An Iontronic Multiplexer Based on Spatiotemporal Dynamics of Multiterminal Organic Electrochemical Transistors

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The seamless integration of electronics with biology requires new bio-inspired approaches that, analogously to nature, rely on the presence of electrolytes for signal multiplexing. On the contrary, conventional multiplexing schemes mostly rely on electronic carriers and require peripheral circuitry for their implementation, which imposes severe limitations toward their adoption in bio-applications. Here, a bio-inspired iontronic multiplexer based on spatiotemporal dynamics of organic electrochemical transistors (OECTs), with an electrolyte as the shared medium of communication, is shown. The iontronic system discriminates locally random-access events with no need of peripheral circuitry or address assignment, thus deceasing significantly the integration complexity. The form factors of OECTs that allow for intimate biointerfacing as well as the electrochemical nature of the communication medium, open new avenues for unconventional multiplexing in the emerging fields of bioelectronics, wearables, and neuromorphic computing or sensing.

1. Introduction

The integration of electronics with biology is triggering significant attention in several fields, including medical diagnostics, personalized and precise medicine, healthcare, bioelectronics, and biorobotics.[1,2] The technological potential of such co-integration stems from the ability of electronics to bi-directionally interact with biological systems, and even more importantly to emulate biological functions, thus, allowing seamless communication and integration.[3,4] Biological systems—including animals and plants—communicate and process information using as carriers ions, small molecules, and electronic charges.[5]

In the nervous system, stimuli are collected from distributed sensory receptors, computation then takes place locally or centrally, and when necessary, feedback signals drive sensorimotor processes.[6] Interestingly, within this loop, biological systems perform sensing, computing, and actuation by means of temporal and spatial multiplexing of information. For instance, nerve fibers in the peripheral nervous system carry and multiplex bio-signals from sensory neurons.[8] Even single neurons are able to discriminate via multiplexing, spatiotemporal sequences of incoming signals that are distributed across their dendrites.[7,8] At a higher level, brain oscillations are known to coordinate or even multiplex the flow of temporally asynchronous biological events,[9–11] but are not strictly considered to be global clocks as they are spatially distributed across the brain and diversify in the frequency domain.[10,11] Such spatiotemporal multiplexing paradigms in biological systems are not only inherently dependent on the structure of biological neural networks, but also on the surrounding medium, namely a global aqueous environment that in the simplest case is salt water.[3,13]

Although conventional multiplexing schemes are successful in numerous applications, it is apparent that seamless integration between electronic and biological systems requires new bio-inspired approaches that, analogously to nature, inherently rely on ionic–electronic (i.e., iontronic) materials and devices, or even system architectures. Inspired from biology and motivated by the need of simplicity and minimalism in multiplexing concepts, iontronic systems that exhibit spatiotemporal response have the potential for compact multiplexing of time-domain signals, as spatial specificity of signal sources is inherently embedded into their device dynamics. Toward this direction, organic electrochemical transistors (OECTs) are promising candidates for bio-inspired electronics because they show stable operation in electrolytes (both liquid or soft solid-state electrolytes), and owing to their volumetric ionic–electronic charge

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interaction taking place in the whole bulk of the polymer, OECTs offer an ideal platform for compact multiterminal configurations\textsuperscript{[38–43]} as well as facile and highly efficient biointerfacing.\textsuperscript{[43]} In addition, the ion transduction of OECTs combined with their neuromorphic properties,\textsuperscript{[21,41]} render them as an ideal technology platform for numerous applications including bioelectronics, neuro-inspired processing, and sensing.\textsuperscript{[21,41,44–46]}

On the contrary, in traditional electronic systems, electron conductive paths (e.g., metal lines) are used to connect the various parts of the system. Applications with restrictions in space, cost, and energy resources require the compact signal transmission between circuit components. For instance, when a multitude of elements is necessary (i.e., sensors, actuators, and memory), the most affordable structure is the cross-bar matrix integration. In this approach, an arbitrary number of $M \times N$ elements are organized in a device grid of $M$ rows and $N$ columns, which require $M + N$ metal lines for the interconnections. Prominent examples of matrix-like electronic multiplexing include displays (e.g., row/column addressing of lighting pixels),\textsuperscript{[14–16]} cross-bar arrays of analogue memories in neuromorphic computing (e.g., dot product or weighted multiplexing of rows across a column),\textsuperscript{[17–22]} electrode arrays for electrophysiological recording/stimulation, and physical (e.g., pressure, temperature, and light), chemical, and biological sensor arrays.\textsuperscript{[22–28]}

