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Multiplexed printed sensors for *in situ* monitoring in bivalve aquaculture†

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Non-intrusive sensors that can be attached to marine species offer opportunities to study the impacts of environmental changes on their behaviors and well-being. This work presents a thin, flexible sensor tag to monitor the effects of dissolved oxygen and salinity on bivalve gape movement. The measurement range studied was 0.5–6 ppm for the dissolved oxygen sensor and 4–40 g kg⁻¹ for the salinity sensor. The curvature strain sensor based on electrodeposited semiconducting fibers enabled measurements of an oyster's gape down to sub-mm displacement. The multiplexed sensors were fabricated by low-cost techniques, offering an economical and convenient platform for aquaculture studies.

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1. Introduction

Understanding the influence of environmental changes on marine life is important to ecological sustainability and aquaculture. For example, rising temperatures reduce the solubility of oxygen in water, leading to more frequent and intense occurrences of hypoxia that devastate aquatic life.¹ Massive algae blooms have exacerbated hypoxia in aquafarms and resulted in poor yields and shellfish toxicity² that distress global oyster industries, which are worth well over \$6 billion US dollars annually.³ As reef-building organisms, oysters also generate habitat for other environmentally and commercially important organisms such as various fish and crustaceans.⁴ Compact, non-intrusive sensors attachable to these marine species are highly desirable for gathering local environmental parameters to correlate organism physiology, migration behaviors, and compensatory responses to adverse conditions.^{5,6}

Building on recent advances in flexible aquatic sensors,^{7–10} this work demonstrates a printed multi-modal sensor tag suit-

able for *in situ* monitoring in complex marine environments. Specifically, the multiplexed sensor tag enables tracking of bivalve gape responses to local changes in dissolved oxygen (DO) concentration and salinity to aid in understanding environmental impacts on oyster behaviors. Sensing fluctuations in DO and salinity are essential because these parameters cause stress regulation and critically affect bivalve growth and mortality.^{11–14} In addition, bivalve closure has been shown to provide early warning of detrimental conditions for shellfish populations.¹⁵ However, prior works on gape measurements were conducted with complex instrumentation such as a magnetic Hall probe^{16,17} or limited to a binary classification between open or closed states.¹⁵ Here we developed a sensitive, miniaturized curvature sensor to monitor gaping activities down to sub-millimeter displacement. The integrated sensor tag with three sensing modalities is flexible and small with an area <3 cm² for non-intrusive placement onto an individual bivalve and can facilitate widespread monitoring at low costs.

The fabrication of our prototype used inexpensive techniques including a digital blade-cutter, lamination, stencil printing, and electrodeposition. The DO and salinity sensor electrodes were cut from flexible graphite foils that are stable in seawater when compared to silver inks used in additive printing.^{18–21} The curvature sensor was based on an organic electrochemical transistor (OECT), and the sensing channel semiconductor was electrodeposited by an alternating current (AC) waveform^{22,23} that controlled micro-fiber formation with nanoporous structures. The robust fibers offer the advantage of operation in seawater without encapsulation. The sensors were connected through a multiplexer circuit to a handheld potentiostat to achieve a portable system. A proof-of-concept demonstration was conducted using the integrated sensor tag

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in laboratory trials to monitor an oyster's gape, as well as ambient salinity and DO concentration down to oxygen concentrations associated with hypoxia. The thin, flexible design enables local measurements in confined spaces that are not accessible using conventional bulky probes.

II. Results and discussion

In the schematic shown in Fig. 1, the concentric rings represent electrodes for DO and salinity detection. Such arrangement minimized the electrodes' footprint and used the same reference and counter electrodes for both DO and salinity measurements. The seawater sample contained the analytes and also acted as the electrolyte in the electrochemical measurements. Of the two working electrodes, one was for sensing the DO level based on the oxygen reduction reaction, and the other was used to measure the seawater conductivity, which at a constant temperature is proportional to its salinity in grams of salt per kilogram of water. The potentials of the working electrodes (WE) were controlled with respect to the reference electrode (REF), and the corresponding currents were collected through the counter electrode (CE). Next to the circular electrodes, an OECT structure was patterned to function as the curvature sensor to monitor gape movements. The drain and gate electrodes were controlled by the two WE channels on the potentiostat, while the source electrode was con-

nected to the REF and CE channels tied together for current read-out. The device dimensions and fabrication procedures are explained in details in the Experimental section and ESI Fig. S1.†

The photographs in Fig. 1 show all three sensors integrated on the flexible poly(ethylene terephthalate) (PET) substrate. The tag was wrapped around the oyster hinge and externally attached to the shell using adhesive without the need to insert anything into the gape since that would be invasive to the organism. The connections to the sensors were directed into the multiplexer circuit, which controlled the sequential read-out of the three sensors by the potentiostat. The timing diagram in Fig. 2 shows the switching of multiplexer control signals (MUX) to each sensor. When the MUX signal was turned on at the selected sensor, the bias on the WE channel was passed to the active device, and the current at the CE channel was measured. In this work, the MUX on-period was 10 seconds, and we used the readout data near the end of each period when transient current spikes from voltage switching settled and the sensor was in a steady state for accurate detection.

