High-resolution label-free 3D mapping of extracellular pH of single living cells

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Dynamic mapping of extracellular pH (pHe) at the single-cell level is critical for understanding the role of H+ in cellular and subcellular processes, with particular importance in cancer. While several pHe sensing techniques have been developed, accessing this information at the single-cell level requires improvement in sensitivity, spatial and temporal resolution. We report on a zwitterionic label-free pH nanoprobe that addresses these longstanding challenges. The probe has a sensitivity > 0.01 units, 2 ms response time, and 50 nm spatial resolution. The platform was integrated into a double-barrel nanoprobe combining pH sensing with feedback-controlled distance dependance via Scanning Ion Conductance Microscopy. This allows for the simultaneous 3D topographical imaging and pHe monitoring of living cancer cells. These classes of nanoprobes were used for real-time high spatio-temporal resolution pHe mapping at the subcellular level and revealed tumour heterogeneity of the peri-cellular environments of melanoma and breast cancer cells.
Cell survival requires the maintenance of a relatively constant neutral extracellular microenvironment. Extracellular acidification occurs due to the stimulation of anaerobic glycolysis under tumour and inflammatory circumstances. For example, acidic extracellular microenvironments can promote tumour metastasis and modulate inflammatory responses. Therefore, the precise measurement of local extracellular pH (pHe) is critical for understanding the role of H⁺ in cell activity and in turn, its implications in cancer diagnosis and treatment. Measuring the local pHe is also crucial in assessing the extent of tumour invasion, and immune reaction. High-resolution pHe mapping can help to better understand the link between pH distribution, cell morphology and cell function. However, the pHe spatial distribution in the cell microenvironment is exceptionally challenging to map, due to the high mobility and rapid diffusion of extracellular protons. There is, therefore, a need to develop an agent- or label-free high-resolution method for effective and sensitive monitoring of pHe changes at the single-cell level.

At present, the most commonly used pH probes are based on conventional microelectrodes and are limited by large footprint and slow response times. Alternatively, fluorescence-based pH probes can be used in extracellular space, but with considerable limitations due to high background levels and rapid photobleaching. Uses of Nuclear Magnetic Resonance Imaging and Positron Emission Computed Tomography have also been reported, however, these methods have a low spatial resolution, and their reliance on the distribution of the probes within tissue makes the quantification of pHe challenging.

Recently, to address part of this challenge, we developed poly pyrrole-based Field Effect Transistors (FET) at the tip of a nanopipette, which could be positioned in well-defined locations and detect pH changes in aqueous environments. However, these probes had slow response times, severely limiting the dynamic mapping of single cells in real-time. The slow response is due to ionic Coulomb blockade, which reduces the rate of hydronium ion diffusion. One promising route in overcoming such limitations is in the use of functional zwitterionic nanomembranes with high conductivity and electroactivity. An example that we propose to take advantage of is in the drying-mediated self-assembly of a poly-l-lysine/ Glucose oxidase (PLL/GOx) hydrogel, which can be crosslinked by using glutaraldehyde vapour. The positively charged quaternary amines of PLL and GOx) hydrogel, which can be crosslinked by using glutaraldehyde.

Characterisation of pH-sensitivity. The structural properties of the membrane are crucial in optimising the pH-sensing capability. Slice and view Focused Ion Beam Scanning Electron Microscopy (FIB-SEM) was used to image the tip of the pipette. Direct slice and view imaging was possible since the PLL/GOx nanomembranes were tightly crosslinked. A series of milling cross-sectional FIB-SEM images showed that the PLL/GOx nanomembrane formed at the tip of the pipette is ~200 nm thick, Fig. 1c and Supplementary Fig. 1.

With these probes, ion selectivity can be achieved, in a similar manner to other types of nanopore sensors, in which the sensing channel opening has comparable dimension to the Debye screening length (nanometre range in physiological solution). The ion selectivity was evaluated by measuring the reversal potential, Supplementary Fig. 2a, b, which was calculated to be between 15–20 mV per pH unit and had an excellent linear response to a range of pH values.

Results
Fabrication of a pH-sensitive single-barrel nanoprobe. A schematic of the drying-mediated self-assembly process at the tip of a glass pipette is shown in Fig. 1b and Supplementary Fig. 1. The volatile nature of glutaraldehyde enables the introduction of free aldehyde by vapour diffusion into the membrane matrix and also facilitates the covalent bonding of the crosslinked functional groups. This also leads to the formation of a stable zwitterionic-like nanomembrane, which exhibits ion-current rectification, Fig. 1e and Supplementary Fig. 2c. Such unique zwitterionic properties can be exploited for ultrasensitive pH sensing by controlling the total surface charge of the nanomembrane to prevent the formation of anionic Coulomb blockade and allow the interfacial H⁺ to be freely and rapidly transferred from the solution phase to the active sensing area.