Many of these multiplexing approaches rely on extensive accessing/read-out peripheral circuitry and accurate timing/synchronization protocols that require significant computational resources. Another possible approach, when applicable, is based on the passive addressing where a matrix is solely made of devices with rectifying properties.\textsuperscript{[22]} Although of compact structure, this addressing scheme requires a concurrent combination of biasing conditions and therefore central clocks and controllers for coordination.\textsuperscript{[22,27,28]} In order to simplify the addressing conditions, methods based on frequency-division multiplexing in passive matrices have been successfully implemented in electronic bio-signal acquisition.\textsuperscript{[29]} However, signal reconstruction requires complex peripheral circuitry for phase detection. Active addressing necessitates access elements in close proximity to the various devices of the matrix, thus significantly increasing the integration complexity. Indeed, such cross-bar arrays minimize undesirable cross-talk/parasitic communication, but still require nonlinear access elements such as transistors or diodes for addressing a device or a whole row/column.\textsuperscript{[22]} Multiplexing with active addressing, in many cases still needs to be coordinated by central clocks.\textsuperscript{[22]} Pseudo-parallel access with high scan-rates of sequential row-by-row addressing (time-division multiplexing) relaxes partially the need for time coordination in biosensing and lighting display applications, but still high-speed peripheral electronics are necessary.\textsuperscript{[15,16,23]}

This pseudo-parallel method requires over-scanning with high-speed rates and, as a consequence, it is power-hungry (with idle power consumption), especially if employed for temporally and spatially sparse or even asynchronous input sources and signals found in biological neural activities and sensory systems.\textsuperscript{[30–31]}

An alternative approach is the event-based signaling—loosely considered as a form of code-division multiplexing—where inputs are encoded and transmitted through a shared medium only when it is necessary, meaning asynchronous or clock-less transmission.\textsuperscript{[34–37]} Therefore, event-based signaling is an energy efficient multiplexing scheme when dealing with sparsity, and has been successfully incorporated in e-skins based on soft electronics for the emulation of tactile sensing.\textsuperscript{[23]} This approach allows for asynchronous communication between sensing and processing modalities by multiplexing input signals via a common conductor.\textsuperscript{[23]} Nevertheless, it is still a multiplexing scheme of address encoding with signal reconstruction that requires off-chip decoding, and the use of the common conductor as a signal collector also implies the absence of spatial specificity of a signal source. On the contrary, spatial specificity is inherently embedded in the dynamics of iontronic devices and systems.

Here, we show a bio-inspired iontronic multiplexer based on spatiotemporal dynamics of multiterminal OECTs. Analogously to biological systems, an electrolyte serves as the shared medium of communication and owing to the spatial dependence of the ionic–electronic coupling, the proposed system is able to discriminate locally random-access events at the input terminals with no need of address encoding or specialized peripheral circuitry. In this way, we demonstrate the multiplexing of 25 signals by means of a simple polymeric channel. The system is modeled by using a physically based circuit-aware approach and the model is implemented in an industrial-standard circuit simulator. The simulations are able to predict the system behavior, providing insight on the key system parameters for spatiotemporal dynamics and the model represents a universal platform for a greater family of materials. The proposed approach opens new avenues for minimalistic, ion-based multiplexing, when circuitry simplicity for signal transfer is essential, including applications in the emerging fields of bioelectronics, wearable electronics, neuromorphic computing, and sensing.

2. Results and Discussion

A schematic of the iontronic multiplexer is shown in Figure 1a. The system consists of a channel made of the conducting polymer blend poly(3,4-ethylenedioxythiophene) doped with poly(styrene sulfonate) (PEDOT:PSS). PEDOT:PSS is a high-performance p-type conductive polymer, consisting of a hole conductive PEDOT blended with the ion conductive PSS.\textsuperscript{[43,47]} Therefore, PEDOT:PSS supports both electronic and ionic transport and is termed as organic mixed ionic–electronic conductor.\textsuperscript{[47]} When a PEDOT:PSS film is in contact with an electrolyte, ions can permeate the polyelectrolyte phase and ionic–electronic charge compensation takes place through the whole bulk volume of the polymer.\textsuperscript{[48]}