II.A. Curvature sensor based on an OECT

To monitor bivalve gape movements, this work developed an OECT with nanoporous semiconducting fibers that are advantageous for bending curvature measurements. Compared to other pressure^{24–26} or strain sensors^{27–31} fabricated by coating

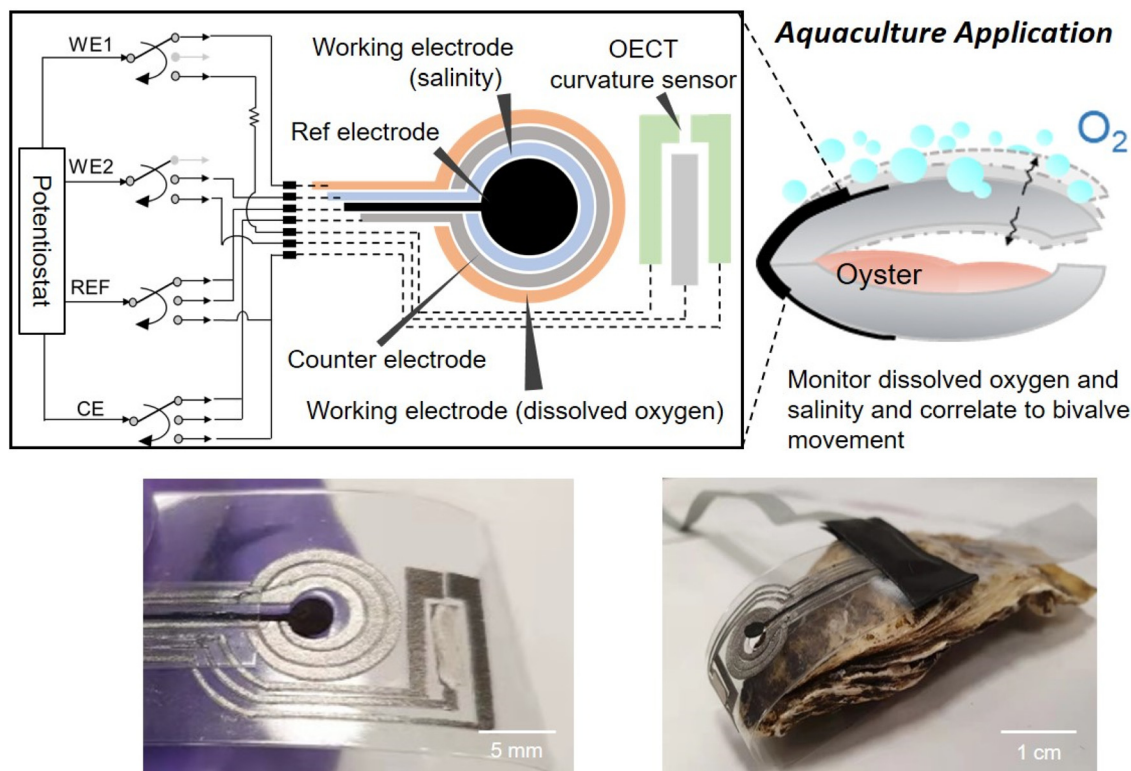


Fig. 1 Schematics and photographs of the printed multiplexed sensors for monitoring bivalve gape response to environmental variables of dissolved oxygen and salinity.

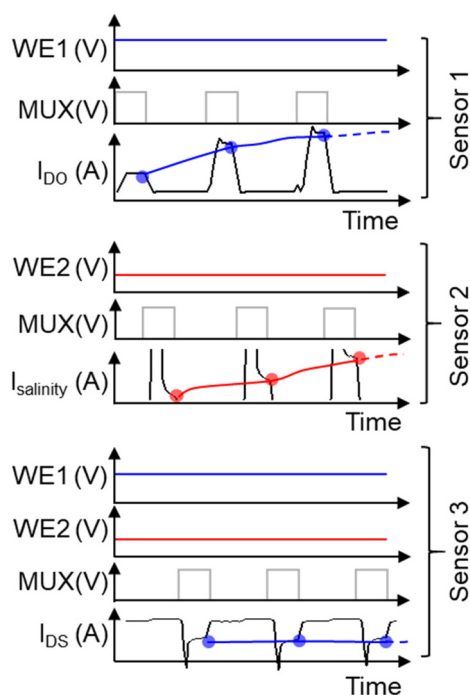


Fig. 2 Timing diagram of control and measurement signals for three sensors, under sequential readout switched by the multiplexer circuit interfacing with the potentiostat. The solid dots indicate steady-state measurement points near the end of each sampling period.

films, the electrodeposition process to form semiconducting fibers facilitated their inherent alignment in that the fibers would grow along the direction of an applied electric field until they bridge the electrodes. Then the semiconducting fibers could be aligned with the bending curvature to maximize their structural deformation and in turn their change in resistance with bending strain. We chose the OECT structure over simple resistor configuration, because the gate bias allowed amplification of current changes to increase the device sensitivity.

