig. 21). It is worth noting that such current-to-voltage signal conversion can be tailored for accommodating the lower glucose concentration in human sweat. This is accomplished by adjusting the load to a higher value, which enhances the signal resolution in lower concentrations in exchange for a wider range. To this end, an optimized load at 250 kΩ was applied to enhance the voltage difference for glucose concentrations over the 0–5 mM range (Fig. 5f(v)(vi)). Similarly, the load was optimized to 10 MΩ to enhance the difference below 1 mM glucose towards the monitoring of sweat glucose, which is typically below 0.5 mM (Fig. 5f(vii)(viii))\(^{44}\). Similarly, the patch with a glucose sensor can be placed on the body to trace the glucose changes over a period of time. Based on the results obtained from the glucose sensors in the 0–0.5 mM range, the MCU was programmed to convert the voltage response of the sensors to the 10 pixel digital display (Fig. 5g(ii) and Supplementary Table 6). The glucose level increased after the intake of sugary drinks during exercise, with the trend agreeing with the changes in blood glucose level (Fig. 5g(iii)(iii)).

Conclusions
We have reported an all-printed epidermal monitoring system with a compact, flexible and stretchable form factor for non-invasive sweat monitoring. The integrated system combines sensors, ECDs and batteries and allows users to obtain accurate electrochemical sensing data and directly and rapidly display it during normal activities, without needing to connect to external devices. Stable and robust elastomers were used as the substrate and binder, which allows the components to be fabricated via the low-cost, high-throughput layer-by-layer stencil printing. The ECDs exhibit colour reversibility for over 10,000 on/off sensing cycles and 1,500 stretching cycles at 20% strain, and the battery could power 14,000 discrete sensing sessions. To illustrate the range of sensing options, potentiometric pH and sodium electrolyte sensors as well as enzymatic glucose and lactate metabolic biosensors were shown to be compatible with the integrated sensing platform. The fully autonomous multifunctional self-sustainable wearable sweat sensing platform could be of use in a diverse range of practical applications in personal wellness management, healthcare monitoring and professional sports. Further development of skin-worn monitoring-display platforms and power sources, such as via flexible microgrids\(^{45}\), could offer full autonomy and improved user experience for next-generation wearable monitoring platforms.
Graphite, toluene, acetonitrile, ethanol, HCl, NaCl, glutaraldehyde, acetic acid, NaFMN, lactic acid, d(+)-glucose, glucose oxidase, Ag flake, AgNO3 powder, Zn, ZnO, bis-muth trioxide (Bi2O3), TiO2, sodium ionophore X, sodium tetrakis[3,5-bis(trifluoromethyl)phenyl]borate, bis(2-ethylhexyl) sebacate, tetrahydrofuran (THF), polyvinyl butyral, poly(vinyl chloride), NaCl, SEBS-MA, methanol, iron (tit) chloride, aniline, sodium phosphate anhydrous, citric acid, sodium PSS, glyceral, b-sorbitol, sodium-N,N′-methylenebisacrylamide, potassium peroxydisulfate, sodium dodecylbenzene sulfonate and bromoresol green were purchased from Sigma-Aldrich. Polyurethane (Techniflex EG-80A) was obtained from Lubrizol. SEBS G1645 trilobed co-polymer was obtained from Kraton. Carbonyl-functionalized multiwalled carbon nanotube (MWNT-COOH) was purchased from Cheap Tubes. The screen-printable PEDOT:PSS paste was purchased from Sun Chemical. LOx was purchased from Toyobo. Capstone fluorosurfactant FS-65 was obtained from DuPont. Poly(vinylfluoride-co-2,3,3-tetrafluoropropylene) binder (GBR6005) was obtained from Daikin.

**Fabrication of stretchable circuit**

A stretchable silver ink was used to fabricate the circuit that connects the various components of the device. The circuit pattern was designed with AutoCAD 2020 (Autodesk) and made into a cut-through metal stencil by Metal Etch Services. The stretchable silver ink was composed of silver flake, SEBS (40 wt% in toluene) and toluene in a 4:2:1 weight ratio.

After the printing of the silver ink layers, a chlorin/lactic acid treatment, adapted from a previous study39, was employed to further functionalize the silver particles and enhance the stability and conductivity of the silver layer during stretching. The treatment involved preparing an aqueous solution of 100 mM NaCl and 50 mM lactic acid, which was repeatedly sprayed onto the printed pattern using a spray bottle and dried in the oven at 80 °C for 5 min for three times, followed by washing with deionized (DI) water and drying in the oven at the same temperature for 10 min. The treatment endowed the silver layer with higher conductivity and stability (Supplementary Fig. 2). The subsequent PEDOT:PSS, Zn, Ag, O and insulator can be printed on the treated silver layer.

**Fabrication of ECD**

The substrate for the ECD was composed of SEBS dissolved in toluene (40 wt%) and was fabricated by doctor blade casting (1,000 μm thick) on the SEBS, followed by curing in the oven at 80 °C for 1 hour.