We found that optimal conditions could be achieved by using a ratio of ~0.4 mg/ml GOx to 0.01% PLL, which gives an optimised ion-current rectification as can be seen in the I−V response of the nanoprobe, Supplementary Fig. 2c (blue line). Under these conditions, the nanomembrane shows preferential permeability for anions at low-pH and cations at high-pH, Fig. 1d. An excellent linear response (R² = 0.99, p < 0.001) between pH 5 and pH 8 could be obtained, Fig. 1e, f and Supplementary Fig. 2d–f. For physiological applications including sensing of living cells, such improved linear sensitivity is likely to be of greater utility than a wider dynamic range. The properties of these nanomembrane sensors (e.g. surface charge, dynamic range, and pH sensitivity) can be further finely tuned by controlling the GOx to PLL ratio, Supplementary Fig. 2c. GOx was chosen as it is easy to obtain, low cost, has good thermal and pH stability, a long shelf life, and high operational stability. However, other charged proteins/enzymes can also be used such as hexokinase and bovine serum albumin, Supplementary Fig. 3.

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The high ion selectivity made it possible to operate the nanoprobe in potentiometric mode, Supplementary Fig. 2b, as in a conventional pH sensor.

Under the influence of an electric field, Fig. 1e, f, a six-point calibration was performed between pH 4.0–9.0. Changes in ion-current rectification can be observed in the I−V curves at varying pH and followed a linear response. The linear ion response can not only be used as a pH indicator, but also as a feedback signal for use in probe-sample distance control for high-resolution 3D imaging when coupled with SICM. Since the spatial resolution of SICM is closely linked to the probe dimensions, the miniaturisation of the nanoprobe is essential to maximising the spatial resolution, Supplementary Fig. 2d–f. For example, at a holding potential of −0.6 V, the ion current flowing through the nanomembrane into the probe ranges from ~−70 nA for a micropipette with ~2.5 µm inner diameter (imaged with SEM in Supplementary Fig. 1) to about −0.8 nA for a nanopipette with a tip inner diameter of ~100 nm as previously investigated by our groups.
Evaluation of SICM feedback-controlled pH sensing. Combined with SICM, the nanoprobe is capable of non-contact surface scanning and accurate positioning at desired locations over a cell or any other \( \text{H}^+ \) releasing source. In order to thoroughly investigate the capability of the sensor, we generated an artificial \( \text{H}^+ \) gradient using delivery via a voltage-controlled nanopipette, as previously published by our group\(^6,21\). This \( \text{H}^+ \) gradient generation has the advantage that the voltage could be rapidly switched on or off, allowing us to induce highly localised positive \( \text{H}^+ \) release or to perform negative controls in nanoprobe pH mapping experiments, Supplementary Fig. 4a–c. The generated \( \text{H}^+ \) gradient can be manipulated by altering the distances from the \( \text{H}^+ \) delivery pipette, Fig. 1g. To illustrate the spatial resolution, we carried out 2D X–Y plane pH mapping of an \( \text{H}^+ \) supplying nanopipette (\(~100 \text{ nm} \) inner diameter), Fig. 1g and Supplementary Fig. 4a, c. For comparison, a larger pH gradient generated with a voltage-controlled \( \text{H}^+ \) supply micropipette were also mapped, Supplementary Fig. 4b.

To evaluate the response time and sensitivity, we used a fast piezo-stage capable of rapidly changing the position of the sensor, Fig. 1g. Response times of \(~2 \text{ ms} \) could be achieved with a sensitivity better than 0.01 pH units and a spatial resolution higher than 50 nm, Fig. 1f.

The magnitude of the pH gradient depends on cell activity, extracellular buffering and \( \text{H}^+ \) diffusion\(^22,23\). In a non-buffered solution, the high diffusion coefficient enables the protonic charge to travel over 100 µm, whereas, in a physiologically buffered solution the distance travelled can be shortened to about 10 nm\(^23\). Since physiological media in living organisms are typically buffered solutions, we first investigated the buffering effects on nanoprobe pH sensing using an artificial \( \text{H}^+ \) gradient model in solution with different buffering capacity, Supplementary Fig. 5. Proton diffusion is heavily limited and localised with the increasing buffer concentration.

High-resolution pHe mapping of living cells. Parietal cells are responsible for the secretion of hydrochloric acid (HCl) in the stomach\(^24\). With the help of SICM distance control, our
single-barrel nanoprobe pH sensor can be navigated and accurately positioned ~1 µm above the cell surface. It was possible to monitor db-cAMP-induced rapid HCl secretion from a single parietal cell, which can be inhibited with the proton pump inhibitor SCH 28080, Fig. 2a. Similar db-cAMP triggered H⁺ release from a single parietal cell was verified by the acidification of extracellular pH-sensitive BCECF dye using confocal microscopy, Fig. 2a.