Fig. 3a shows the chemical structure of the polymer poly(3,4-ethylenedioxythiophene)-hexafluorophosphate (PEDOT:PF₆), which was electrodeposited using an AC waveform²³ applied across the source and drain electrodes in a solution of monomers and salt mixture (details described in the Experimental section). In brief, the 3,4-ethylenedioxythiophene (EDOT) monomers were oxidized forming radical cations, which were charge balanced by PF₆⁻ anions. The sequential reaction with other EDOT⁺ monomers enabled the growth of the polymer chain following the electric field lines, with fiber tip growth more kinetically favorable because of stronger local field compared to flat edges of electrodes. The fiber morphology depended on the amplitude and frequency of the electrodeposition voltage waveform, as shown in ESI Fig. S2.† A waveform frequency higher than 50 Hz resulted in more bifurcation and branching in the fibers, but low frequency less than 5 Hz led to dense and wide strips. We used a triangular waveform of 12.5 Hz and ±10 V to form thin fibers

with a porous surface as seen in the scanning electron microscopy image in Fig. 3a. The OECT channel conductance was measured to be 40 mS cm⁻¹ and reproducible over multiple deposition runs (ESI Fig. S2†).

The current–voltage characteristics of the PEDOT:PF₆ OECT were taken in simulated seawater (Instant Ocean®) or when covered by a hydrogel electrolyte with a seawater-equivalent ionic strength of 0.6 M NaCl. There was negligible difference between the two measurements (ESI Fig. S3a†). In Fig. 3b–g, characterization was carried out with the hydrogel for convenience, while the data in all other figures were measured with the device submerged in seawater. The OECT operated in depletion mode, and applying positive gate bias de-doped the semiconductor and reduced the channel conductance as seen in Fig. 3c and d. The maximum transconductance was $g_m = 0.88$ mS, and the device on/off ratio was ~500. The mobility–capacitance product (μC^*) was extracted²³ to be 1.24 F cm⁻¹ V⁻¹ s⁻¹, assuming a geometry of fiber width $w = 75$ μm, channel gap $L = 250$ μm, and fiber channel number $n = 2$. In OECTs, it is preferable that the tunable channel resistance exceeds all other elements combined, including contact resistance^{32,33} between polymer/electrode interfaces and wire connections. The contact resistance of our OECT was estimated by fitting to the Y-function model^{34,35} to be 240 Ω, similar to the value extracted by electrochemical impedance spectroscopy (ESI Fig. S3b and S3c†). Meanwhile, the OECT channel resistance was ~1 kΩ at the gate bias of $V_{GS} = 0$ V. Thus, the resistance across the channel was higher than at contacts, which was favorable for strain sensing to obtain large resistance change with deformation of channel materials.

Fig. 3e displays the OECT source–drain current I_{DS} decreased as the substrate was bent to tighter radii of curvature from $r = 9$ cm to 2 cm, corresponding to the curvature $1/r = 0.11$ cm⁻¹ to 0.5 cm⁻¹, respectively. The sensor current was reproducible at different bending speeds from 0.5 cm s⁻¹ to 5 cm s⁻¹ (ESI Fig. S4a†). The I_{DS} exhibited excellent recovery after bending cycles, and the device in a flat state returned to the baseline with only a small drift of <6% over 6 min, potentially due to bias stress^{36,37} that can be further reduced by changing the duty cycle or compensated by drift subtraction. While the channel current I_{DS} was clearly influenced by bending strain, the gate leakage (I_{GS}) was a thousand-fold lower than I_{DS} and showed no apparent response to bending (ESI Fig. S4b†). The steady I_{GS} confirmed that strain in the hydrogel, graphite electrodes, and interconnects did not induce significant current response, and only the semiconducting fibers were the major contributor to the I_{DS} change upon bending. In Fig. 3f, the transconductance was reduced with increasing curvature, comparable to other strain measurements on PEDOT films.^{27,38} This trend was attributed to percolation effects, that the applied strain stretches the semiconductor, disrupts conductive paths, and hence decreases the transconductance.

Fig. 3g provides the calibration curves of OECT curvature sensors. The ratio $\Delta I/I_0$ was defined as the change in I_{DS} between the baseline flat state and bent state [$\Delta I = I_{DS}(0)$