As a result, the electronic (hole) charge density into the PEDOT semiconducting phase of the polymer is electrochemically modulated by injecting ionic carriers from/to the electrolyte into the PSS phase of the polymer, depending on the voltage of the gate electrode. This ionic-to-electronic current modulation constitutes the principle of operation of OECTs.\textsuperscript{[43,49]} Here, the PEDOT:PSS channel is electrostatically coupled via an electrolyte (NaCl at various concentrations) with a grid of $5 \times 5$ electrodes which acts as gates. Therefore, the iontronic multiplexer consists of a multiterminal OECT. The gate electrodes are made of gold (Au) covered with PEDOT:PSS providing electronic-to-ionic signal conversion. The
device microfabrication process is described with more details in the Experimental Section. In this system configuration, input voltage signals ($I_n$) are applied at a gate electrode, $G_n$, while the output current ($O$) is measured at the drain electrode of the multigate OECT. It is worth noting that in a real biointerfacing or biological system the gate could be replaced by a distributed sensor or a local bio-signal generator entity. The shared electrolyte between the gates, $G_n$, and the channel allows for “parallel” coupling of the channel with the various gates, $G_{n}$. An actual microfabricated device is displayed in Figure 1b.

A subsection of the multiterminal OECT comprising a gate, $G_n$, and the channel is depicted at Figure 1c. When an input voltage signal is applied at the gate electrode, ions drift from the bulk of the electrolyte toward $G_n$ and the channel. More in detail, a positive bias at the gate leads to the drift of cations via lateral gating into the channel. Cations compensate the sulfonic acid groups of PSS$^-$ (fixed anions) close to the PSS/PEDOT interface and, in turn, leads to a decrease of the hole density into the PEDOT. The opposite situation is observed at the $G_n$ gate electrode that is also covered with PEDOT:PSS. The ions flowing into the electrolyte can be described by an ionic resistor which has an intrinsic property of the electrolyte, while the area and length ($d$) of the resistor can be selected by design. As a result, the electrostatic ionic–electronic coupling of the gate–electrolyte-channel can be described by the simplified circuit diagram showed in Figure 1c. The volumetric electric-double-layer capacitance at the gate is described with a capacitor $C_{Gn}$ and the resistor $R_{Gn}$ connected on parallel expresses any electronic or ionic loss mechanism (in Au/PEDOT:PSS electrodes $R_{Gn}$ is usually high) and could be omitted. The drift of ions into the electrolyte is described by a distance-dependent resistor $R(d)$, and the ionic–electronic coupling between the electrolyte and the channel is described with a volumetric transistor. Owing to the spatial distribution of the gates in respect to the channel, the electrolyte resistance depends on the input–output distance, $d$, between $G_n$ and the channel. Hence, $R(d)$ in the multiterminal OECT system configuration is a variable (distance dependent) resistor. Owing to the common nature of the electrolyte, this ionic–electronic coupling holds in parallel for every $G_n$–channel pair of the multiterminal OECT.

The device geometric configuration leading to spatiotemporal response is investigated in Figure 2. The electrochemical parameters of the system are measured by probing the impedance spectrum (IS) of the multiterminal OECT device with channel of variable-sized active area. In Figure 2a,b, a large channel device with an area ($W \times L$) of $500 \times 15000 \mu m^2$ is investigated, and in Figure 2c,d, a small channel device with an
demonstrate that the IS is dominated by the electrolyte resistance $R$, which increases by increasing the gate–channel distance (Figure S2, Supporting Information). We found that in large channel devices the maximum relative variation amounts to $\Delta R/R_{\text{MIN}} = 560\%$. In contrast, in the case of small channel devices, Figure 2c shows that the IS spectra are almost perfectly overlapping for all $G_n$–channel pairs ($n = 1–25$), and the corresponding spatial mappings of $R$, $C$, and $\tau$ are only weakly dependent on the spatial device configuration. In more detail, Figure 2d shows that the maximum relative variation of is: $\Delta R/R_{\text{MIN}} < 20\%$, $\Delta C/C_{\text{MIN}} < 10\%$, and $\Delta \tau/\tau_{\text{MIN}} < 20\%$, respectively. The measured values are displayed in Figure S2, Supporting Information. In both cases of small and large channels, C is practically independent of the $G_n$ positions ($\Delta C/C_{\text{MIN}} < 10\%$). This is attributed to the fact that interfacial capacitance $C$ corresponds to the low frequency regime of the spectrum, therefore it is defined by the interfaces (practically the same interfaces for different $G_n$–channel couples) and not by the electrolyte. The response time, $\tau$, follows the trend of $R$. Comparing the measured $R$ and $C$ displayed in Figure S2b,d, Supporting Information, we found that the ionic resistance $R$ of the small channel system is about one order of magnitude larger than $R$ measured in the large channel system. This explains the limited relative variation, $\Delta R/R_{\text{MIN}}$, in the case of the small channel system, and indicates that the ionic resistance significantly depends on the geometrical properties of the channel. This experimental analysis clearly shows that the dimension of the polymeric channel is a key system parameter, since it significantly impacts on both the gate–channel ionic resistance and the channel capacitance itself. Importantly, the proper sizing of the channel enables the enhancement of the spatiotemporal response of the system while the electrolyte connectivity medium tunes the spatiotemporal response.