Unlike the acidic microenvironment of parietal cells, a significant rise in cell surface pH in algae exposed to light is expected due to photosynthetic uptake of dissolved inorganic carbon. Fluctuations of around 0.3 pH units were observed at 1 µm above the surface of marine diatom *Coscinodiscus wailesii* within 200 s of light exposure, Fig. 2b. No such change in pH could be detected 100 µm away from the cell surface, which was attributed to previous observations that light-induced pH change could be detected 100 µm away from the cell surface.

In SICM, the probe to sample distance is controlled via the feedback-control signal for scanning, which is not only sensitive to approximately one probe radius separation between nanoprobe–cell surface, but also to the extracellular pH changes and can induce ball-like topographical artefact at the tip of the H⁺ supply pipette (dotted-circle highlighted in Supplementary Fig. 6d–g). Although such interference of pH sensing can be partially minimised with constant-height (Supplementary Fig. 6h, i) or feedback-controlled iceberg SICM scanning mode, Supplementary Fig. 7, as will be discussed, this limitation can be overcome with the use of double-barrel probes.

**Fig. 2** SICM feedback-controlled real-time pH detection of living cells. a Bright filed microscopy image showing a pH-sensitive nanoprobe over a buffered single gastric parietal cell (left top). The cartoon at the bottom illustrates the construction of extracellular pH measurements (left bottom). Feedback-control allows for positioning of the pH nanoprobe to accurately detect 100 µm db-cAMP (cAMP analogue) triggered via rapid HCl secretion from a single gastric parietal cell. The secretion can be inhibited with selective H⁺, K⁺ ATPase inhibitor SCH28080 (middle). The db-cAMP-induced HCl secretion from the gastric parietal cells was confirmed with pH-sensitive BCECF fluorescent dye by confocal microscopy (right). Changes in fluorescent intensity of pH-sensitive dye can be mapped out either by x-y or by x-z scanning. The dotted crossline marks the Z position in the XZ image and the Y position in the XY image, respectively. b pH detection from a single low-buffered *Coscinodiscus wailesii* cell. Changing light conditions from illumination (yellow stripes represent light illumination) to darkness (grey stripes represent darkness), and vice versa revealed rapid pH changes ~1 µm above the cell surface. Change in pH is almost undetectable when the probe is 100 µm away from the cell surface.

*High-resolution 3D pH mapping of live cancer cells.* To decouple the SICM scanning ability from the pH sensing, we fabricated a double-barrel nanoprobe. As demonstrated in the operational (Fig. 3a) and fabrication (Fig. 3b) schematics, the double-barrel SICM-pH nanoprobe consists of an unmodified open barrel (SICM-barrel) for SICM control and another barrel with a pH-sensitive PLL/GOx ommembrane (pH-barrel), which enables both pH measurement and SICM topographical imaging simultaneously and independently. The ion-current flowing into the two independent barrels of the double-barrel nanoprobe showed very different I-V responses at varying pH, Fig. 3c. Much like the single-barrel case, the dynamic range, linearity, and sensitivity were similar. In order to measure local pH accurately, a self-referencing 3D mapping protocol that is used in multifunctional SECM-SICM was employed. Note that such self-referencing measurements allow the response of local pH near to the cell surface (about 100 nm) to compensate for the possible pH drift in bulk (~10 µm over) at every pixel of SICM 3D pH mapping.

We further applied the double-barrel SICM-pH nanoprobe to measure the dynamic change of pH gradients in breast cancer MCF7 cells at the single-cell level, Fig. 3d–f. Previously, it has been demonstrated that CD44<sup>high</sup> breast cancer cells bear similarity to cancer stem cells and are involved in aggressive phenotypes. Estrogen deprivation therapy is used in the treatment of breast cancer; our previous study demonstrated that only a small fraction of CD44<sup>high</sup> cells could adapt to such estradiol-deprivation. Using MCF7 cells engineered to drive GFP expression under the promoter of the CD44 gene, estradiol-deprived (−E<sub>2</sub>) CD44<sup>GFP-high</sup> cells showed a strong pH gradient towards acidification, Fig. 3d, as opposed to estradiol-supplemented (+E<sub>2</sub>) CD44<sup>GFP-high</sup> cells, which do not generate any noticeable pH gradient, Fig. 3f. However, pH mapping across these −E<sub>2</sub> CD44<sup>GFP-high</sup> cells further corroborates the phenotypic heterogeneity in this subpopulation of MCF7 cells, Fig. 3e. This is in line with the heterogeneity in single-cell transcriptomics data, Supplementary Fig. 8, and our previous observations at the transcriptional level.