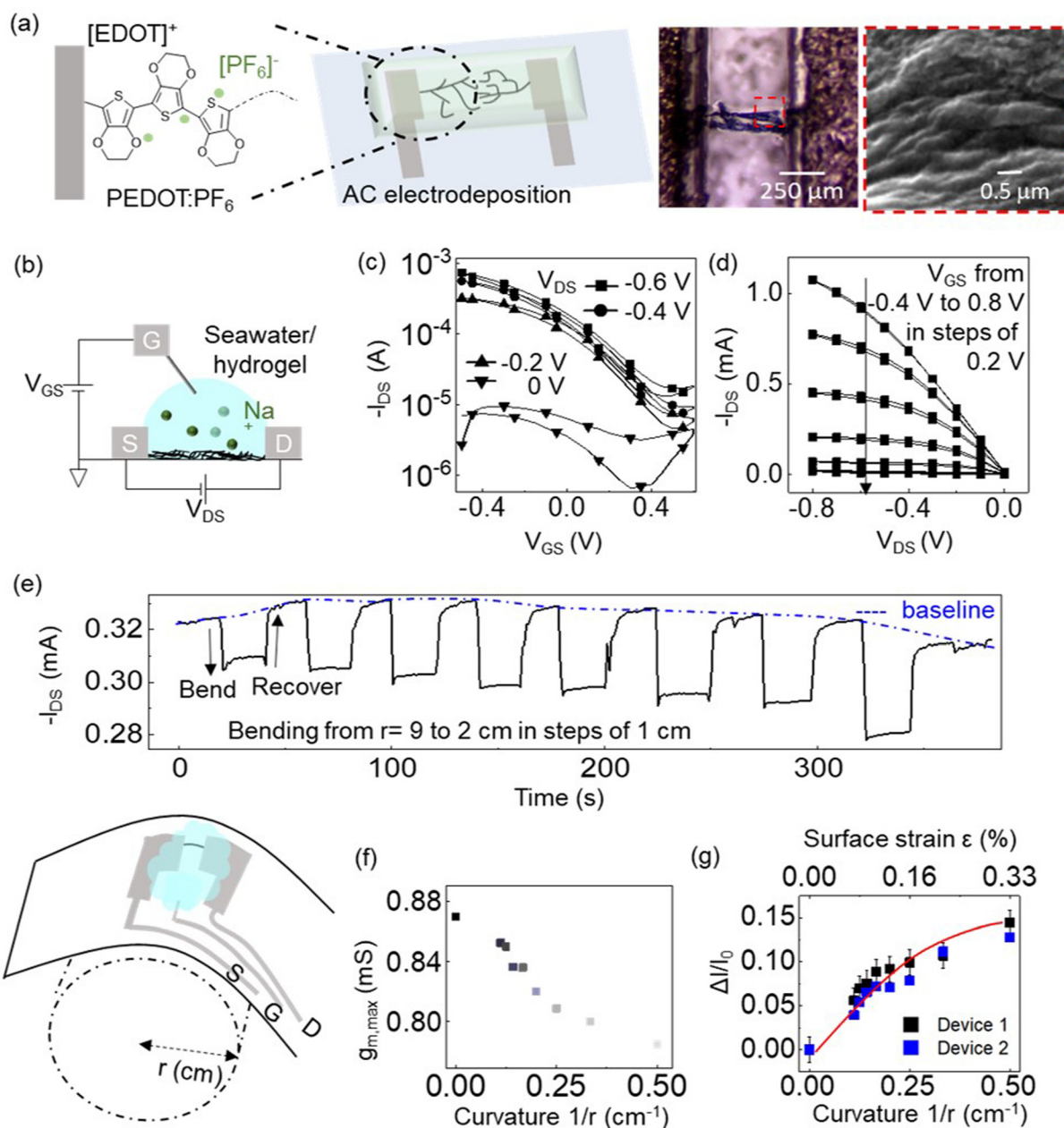


Fig. 3 (a) Schematics of PEDOT:PF₆ formed by AC electrodeposition. Optical microscope and scanning electron microscopy image of PEDOT:PF₆ fibers. (b) Schematics of an organic electrochemical transistor with seawater or hydrogel as the electrolyte. (c) Transfer and (d) output characteristic of a PEDOT:PF₆ electrochemical transistor in the flat state. (e) Source–drain current versus time under various bending radii, with an illustration showing the experimental configuration for calibrating the curvature sensor. (f) Transconductance and (g) ratio of current change as a function of bending curvature. Red line indicates a fit to eqn (1). Error bars accounted for the uncertainty level in the baseline drift. For parts (e) and (g), V_{DS} = -0.5 V, V_{GS} = -0.2 V.

$-I_{DS}(r)$ divided by the baseline I_{DS} , where $I_{DS}(0) = I_0$. The data $\Delta I/I_0$ versus curvature $1/r$ was fitted to an empirical exponential equation

$$y = -a \exp\left(-\frac{x}{b}\right) + c, \quad (1)$$

where fit parameters $a = 0.153$, $b = 0.228 \text{ cm}^{-1}$, $c = 0.155$, with an excellent coefficient of determination $R^2 = 0.97$. The

gauge factor GF indicating the device sensitivity was calculated by²⁸

$$\frac{\Delta I}{I_0} = \text{GF} \times \epsilon \text{ with } \epsilon = \frac{d}{2r}, \quad (2)$$

where ϵ is the surface strain and d is the substrate thickness equivalent to 130 μm in this work. The sensor GF was ~35 when the radius of curvature was greater than 5 cm ($1/r <$

0.2 cm^{-1}), and then GF dropped to 20 with bending radius below 5 cm. While this GF is not the state of the art, it falls within the range of most OECT strain sensors, and this unencapsulated device provided the additional advantage of stable operation in seawater. The OECT curvature sensor also delivered a uniaxial strain response, with much lower resistance change when the applied strain was perpendicular to the fiber length as seen in ESI Fig. S4e.† This result indicated the benefit of fiber alignment to the bending strain direction, maximizing the effect of structural deformation to enhance GF. In the following demonstration, because the oyster shell was typically 4–5 cm thick, the curvature measurement centered around $r = 2.5 \text{ cm}$ and the sensor GF was typically 20 in this work.