The scaling behavior of the ionic resistance $R$ as a function of the $G_n$–to-channel distance $d$ (central column of the $5 \times 5$ gate grid) is presented at Figure 3a, in the case of the large channel and small channel as reference. The spatial scaling of the ionic resistor is evident for the large channel, while the effect is negligible for the small channel configuration. Figure 3b focuses on the large channel, displaying the spatial scaling of $R$ for various electrolyte concentrations ($c = 5–500$ mm NaCl). As expected, $R$ scales linearly with $d$ for all ionic concentrations, $R \propto d$. Despite the fact that the two interfaces, $G_n$/electrolyte and electrolyte/channel are lateral (integrated on the very same substrate), with different areas, and with a “virtual” cross section of the ionic conductor (i.e., the electrolyte) to be loosely defined, linear scaling indicates an exemplar Ohm’s law behavior. The configuration of the multiterminal OECT device with the $5 \times 5$ $G_n$ grid, suggests the formation of 25 “virtual” and parallel wires between the $G_n$–channel terminals via the electrolyte medium. In addition, it is worth noting that also all the various gates are connected (25 $\times$ 25 connections) through the electrolyte. For evaluating the linear scaling behavior in Figure 3b, a modified formulation of a conductor with variable cross-sectional area, $A_1$ to $A_2$, is used ($R = \rho \cdot d/\sqrt{A_1 A_2}$, refer to the Supporting Information), with $R_0$ the residual resistance obtained when $d = 0$. The resistivity $\rho$ of the “virtual” wire is calculated for $c = 100$ mm NaCl and amounts to $\rho = 8.1 \text{ Ohm} \cdot \text{cm}$. It should
Figure 3. Spatial scaling of the connectivity medium. a) Ionic resistance $R$ ($X = 3$, $Y = 1–5$) as a function of the gate-to-channel distance, $d$, for the large and small channel ($c = 100 \text{ mm NaCl}$). The dependence of $R$ on $d$ is much stronger in the case of the large channel. b) Ionic resistance, $R$, as a function of $d$ for $c = 5–500 \text{ mm NaCl}$, in the case of the large channel. $R$ scales linearly with $d$ for all concentrations suggesting a simple Ohm’s law behavior (here shown in logarithmic scale for clarity). c) Maximum resistance difference $\Delta R$ of the central gate column between the most distant and the closest gate electrode as a function of $c$. Spatiotemporal response and connectivity medium. Spatial mapping of the relative variation of $R$, $C$, and $\tau$ for all $G_n$–channel pairs and for $c = 5–500 \text{ mm NaCl}$. d) $c = 5 \text{ mm NaCl}$. e) $c = 500 \text{ mm NaCl}$. Large channel device: channel dimensions ($W \times L = 500 \times 15000 \mu\text{m}^2$), gate dimensions: $(A_G = 2000 \times 2000 \mu\text{m}^2)$. Small channel device: channel dimensions ($W \times L = 20 \times 20 \mu\text{m}^2$), gate dimensions $(A_G = 2000 \times 2000 \mu\text{m}^2)$.

be noted that the calculated value is an effective resistivity for the designed system and it cannot be directly related to the intrinsic electrolyte resistivity ($100 \text{ Ohm-cm}$ for $c = 100 \text{ mm NaCl}$), as the system consists of in-plane interfaces; a situation that differs from an ideal parallel plate capacitor. Moreover, the analysis provides the residual resistance which amounts to $R_0 = 68 \text{ Ohm}$.

The electrolyte serves as connectivity medium and it can tune the spatiotemporal response of the multiterminal system when the system is suitably designed. The maximum variation of the electrolyte resistance $\Delta R$ across the $Y$-direction as a function of the electrolyte concentration, $c$, is shown in Figure 3c. The measurements show that for highly conductive connectivity media that approach metal conductivity, the spatiotemporal response of the system is heavily suppressed. More specifically, Figure 3d,e shows the spatial mappings of relative variation of $R$, $C$, and $\tau$ resulting from the equivalent circuit analysis of the IS spectrum for all $G_n$–channel pairs, for two cases of electrolyte concentration (low: $c = 5 \text{ mm}$, high: $c = 500 \text{ mm NaCl}$). As expected, in both cases the relative variation of the volumetric capacitance ($\Delta C/C_{\text{MIN}}$) across the device footprint is minor, as it primarily depends on the low frequency IS response of a $G_n$–channel pair. In contrast, the relative variation of the ionic resistance $\Delta R/R_{\text{MIN}}$ (and therefore of the corresponding $\tau$) is more pronounced for lower ionic concentrations of the electrolyte. This can be explained as follows. At low ion concentration, the ionic resistance dominates in a wide range of the IS compared to the volumetric capacitance, and the spatiotemporal dynamics of the multiterminal system are emerging. Figure 3d,e demonstrates clearly that spatiotemporal phenomena are more pronounced for less-conductive connectivity media and in contrast with current approaches, spatial specificity (sensory) signal sources can be introduced locally by collecting signals via media of relatively low ionic conductivity.