It has been proposed that thymosin beta-4 × -linked (TMSB4X) may positively regulate the activity of ATP-synthase to transport proton from the intracellular to the extracellular space of cancer cells. Moreover, an ATP-driven, vacuolar proton pump (V-ATPase) has been suggested to be involved in the regulation of extracellular acidification by cancer cells. Using single-cell RNA-sequencing, we found a significantly higher expression of TMSB4X and ATP6V1F in estradiol-deprived (−E<sub>2</sub>) CD44<sup>GFP-high</sup> compared
**Fig. 3** Independent SICM feedback-controlled scanning and simultaneous 3D pH mapping of living cells. 

**a** A schematic showing the operation of double-barrel nanoprobe for simultaneous SICM imaging and pH measurement. 

**b** A pH-sensitive nanomembrane is formed inside one barrel (pH-barrel) of a double-barrel θ quartz glass nanopipette, while the second barrel (SICM imaging-barrel) is kept open via applied back pressure during fabrication. 

**c** The ion-currents flowing into two separated barrels of the generated double-barrel nanoprobe show different I-V responses to pH. 

**d** SICM imaging and 3D pH mapping of a group of low-buffered CD44GFP-high breast cancer MCF7 cells in estradiol-deprived medium (−E2). The SICM topographical images (left), fluorescence image (GFP, middle), and 3D pH distributions (right) can be simultaneously obtained from a single scan. 

**e** Same as d but using a different group of estradiol-deprived (−E2) CD44GFP-high cells. 

**f** Same as d but using a group of CD44GFP-high cells under estradiol-supplemented culture (+E2). Scale bars represent 20 μm. Intensity of fluorescence images have been normalised.
to estradiol-supplemented (+E2) CD44\textsuperscript{GFP-high} cells, Supplementary Fig. 8, which may explain the observed difference in pH\textsubscript{e}, Fig. 3d–f. Adding 10 mM HEPES buffer eliminated most of the observed pH\textsubscript{e} gradient of these estradiol-deprived (−E2) CD44\textsuperscript{GFP-high} MCF7 cells, Supplementary Fig. 9. Three experiments with continuous repetitive 3D pH\textsubscript{e} mapping revealed profound effects of buffer on the profile of pH gradients and demonstrated noticeable dynamically changes in pH\textsubscript{e}, while the changes in topographical and fluorescence images were virtually undetectable, Supplementary Fig. 9. Such buffer treatments may help us to study the dynamic changes in local H\textsuperscript{+} distribution near to cell surface in the future.

Since melanoma represents one of the most aggressive and heterogeneous forms of cancer, exhibiting the most diverse cell subpopulations and showing extracellular acidity\textsuperscript{32}, we further applied our nanoprobe for pH\textsubscript{e} mapping to living melanoma A375M cells, Fig. 4. The high-resolution SICM topographical images (left panel of Fig. 4) and pH\textsubscript{e} 3D mappings (right panel of Fig. 4) of the same melanoma cells could be obtained simultaneously in a single SICM scan. As expected, high-resolution 3D pH\textsubscript{e} mapping among these melanoma cells demonstrated a highly variegated distribution pattern, which is in agreement with the most heterogeneous nature of melanoma cells. In contrast, the pH\textsubscript{e} mapping of non-cancerous melanocytes did not generate a noticeable pH\textsubscript{e} gradient, Supplementary Fig. 6a–c.

**Discussion**

It is becoming clear that an acid pH\textsubscript{e} plays an essential role in cancer cell progression, invasiveness and resistance to therapy\textsuperscript{1,33}. Monitoring the pH\textsubscript{e} variation of cancer cells is likely to provide critical information for understanding tumour heterogeneity during pathological processes such as epithelial-mesenchyme transition\textsuperscript{33}. However, a high diffusion coefficient and heterogeneous distribution of extracellular protons make real-time monitoring of pH\textsubscript{e} more difficult\textsuperscript{34,35}. The development of label-free high-sensitivity high-resolution methods to detect pH\textsubscript{e} at the single-cell level is urgently needed\textsuperscript{7}.

In this study, pH-sensitive nanoprobes have been developed by crosslinking GOx and PLL to produce a drying-mediated self-assembly nanomembrane at the tip of glass nanopipettes, resulting in high spatial resolution of 50 nm, about 2 ms rapid response time, 0.01 pH sensitivity, and with minimal disturbance of the samples. A typical probe can remain functional for ~180 days (dry shelf-life at room temperature) and can be used under varied ionic strength conditions from freshwater to seawater as shown in Supplementary Fig. 10 and Supplementary Fig. 11.

High H\textsuperscript{+} sensitivity of probe is likely due to the pH-dependent modification of the surface charge of the zwitterion-like PLL/GOX nanomembrane\textsuperscript{16,36} or intramolecular interactions among amine and carboxyl groups\textsuperscript{37,38} within PLL/GOX nanomembrane, Fig. 1d. The higher surface-to-volume ratio of the porous nanoneedle may also help the accumulation and adsorption of H\textsuperscript{+} and consequently, the pH sensitivity. Although these hypotheses may explain the sensitivity and precision of our pH-sensitive nanoprobe, further modelling and investigations are still required.