II.B. Salinity and dissolved oxygen (DO) sensors based on amperometry

The salinity of seawater is proportional to the amount of sodium chloride dissolved in the sample at a constant temperature and can be determined from ionic conductivity measurements. Here the salinity sensor detected the ionic current I_{salinity} through seawater samples using graphite electrodes (WE and CE in Fig. 1) with a bias of -0.2 V between them. The data in Fig. 4 was collected through the multiplexer

controlling the potentiostat. In Fig. 4a as more salt was added to the sample, the change in I_{salinity} was recorded simultaneously alongside a commercial multimeter (Hanna model #HI98194). The calibration of I_{salinity} versus salinity readings from the commercial meter is shown in Fig. 4b at the temperature of $23 \text{ }^\circ\text{C}$. The sensitivity was $0.013 \mu\text{A cm}^{-2}$ per g kg^{-1} . The measured noise standard deviation in our multiplexed system was $\sigma = 3.5 \text{ nA}$ (ESI Fig. S5†). Based on the signal-to-noise ratio, the salinity sensor showed a detection limit of 4.3 g kg^{-1} [detection limit = $3.3 \times \sigma/S$, where is the noise standard deviation = 3.5 nA and S is the sensitivity = $2.68 \text{ nA per g kg}^{-1}$]. Typical salinity in the ocean³⁹ is $33\text{--}37 \text{ g kg}^{-1}$, although oysters live in shallow coastal water where the salinity can be lower. Our sensor can cover the relevant range and was demonstrated to work from $4\text{--}40 \text{ g kg}^{-1}$.

For the DO sensor, since the detection mechanism was through oxygen reduction reactions (ORRs), the working electrode was electrodeposited with platinum Pt particles, in order for the Pt to catalyze the ORRs on the graphite electrode. The reaction current I_{DO} was measured under a potential of -0.8 V with respect to the reference electrode of activated carbon. The change in I_{DO} was recorded as the DO concentration was adjusted by gas purging, with simultaneous measurements by the commercial multimeter as seen in Fig. 4c. While adjusting