The ECD was fabricated using layer-by-layer screen printing with customized four kinds of ink: the electrochromic PEDOT:PSS:ink, the silver ink for interconnection, the opaque insulator ink composed of SEBS and TiO2, and the sodium-PSS-based electrolyte ink. The PEDOT:PSS ink was prepared using 1 g PEDOT:PSS paste, 0.2000 ml toluene, 0.1500 ml sodium dodecylbenzene sulfonate (75 mg ml−1 in DI water) and 0.0135 ml Capstone fluorosurfactant FS-65. The use of toluene enables better bonding between PEDOT:PSS and the substrate, and the addition of sodium dodecylbenzene sulfonate and Capstone surfactant was used to improve ink wettability to avoid contraction of the printed patterns due to surface tension of the ink. Furthermore, the addition of the surfactant can further improve PEDOT:PSS conductivity due to enhanced phase separation46. The opaque insulator ink was composed of TiO2 and SEBS (40 wt% in toluene) in 1:10 weight ratio. The PEDOT:PSS:ink was formulated by mixing sodium PSS, b-sorbitol, glyceral, TiO2, and polyacrylamide (PAM) precursor solution in a 4.0:1.0:1.0:0.8:2.0 weight ratio. The PAM solution was formulated by mixing acrylamide, DI water, potassium peroxydisulfate and N,N′-methylenebisacrylamide in a 2.00:10.00:0.05:0.02 ratio. Here TiO2 was used to form a white, opaque electrolyte layer to enhance the color-change contrast, whereas the PAM solution was used to enhance the electrolyte viscosity to avoid the bleeding and short-circuiting of the printed PSS electrolyte ink among the pixels (Supplementary Fig. 12). All the inks are mixed in the planetary mixer at 1,800 rpm for 10 min or until homogeneous.

The ECD panel was composed of the colour-changing top panel and the bottom panel to control the regional colour change. The PEDOT:PSS ink was printed first onto the SEBS substrate and cured in the oven at 100 °C for 2 hours. Then, the silver interconnection and opaque insulator layers were printed, with each layer cured at 80 °C for 5 min. Before assembly, the electrolyte was printed onto the bottom panel and briefly heated in the oven for 15 s at 80 °C to crosslink the PAM in the electrolyte.

**Fabrication of stretchable battery**

The formulation of the battery inks and electrolyte was adapted from previous studies23,24. The printing of the battery relied on four inks: Ag ink for the current collectors, Zn ink for the battery anode, Ag2O ink for the battery cathode and separator ink for the battery separator. The Zn ink is prepared using Zn, ZnO, Bi2O3 and GBR6005 binder (25 wt% in acetonitrile) in a 9.00:0.50:0.50:0.267 weight ratio. The Ag2O ink was prepared using Ag2O, Super P carbon black and GBR6005 binder (33 wt% in acetonitrile) in a 1.90:0.10:2.66 weight ratio. The separator ink was composed of TiO2, Triton X-114, SEBS (40 wt% in toluene) and toluene in a 3.0:0.1:1.5:1.5 weight ratio. All the inks were mixed in the planetary mixer at 2,500 rpm for 10 min or until homogeneous. The PVA electrolyte gel was initially crosslinked by dissolving 0.9 g PVA into 9.1 ml DI water at 90 °C, followed by mixing with 14.0 g of 16.7 wt% NaOH solution and desiccating the mixture in a vacuum desiccator until remaining 67% of the total weight. The formed hydrogel was then cut using a three-dimensionally printed hole punch into the same shape as the electrodes and soaked in 9 M NaOH solution for over 12 hours before use. Here NaOH was used instead of potassium hydroxide to reduce self-discharge and extend the shelf life of the cell.

The stretchable battery was composed of two pairs of Ag2O–Zn batteries separately printed on the bottom and top panels. The layer-by-layer printing steps of the battery were followed by using Ag ink, Zn anode or Ag2O cathode ink, and separator ink, with each layer cured at 80 °C for 5 min (Supplementary Fig. 1). Before assembly, a PVA electrolyte gel was placed between the anode and cathode, and a concentric circle switch was placed between the Ag interconnection of two panels. After aligning and combining the two panels, the battery can effectively work and power the MCU by pressing the switch.

**Fabrication of potentiometric sensors**

The sodium and pH sensors were fabricated by using flexible silver and carbon inks. The formulation of the Ag ink is described above. The carbon ink was prepared using graphite, Super P carbon black, SEBS (40 wt% in toluene) and toluene in a 6.0:1.0:8.4:2.1 weight ratio. Both inks were mixed in a planetary mixer at 1,800 rpm for 10 min or until homogeneous.

The silver ink and carbon ink were printed onto the top-panel SEBS substrate in a layer-by-layer manner and were covered using SEBS resin to define the electrode area, exposing 2 mm2 of carbon electrode as the working electrode and 1 mm2 of the silver electrode as the reference electrode. A 0.1 M iron (III) chloride solution was drop cast onto the silver electrode to chlorinate the surface and form AgCl.