SICM feedback-controlled pH-sensitive single-barrel probes have been used to detect spatiotemporal pH\textsubscript{e} around a single acid-releasing parietal cell and photosynthesis-induced alkaline of algae. Although the single-barrel probe demonstrated the possibility to map pH\textsubscript{e} over a single immune cell or a group of melanoma cells with constant-height or iceberg SICM operational mode, the interference of pH in feedback ion currents made accurate 3D pH\textsubscript{e} mapping of living cells challenging to achieve. We therefore fabricated a double-barrel SICM-pH nanoprobe to combine pH sensing with robust independent SICM feedback control to perform reliable 3D pH\textsubscript{e} mapping around living cells. In order to improve the accuracy of local pH\textsubscript{e} detection around a single cell, a self-referencing approach was introduced to compensate for possible pH drift in bulk solution of the living system\textsuperscript{26}. Previous studies have demonstrated that tumour cells can generate 0.2–0.6 units lower pH\textsubscript{e} gradients than the normal tissues at single-cell level\textsuperscript{39}. We detected about 0.7 units lower pH\textsubscript{e} gradients of a group of estradiol-deprived (−E2) CD44\textsuperscript{GFP-high} breast cancer MCF7 cells (Fig. 3d, e), and about 0.2 units lower pH\textsubscript{e} gradients of a group of A375M melanoma cells (Fig. 4). Intratumoral heterogeneity has long been recognised as a general characteristic of most cancers, particularly melanoma, which is among the most aggressive and therapy-resistant of human cancers\textsuperscript{40}. The spatial resolution of 3D pH\textsubscript{e} mapping of A375M cells shows a variegated pH gradient pattern (Fig. 4), which may indicate the exceptional level of intratumor heterogeneity and the presence of cell subpopulations with different phenotypes and biological behaviour\textsuperscript{32,40}. Such heterogeneous distribution of pH\textsubscript{e} observed from melanoma cells may also relate to the inhomogeneously distributed proton transporters to the cell border or leading-edge pseudopodia of these invasive cancer cells\textsuperscript{41}.

Although single-cell genomics makes it possible to analyse the intratumoral heterogeneity, understanding the cellular heterogeneity of cancer cells is still a big challenge\textsuperscript{42}. Using CD44 as a biomarker to select breast cancer stem cells for single-cell sequencing, we have demonstrated that CD44\textsuperscript{GFP-high} breast cancer MCF7 cells consist of multiple subpopulations with marked heterogeneity\textsuperscript{28}. In this study, pH\textsubscript{e} 3D mapping also showed such heterogeneity among CD44\textsuperscript{GFP-high} cells under
culture when they were estradiol-deprived (−E2) or estradiol-supplemented (+E2), and even heterogeneity among these estradiol-deprived (−E2) CD44<sup>+</sup>SP-high aggressive and plastic cell subpopulations (Fig. 3d, e). The single-cell RNA-sequencing showed upregulated gene expressions of extracellular acidification related proteins TMSB4X and V-ATPase<sup>9,20</sup> (Supplementary Fig. 8), which may partially explain our observed pH heterogeneity of MCF7 cells. In future work, we plan to assess the heterogeneity of these cancer cells with our double-barrel nanoprobe functional pH mapping and then select cell subpopulations to perform nanobiopsy sampling from these individual cells<sup>34,41</sup>. This will enable a method for function-driven single-cell sequencing and thus further understanding of cancer heterogeneity, cellular plasticity and therapeutic resistance at the single-cell level.

Extracellular acidic pH enhances the invasive behaviour and the migration of cancer cells and may define an additional barrier for therapy<sup>9</sup>. Normalising the pH gradient through inhibition of various proton transporters or by systemic buffering with bicarbonate is of therapeutic benefit to cancer treatment<sup>14</sup>. The double-barrel SICM-pH nanopores, self-referencing sensing platform enables real-time feedback-controlled dynamically 3D mapping of pH heterogeneities of cancer cells label-free and at subcellular resolution. This method could help with a cancer diagnosis, prognosis, and in evaluating acidic pH targeted therapies.

Methods

Chemicals. Poly-l-lysine (PLL, P4707), glucose oxidase (GOx, G2133), HCl and glutaraldehyde were purchased from Sigma–Aldrich. Phosphate buffered saline (PBS) solution was prepared from 7.2 mM Na<sub>2</sub>HPO<sub>4</sub>, 2.8 mM KH<sub>2</sub>P<sub>4</sub>, 137 mM NaCl and 2.7 mM KCl (pH 7.4). Hanks’ Balanced Salt Solution (HBSS) containing 5 mM bicarbonate and 0.8 mM phosphate buffer (pH 7.4) was purchased from Gibco.