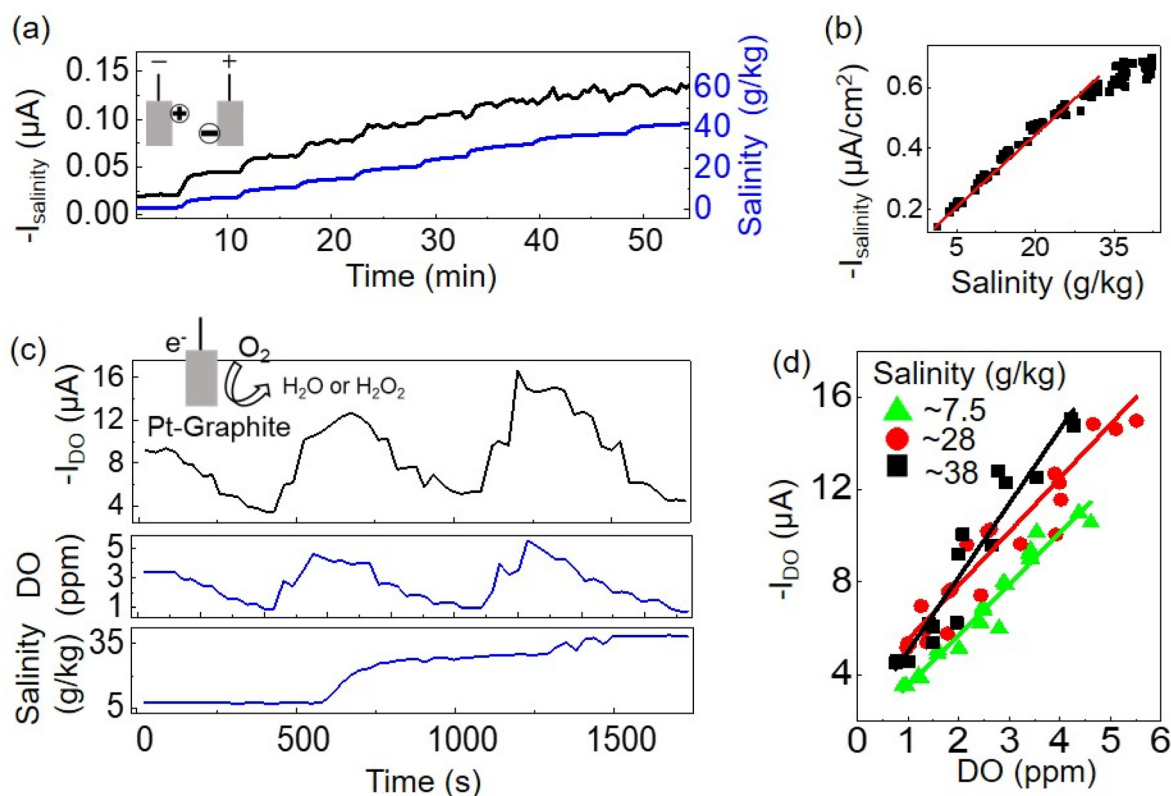


Fig. 4 (a) Current measurement by the printed salinity sensor biased at -0.2 V (black) and readings from the commercial meter (blue) as the sample salinity was increased over time. The inset illustrates the ionic current between sensor electrodes. (b) Calibration characteristics showing the sensor current density versus salinity. (c) Current measurement by printed dissolved oxygen sensor biased at -0.8 V (black) and readings from the commercial meter (blue), as oxygen concentration and salinity were adjusted. The inset illustrates oxygen reduction reaction at the sensor working electrode. (d) Calibration characteristics showing the sensor current versus dissolved oxygen concentration at three levels of salinity.

DO, we checked the effect of salinity on DO sensor measurements. The sensitivity of DO sensor was $2.42 \mu\text{A ppm}^{-1}$ in Fig. 4d, with no significant interference from changing salinity between $7.5\text{--}28 \text{ g kg}^{-1}$. Only when salinity exceeded 38 g kg^{-1} did the DO sensor response slightly increase due to higher background ionic current. The detection limit for this amperometric DO sensor was 0.5 ppm . The DO concentration lower than 2 ppm has often been defined as hypoxic,¹⁵ and the sensor here would be sufficient to monitor even severe hypoxia ($\text{DO} < 0.6 \text{ ppm}$).

II.C. Using the integrated sensor tag to monitor environmental effects on oyster gape behavior

In Fig. 5a, the printed sensor tag was attached to an oyster in a laboratory tank, with simultaneous video recording of the oyster movement to validate the curvature sensor readout. The DO level in the tank was adjusted by purging with gas, and the presented data excluded the gas-purging periods to avoid arti-

facts from gas turbulence. The oyster was active in opening and closing its gape when the DO level was above 3 ppm , as indicated by the increasing fluctuation frequency and amplitude detected by the curvature sensor in Fig. 5b. This behavior is also clearly observable in the ESI Video V1.† Such effect of DO on gaping behavior has been reported in prior work^{15,40} showing that oysters shut their gape under hypoxic conditions to lower their metabolism. In comparison to video-recording, the curvature sensor consumed much less power and can operate in the dark so as to not disturb the circadian cycle of the organisms.

Another observation period of over 80 minutes using the integrated sensor tag is shown in Fig. 5c, with the DO, salinity, and curvature measurements carried out in parallel. Fig. 5c presents the simultaneous real-time outputs of all three sensors, with the black data lines from each sensor (top: curvature sensor; middle: dissolved oxygen sensor; bottom: salinity sensor), and the blue data lines were from the commercial