**Fabrication of sodium sensor**

The sodium sensor was based on screen-printed electrodes modified via drop casting. A cocktail composed of polyvinyl butyral (78.1 mg ml−1) and NaCl (50.0 mg ml−1) dissolved in methanol was drop cast onto the chlorinated reference surface (1.5 μl mm−2). A polyurethane resin (1 g in 20 g THF) was then drop cast onto the dried cocktail layer (2 μl mm−2) to prevent leaching. A cocktail for the Na+-selective electrode was formulated by dissolving 1.00 mg sodium ionophore X, 0.77 mg sodium tetraakis[3,5-bis(trifluoromethyl)phenyl]borate ion exchanger, 33.00 mg poly(vinyl chloride) and 66.00 mg bis(2-ethylhexyl) sebacate in 660 ml
nitrogen-purged THF. The resulting cocktail was subsequently drop cast onto the carbon electrode (2 µl mm⁻²).

Fabrication of pH sensor
Following the screen-printing fabrication protocol, the working and reference electrodes were created. The pH sensor was prepared by the electropolymerization of aniline onto the working carbon layer. The carbon electrode was first electrochemically cleaned via ten cyclic voltammetric scans in 0.5 M HCl (40 µl) from 0.3 to 1.1 V versus Ag/AgCl at a scan rate of 0.1 V s⁻¹. Subsequently, polyaniline electropolymerization was performed with a solution containing 0.1 M aniline in 1.0 M HCl by cyclic voltammetry from −0.2 to 1.0 V (versus Ag/AgCl) at 0.05 V s⁻¹ to an accumulative gain of around 100 mC charge on the carbon electrode. Subsequently, the electrode was washed with DI water and was allowed to air dry.

Fabrication of enzymatic sensors
The glucose sensor and lactate sensor were fabricated by using flexible silver ink for the connection, carbon ink for the enzymatic anode and Ag₂O ink for the sensor cathode. The Ag₂O ink is prepared by mixing Ag₂O, Super P carbon black and GBR binder (21 wt% in acetone) of the electrode was covered using SEBS resin to define the electrode area, exposing 2 mm² of carbon electrode as the anode and 2 mm² of the Ag₂O electrode as the cathode.

Modification of glucose sensor
The glucose sensor was fabricated by drop casting 0.2 M NQ solution dissolved in an acetone:ethanol (9:1) mixture containing 2 mg ml⁻¹ MWCNT-COOH onto the carbon electrode (1.5 µl mm⁻²). Then, 40 mg ml⁻¹ glucose oxidase solution dissolved in phosphate-buffered saline (PBS) containing 10 mg ml⁻¹ bovine serum albumin was drop cast after the first layer was dried (1.5 µl mm⁻²). Finally, a 1.50 µl mm⁻² chitosan layer (1 wt% in 0.1 M acetic acid) and a 0.75 µl mm⁻² glutaraldehyde layer (1% in water) were drop cast on the top for improved immobilization. A surface-mount resistor (75 kΩ, 150 kΩ or 10 MΩ) was used to temporarily soften the interconnecting silver traces followed by placing the MCU. The sensor was allowed to evaporate at room temperature for 5 min to facilitate the physical crosslinking of the SEBS within the Ag ink and the substrate to the metal contact pads of the MCU chip. This rapid solvent-welding process, thus, eliminates the use of any bonding layer or conductive paste. The connected MCU was sealed by 0.1 ml SEBS (40 wt% in toluene) and cured in the oven at 80 °C for 30 min after drying at room temperature. Two PVA gels were placed onto the Zn anode and Ag₂O cathode of the bottom panel as the electrolyte of the battery. The top panel was then aligned and covered with the bottom panel. Last, the assembled device was heat sealed on all the four sides to finish the device assembly.

On-body sensing with the patch
A stretchable adhesive (3M Adhesive Transfer Tape 468MP) was used to adhere the patch to the skin surface. A simple fluidic pattern was designed, allowing the adhesive layer to guide the sweat through the sensor and avoid sweat accumulation below the patch. The adhesive was initially cut using a computer-controlled cutter (Explore Air 2, Cricut) and attached to the back panel of the patch. The patch can thereafter be attached to the subject’s arm during on-body trials. During the trial, the subject was asked to ride a stationary bike until sweating before applying the patch. To perform one sensing session, the subject can simply press the switch mechanism for ~1 s to refresh the display. To study the accuracy and temporal profile of sweat lactate, the sensing session was performed every 5 min. The sensor output was sent to the corresponding author upon reasonable request.

Data availability
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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**Author contributions**
L.Y., K.N.K. and J.W. conceived the idea for this work. L.Y., M.C., K.N.K., A.T. and J.W. designed the experiments and wrote the manuscript. L.Y., M.C., J.-M.M., J.R.S., C.W., A.T., F.Z. and H.H. conducted the experiments. L.Y., M.C., J.Y. and R.L. fabricated the samples. M.L. designed and programmed the electronics. J.R.M. conducted the fluid simulations. A.T., J.G., S.X. and J.W. provided suggestions for the experiment designs. L.Y., M.C., K.N.K., J.R.S. and J.W. designed the figures and revised the manuscript.

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

**Additional information**

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