Fabrication of the glass pipettes. Single barrel: pH-sensitive nanomembrane probes were fabricated by pulling a borosilicate glass capillary (O.D. 1 mm, I.D. 0.58 mm) to micropipettes with a tip diameter of ~1 μm or nanopipettes with a diameter of ~100 nm using a laser-based puller (Model P-2000, Sutter Instruments Co., USA). Nanopipettes were pulled with a two-step protocol. Briefly, for the initial step, the parameters were: heat 350, filament 3, velocity 30 and delay 200. For the second step, they were: heat 350, filament 2, velocity 27, delay 160 and pull 250. It should be noted that these values are instrument-specific, and parameters would have to be optimised for each instrument in order to obtain similar nanopores.

Double barrel: Double-barrel pH nanopores were constructed from a double-barrel pulled double-barrel quartz theta capillary (O.D., 1.2 mm, I.D., 0.9 mm, Sutter Instruments), which was pulled with a laser-based P-2000 pipette puller (Sutter Instruments) using a single line program (heat 700, filament 3, velocity 45, delay 130, and pull 93) to produce sharp double-barrel nanopipettes. The size of each barrel of this pulled double-barrel nanopipette was about 100 nm.

Assembly of the pH-sensitive nanomembrane. Glucose oxidase (GOx) has a pI of 4.2 and is therefore negatively charged at physiological conditions. GOx was firstly inactivated by denaturation at 70 °C for 10 min before nanomembrane fabrication. This was then dissolved in 0.01% (v/v) poly-l-lysine (PLL) at a concentration of 0.4 mg/ml. Further details can be found in Supplementary Note 1. The positively charged quaternary amines of PLL and negatively charged carboxylic acid residues of GOx facilitate the formation of a self-assembled hydrogel at the tip of nanopipettes. Dry-transported self-assembly of the PLL/GOx hydrogel can then be crosslinked with glutaraldehyde vapour, Fig. 1b. Briefly, the pulled glass pipette was filled to a length of about 1 mm with a mixture of PLL and GOx by capillary action. The crosslinking reaction between PLL and GOx at the tip of the pipette was initiated and was allowed to proceed overnight in the vapour of 2% (v/v) glutaraldehyde at room temperature. Following this slow evaporation of the aqueous solution and co-condensation at the air-water interface (at the pipette tip), strong covalent crosslinking between the amino groups of PLL and GOx with the glutaraldehyde were formed, and a self-assemble nanomembrane at the tip of glass pipette was generated. Before using the nanomembrane probes were washed with 100 mM pH 7.0 KCl to remove unreacted glutaraldehyde and any un-crosslinked reagents. The nanomembranes are very stable, and we have observed no degradation of pH stability over the duration of continuous experiments in solution (up to 3 h), Supplementary Fig. 4d.

For experiments using double-barrel nanopipettes, the procedure was similar; however, one of the barrels was closed with “Blu-Tack” (Bostik, UK). The second, open barrel was kept away from sucking in PLL/GOx solution by holding a positive ~200 kPa pressure to compensate for the negative capillary force. As a result, only one barrel was modified with crossed PLL/GOx nanomembrane, another unmodified open barrel was used for SICM feedback control, Fig. 3a.

Scanning ion conductance microscopy. All experiments were performed using a customised SICM setup and operated using an ICAPPC Controller (IC-UN-001, ICAPPC Ltd., UK). The SICM scan head consisted of a PIHera P-621.2 XY Nano-positioning Stage (Physik Instrumente, Germany) with 100 × 100 μm travel range that moved the sample and a LISA piezo actuator P-753.21 C (Physik Instrumente, Germany) with travel range 25 μm for pipette positioning along the Z-axis. Coarse positioning was achieved with translational stages M-111.DG (XY directions) and M-112.DG (Z-axis) (Physik Instrumente, Germany). Piezo actuators were powered by high voltage amplifiers E-503 and E-505 and servo module E-509 (Physik Instrumente, Germany). SICM control, data acquisition and analysis software were written and kindly provided by Dr Pavel Novak, ICAPPC Ltd. Ion current was detected and recorded using a MultiClamp 700B amplifier (Molecular Devices, UK) using a 2 kHz low-pass filter. A typical external holding voltage of ~200 mV was supplied to the scanning probe. The ion current and output of the capacitive sensors from all three piezo elements were monitored using an Axon Digidata 1322 A digitiser and Clampex 9.2 software (Molecular Devices, UK).