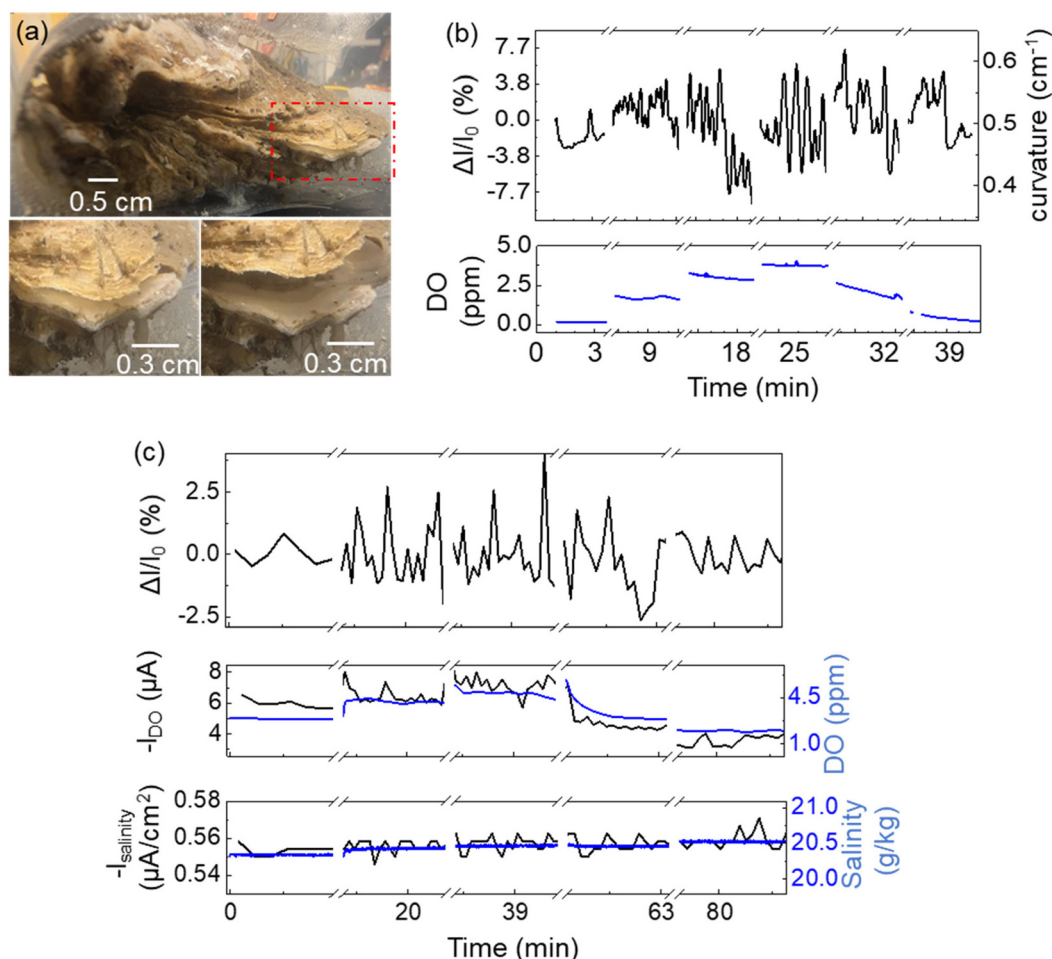


Fig. 5 (a) Photographs of oyster gape movement. (b) Changes in curvature tracking the oyster gape under different dissolved oxygen levels with data sampling at 1 Hz . (c) Measurements by the integrated sensor tag to monitor gape responses to environmental parameters of dissolved oxygen concentration and salinity, sampling at 0.016 Hz . The applied bias on OECT was $V_{\text{GS}} = -0.2 \text{ V}$ and $V_{\text{DS}} = -0.5 \text{ V}$, while the salinity sensor was biased at -0.2 V and DO sensor at -0.8 V . The data in blue represent measurements taken with the commercial multimeter to confirm the accuracy of printed sensors.

Table 1 Performance summary of sensors in this work

Sensor	Detection accuracy	Detection limit	Detection range	Sensitivity	Device size
Curvature	$\pm 0.5\%$	1 cm^{-1}	$0.11\text{--}0.5 \text{ cm}^{-1}$	GF of 35 to 20	0.3 cm^2
Salinity	$\pm 0.7 \text{ g kg}^{-1}$	4.3 g kg^{-1}	$4\text{--}40 \text{ g kg}^{-1}$	$0.013 \mu\text{A cm}^{-2}$ per g kg^{-1}	0.2 cm^2
Dissolved oxygen	$\pm 0.2 \text{ ppm}$	0.5 ppm	$0.5\text{--}6 \text{ ppm}$	$2.42 \mu\text{A ppm}^{-1}$	0.32 cm^2

sensors nearby. The recordings were taken through the MUX switching with the timing sequence as shown in Fig. 2. Like the earlier trial, higher DO in the environment led to more oyster gape movement. The salinity level was not adjusted in this trial, and the salinity readings remained constant and unaffected by gas purging.

The readout channel for the curvature sensor had a noise level of $0.01 \mu\text{A}$ with the current signal centered at $52 \mu\text{A}$. From this ratio of $I_{\text{noise}}/I_0 = 0.01/52 = 2 \times 10^{-4}$, the noise around the curvature measurement was calculated by using eqn (1) to be $1/r \pm 0.002 \text{ cm}^{-1}$. That is, for a case in which the measured $\Delta I/I_0 = 0.136$ corresponding to a curvature of 0.477 cm^{-1} , the curvature accounting for noise uncertainty was $0.477 \pm 0.002 \text{ cm}^{-1}$. Thus, the upper bound of curvature $1/r$ was 0.479 cm^{-1} and lower bound was 0.475 cm^{-1} , translating to radii of 2.088 mm and 2.105 mm , respectively. The curvature sensor was capable of distinguishing gape displacement down to sub-mm levels. While the long-term stability of the sensor tag will be assessed in future studies, the materials in this work have exhibited promising stability in other week-long studies in simulated marine environment¹⁰ or in saline culture medium.⁴¹

III. Conclusions

With the sensing performance summarized in Table 1, this work demonstrated the feasibility of monitoring multiple parameters through an integrated sensor system that is low-cost and enables *in situ* monitoring for bivalve aquaculture research. In addition to salinity and DO detection, the multiplexed setup offers the potential to easily add sensors for other environmental parameters such as pH and temperature. The sensor tag could be improved by integrating a temperature sensor that tracks the local variations in temperature and thus compensate the change in sensor outputs due to temperature effects. The curvature sensor based on electrodeposited semiconducting fibers was mechanically robust and remarkably stable in seawater without the need for encapsulation. The small, non-intrusive sensor tags do not take up much space and in the future can be adapted for field use and attached to multiple bivalves to facilitate group behavior studies. Upon accounting for the statistical variations in the sensors, if a high percentage of bivalves close their gapes, that might be a prominent indicator of severe environmental conditions. Overall, this report shared the design and fabrication processes to implement flexible electronics as compact data collection tools in marine studies.

IV. Experimental procedures

IV.A. Materials

The substrates were poly(ethylene terephthalate) (PET) of 5 mil ($\sim 130 \mu\text{m}$) thickness with lamination adhesive on the surface and were purchased from Immuson. Graphite foils were purchased from Emitac New Material Technology ($16\,000 \text{ S cm}^{-1}$ in conductivity, $16 \mu\text{m}$ thickness). A platinized carbon cloth was purchased from FuelCellStore (0.3 mg cm^{-2} , 40% Pt). Instant super glue for attaching the sensor tag to the bivalve shell was purchased from Gorilla Glue. The other chemicals were purchased from Sigma-Aldrich and used as received. The preparation procedures of hydrogel electrolyte and activated carbon (AcC) slurry were based on our previous work.¹⁰ In brief, the hydrogel electrolyte was a mixture of 1 g xanthan gum powder with 10 mL of simulated seawater (instant ocean salt, 31 mg in 1 L of deionized water). The AcC slurry was prepared by mixing activated carbon powder (Kuraray Chemical YP50F) and polyvinylidene fluoride in 9:1 ratio, with *n*-methyl-2-pyrrolidone as the slurry solvent.

IV.B. Device fabrication

The design of concentric-ring electrodes for amperometric DO and salinity sensors minimized the device footprint. Each ring was separated by a gap of $125 \mu\text{m}$ from neighboring rings. The electrode patterns in ESI Fig. S1† were cut from graphite foils by a digital blade-cutter (Silhouette Cameo). The graphite cut-outs were laminated onto the PET substrate by pressing them together at $150 \text{ }^\circ\text{C}$ for ten seconds. Then AcC was deposited onto the surface of the reference electrode by stencil printing to increase electric double layer capacitance in the reference electrode for DO and salinity sensors. The gate electrode of the OECT was modified by stencil printing Ag/AgCl ink (Ercon part #E2414),⁴² which reduced the series resistance of the gate for an applied voltage to drop mainly across the electrolyte and semiconductor channel. The stencil-printed materials were dried at $85 \text{ }^\circ\text{C}$ for 20 minutes.

Subsequently, two electrodeposition processes were carried out, one on the working electrode of the DO sensor and the other on the OECT to form semiconducting fibers for the transistor channel. For the DO sensor, the reaction catalyst was platinum, and the working electrode surface was modified by electrodepositing platinum nanoparticles from a platinized carbon cloth (40%, Pt-C) through a linear sweep voltammetry scan, applying 1.2 to 0.8 V for five cycles scanning at -50 mV s^{-1} on the platinized carbon cloth.¹⁰ Electrodeposition of semiconducting fibers to bridge the OECT channel ($250 \mu\text{m}$ in length) was done through a solution

of 10 mM 3,4-ethylenedioxythiophene (EDOT) and 1 mM tetrabutylammonium hexafluorophosphate (TBAPF₆) in propylene carbonate solvent. The deposition voltage was a triangular waveform of ± 10 V switching at 12.5 Hz, applied between the two graphite source and electrodes. The deposition time was around 10 minutes until the deposition current reached saturation. Finally, the graphite interconnects were connected to aluminum wires through silver paste. The bonding area between the sensor tag and the aluminum wires was encapsulated by silicone elastomer (Ecoflex) to avoid contact between the wires and seawater during measurements.

IV.C. Multiplexer circuit and wiring

The circuit components included four 8-to-1 multiplexer chips (74HC4051) and the Nano-Arduino purchased from Sparkfun. The multiplexer channels were controlled by Nano-Arduino through three select-lines (S0, S1, and S2, connected to the Arduino analog outputs of A0, A1, and A2). The outputs of multiplexers were connected to a portable potentiostat (Metrohm μ 300) for data acquisition. The 5 V used to power Nano-Arduino and multiplexers were supplied by computer. The multiplexer circuit was soldered on a small piece of gold-plated printed circuit board. The multiplexers served as digital switches between channels, controlled by a custom Arduino program shown in ESI Fig. S7.†

IV.D. Device characterization

Electrical measurements were taken with electrometers (Keithley 2000) controlled by LabVIEW software or with a portable potentiostat (Metrohm μ 300) controlled by DropView 8400. Cyclic voltammetry, linear sweep voltammetry, and electrochemical impedance spectroscopy were carried out using a benchtop potentiostat (BioLogic SP-200). Scanning electron microscopy images were taken with FEI microscope operating at 5 kV. In the bending measurements, the device curvature was adjusted by laying it on a printed mold with various radii of curvature (ESI Fig. S7†); the mold was fabricated using the printer Raise3D N2 platform.⁴³ The concentration of dissolved oxygen was adjusted by purging oxygen or nitrogen and monitored by a multimeter (Hanna model #HI98194) in real-time.

Conflicts of interest

The authors declare no conflict of interest.

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