Cyclic voltammetry and signal recording. The nanoprobe pH sensor was back-filled with 100 mM pH 7.0 KCl, contacted with an Ag/AgCl wire and immersed into a 2 mL physiological bulk solution (PBS, low-buffered solution, or buffered solution). Another Ag/AgCl electrode was placed in bulk solution acting as a reference electrode. All potentials were quoted against this reference electrode. Both electrodes were connected to a MultiClamp 700B amplifier and digitised with an Axon Digidata 1322 A (Molecular Devices) and Clampex 9.2 (Molecular Devices, UK). The potential was cycled between −0.6 V and +0.6 V vs. Ag/AgCl at a scan rate of 650 mV s<sup>−1</sup>. The recorded ion currents flowing in the nanoprobe were low-pass filtered at 1 kHz and analysed with pClamp 10 software (Molecular Devices). All measurements were performed in an air-conditioned room. The solution temperature was monitored throughout with a TC-344B Automatic Temperature Controller (Warner Instrument Corporation) and was maintained at 26 °C during pH experiments.

SICM feedback-controlled pH mapping. Scanning and local pH measurements were performed with a customised SICM setup and controller (ICAPPC Ltd., UK). For pH mapping, each pH nanoprobe was firstly calibrated with pH adjusted solutions. Then the pH-sensitive nanopores were typically held at ~200 mV vs. Ag/AgCl and used like any other standard SICM scanning probes. The SICM probe-sample distance was regulated by monitoring ion current through the nanoprobe pH sensor, which can be accurately positioned towards the cell surface under feedback control (without contact from the nanoprobe itself), and a pH measurement at each SICM topographical scanning point can be used to generate a 3D map of the pH distribution. All mapping, data acquisition and analysis software were developed by Dr Pavel Novak, ICAPPC Ltd.

Calibration and evaluation of the nanopore pH sensor. The pH-sensitive nanomembrane probes were calibrated using standard pH buffered solutions at different pH values and benchmarked against commercially available pH metre (MP 220, Mettler Toledo). Only nanopores that exhibited a linear response within the pH range used in further studies. To test the platform, artificial studies were reversibly generated using tunable voltage pulses. The delivery pipettes were immersed into the PBS (pH 7.4) solution and backfilled with 100 mM KCl in order to generate an H<sup>+</sup> gradient. A positive voltage of 0.5 V vs Ag/AgCl was locally delivered H<sup>+</sup>, whereas, a retaining voltage of ~0.5 V was applied to prevent a release and could be used as a negative control for nanoprobe pH detection. SICM feedback position control and sub-ms fast response of the Linear Piezo Stage in our SICM setup allowed for rapidly moving the pH sensor along the generated pH gradient by changing the distance from the H<sup>+</sup> source. The response time was defined as the time required to reach the equilibration output of the nanoprobe.

Double-barrel SICM-pH nanopore 3D pH mapping. Cells were placed on the bottom of a glass microscopy dish (35 mm diameter) and imaged using a Nikon inverted microscope. Measurements of pH on living cells were performed in low-buffered media by adding 1/4 HBSS to 137 mM NaCl 5 mM KCl and 25 mM D-glucose (pH 7.4), which helps to maintain the pHe gradients. A schematic diagram shows the operating principles for both SICM imaging and pH being measured simultaneously, Fig. 3a. When combined with SICM, the open barrel (SICM-barrel) was used to measure a probe-cell surface distance in the same fashion of traditional SICMs. The second pH-barrel was functionalised with PLL/GOx nanomembrane to investigate the pH distribution around scanned cells. As with the single-barrel pH nanoprobe, the nanoprobe was backfilled with 100 mM KCl (pH 7.0). A typical ~200 mV bias was applied to the open barrel (SICM-barrel) to induce an ion current for SICM feedback control and imaging; a typical 200 mV bias was applied to induce ion current flowing into the pH-barrel for H<sup>+</sup> gradient mapping, Fig. 3c.

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ARTICLE
SCM hopping mode allows self-referencing measurements as previously used in multifunctional SESC-SICM to compare the response in bulk and near the surface at each pixel. We hereby detail if the nanoprobe signal drifts over time in a controlled environment (illumination: 120 μmol·m⁻²·s⁻¹), light/dark: 16 h/8 h; temperature: 15 °C) at Silwood Park, Imperial College London. To prepare the AQUIL laboratory, we used acclimated (24 h in 10% HNO₃, Fisher Scientific) and rinsed with Milli-Q water at least four times. Chemicals of ACS grade or higher purity were purchased from Sigma–Aldrich, and the medium containing synthetic ocean water and major nutrients was sterilized at 121 °C for 15 min before adding 0.2-µm filtered (polycarbonate filters, Merck Millipore Ltd.) solutions of EDTA–metals and vitamins. Diatoms were placed in a glass-bottomed microscopy plate (35 mm diameter, Commun and observed using an inverted micro-scope. During the measurements, the diatoms were kept in low-buffered artificial seawater (0.5 M NaCl + 1/10 PBS, pH 7.65) exposed to light or darkness. Cells were illuminated at 200 μmol·m⁻²·s⁻¹ using a blue light source, and there was no change in temperature within the dish during illumination.

Focused ion beam milling and SEM imaging. Nanomembrane probes were cut to the desired length, mounted on sample stubs and coated with all sides with 10 nm of chromium in a sputter coater (Q150T S Quorum). The probes were imaged using a FIB-SEM (CrossBeam Workstation Auriga, Carl Zeiss). Tips on the nanomembrane were located using secondary electron imaging with an accelerating voltage of 5 keV. The sensor tip was milled using a milling current of 120 pA at a working distance of 5 mm. Sections of 10 nm were ion milled and imaged by SEM at 1.6 keV, using a secondary electron in-lens detector.

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

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Cell culture. The human malignant melanoma cell line A375M and human immortal melanocyte line Hermes 3A were both obtained from the Wellcome Trust Functional Genomics Cell Bank (St George’s, University of London, UK). A375M cells were grown in RPMI 1640 (Gibco). Culture medium was supplemented with 10% fetal bovine serum (Sigma–Aldrich), 100 U/mL streptomycin and 100 U/mL penicillin in an atmosphere of 10% CO₂ at 37 °C. Hermes 3A melanocytes were grown in RPMI 1640 medium supplemented with 10% fetal calf serum (Sigma–Aldrich), 200 mM 12-tetradecanoylphorbol acetate, 200 μM choler toxin, 10 ng/mL human stem cell factor and 10 mM β-2-mercaptoethanol. The Wellcome Trust Functional Genomics Cell Bank (St George’s, University of London, UK) recommends using 10% CO₂ for growing cells of melanocytic lineage.

Primary gastric parietal cell isolation and culture. Isolated rat stomach was turned inside out, and it was treated with a C₆H₁₄–free DMEM containing 20 mM HEPES (pH 7.4), 0.5% BSA, 0.1 mg/ml trypsin inhibitor and 1 mg/ml Pronase E for 20 min at 37 °C. After the cell debris was removed, the stomach was further digested in DMEM containing 20 mM HEPES (pH 7.4), 0.5% BSA, 0.1 mg/ml trypsin inhibitor, 1.5 mM Pronase E, and 1 mg/ml collagenase for 30 min at 37 °C. The mixture was then filtered through a cell strainer and washed with DMEM. Isolated cells were incubated with 1 mg/ml amphotericin B for 15 min at 37 °C. After washing, the cells were loaded onto a discontinuous Optiprep (Axis-Shield) gradient. After centrifugation at 800 × g for 8 min, the fraction enriched in parietal cells (around 70% of the total) was collected and plated onto collagen-coated coverslips in 12-well plates and incubated at 37 °C in DMEM containing 2 mg/ml BSA, 10 mM glucose, 5 mg/ml ITES, 8 mg/l EGF, 5 mg/ml G418, 400 mg/ml gentamicin, 20 mg/ml novobiocin, 10 mM hydrocortisone, 100 U/ml penicillin/streptomycin and 20 mM HEPES (pH 7.4). The experiment using the nanomembrane pH probe sensor, parietal cells were identified by using anti-H⁺, K⁺-ATPase β-subunit antibody (D032-3H, 2B6, Medical & Biological Laboratories) and diluted 1:250 Alexa Fluor 488-conjugated anti-mouse IgG antibody (ab150105, Abcam).

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pH-sensitive fluorescent dye. Fluorescein derivative 2-[(2-carboxyethoxy)-5- (and-6)-carboxyfluorescein (BCECF) is the most widely used pH probe, whose excitation ratios at two different wavelengths are correlated to pH. Since BCECF does not permeate the cell membrane, and fluorescence can be measured outside the cell. BCECF (Dojindo) was chosen to evaluate the extracellular pH changes of gastric parietal cells. The fluorescence levels of BCECF around parietal cell were measured with a Leica TCS-SP5 Confocal Microscope. Z stack images were acquired using a 1 μm Z step, and the resolution was calculated to be around 0.2 μm/pixel.
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Author contributions
Y.Z., E.V.S., J.B.E., Y.K. designed and supervised the research. Y.Z., Y.T., S.P.H., F.L., S.G., I.B. performed the experiments and analysed the data. J.B., P.S.G., H.S., Y.S., T.F., A.E., P.G., A.M. prepared the cell samples. P.N., A.S., A.P.I., L.M. helped with the experiments and analysed the data. Y.Z., E.V.S., J.B.E., Y.K., D.J.W., C.E., D.K., A.P.I. prepared the manuscript. All the authors contributed to the results and commented on the manuscript.

Competing interests
P.N., A.S., A.E., P.G., C.E., D.K. and Y.K. are shareholders in ICAPPIC Ltd, a company commercialising nanopipette –based instrumentation. All other authors declare no competing interests.

Additional information
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