By closely observing representative IS spectra of the multiterminal OECT for the $G_n$–channel pairs, specific frequency-dependent modes of operation can be identified (Figure 4). Convectional current–voltage ($I–V$) curves with slow sweep rates ($<200 \text{ mV s}^{-1}$, $f_{\text{eff}} < 0.3 \text{ Hz}$), such as output characteristics of the multiterminal device (drain current vs gate voltage, $I_D$ versus $V_G$), are effective low-frequency measurements. On the contrary, pulsing the $G_n$ gate(s) effectively probes the high-frequency device dynamics (for a pulse width $\leq 100 \text{ ms}$, $f_{\text{eff}} \geq 10 \text{ Hz}$) in the $I_D$ versus time response. Under such temporal conditioning, the response of the multiterminal OECT displays readily observable spatiotemporal response. In the case of the large channel (Figure 4a), the IS spectra for three representative $G_n$–channel pairs (small, medium, and far
Figure 4. Frequency-dependent modes of operation. a) Large channel. IZI spectrum for three gate–channel pairs: close distance (X = 3, Y = 1), medium distance (X = 3, Y = 3), and far away distance (X = 3, Y = 5). IZI versus f merges into a single curve for low frequencies f, while it is increasing with the Gsc–channel distance at higher f. Low f (quasi-static) output characteristics I0 versus V0, and high f pulsing response (I0 vs time) of the multiterminal OECT. Low f, I0 versus V0 characteristics are independent on the Gsc–to-channel distance, while the high f pulsing response (I0 vs time) depends on the Gsc–to-channel distance. In the latter regime of operation, the system exhibits spatiotemporal properties. b) Small channel. IZI versus f is independent of the spatial configuration of the multiterminal system. The electrical characteristics (I0 vs V0, and pulsing response) are independent of the Gsc–to-channel distance. For low f I–V measurements, step delay time for voltage sweep is equal to 2 s. The high f pulsing measurements parameters are the following: gate pulse voltage amplitude = 500 mV, time width = 100 ms, time between pulses = 5000 ms. V0 = 0 mV, V0 = −100 mV. Large channel device: channel dimensions (W × L = 500 × 15 000 µm²), gate dimensions: (A0 = 2000 × 2000 µm²). Small channel device: channel dimensions (W × L = 20 × 20 µm²), gate dimensions (A0 = 2000 × 2000 µm²).

distance between Gsc and channel) are converging in the low frequency regime, and the (low frequency) I–V characteristics are independent of the spatial device arrangement. In the high-frequency regime, however, the IS spectra are divergent and the high-frequency pulsing response obtains spatial properties. For a small channel (Figure 4b), the IS spectra merge into a single curve, and the corresponding low- (I–V) and high-frequency response (pulsing) is practically independent of the spatial configuration of the system. It is clear from Figure 4 that the system geometry and its probing conditions define the emergence of spatiotemporal dynamics. It should also be noted that the transconductance g m of the large and small OECT is g m(l) = 0.13 mS and g m(s) = 3.6 mS, respectively, and thus g m(l)/g m(s) = 0.036. The small g m obtained in the case of the large OECT is due to the OECT geometries. Indeed, g m is proportional to the transistor form factor, W/L. The dimensions of the large OECT device are W1 = 500 µm and L1 = 15 000 µm, which result in a form factor, W1/L1 = 0.033 while the dimensions of the small OECT are W2 = 20 µm and L2 = 20 µm, which result in a form factor, W2/L2 = 1. The reduction of the form factor from 1 to 0.033 quantitatively supports the reduced g m. The key OECT parameter affecting the spatiotemporal response are the OECT dimensions. More in detail, i) the polymer should extend from the first to the last columns of the gate matrix in order to locally convert ions to electronic charges that are then transported into the semiconducting electronic phase of the polymer. ii) The polymer area that, in turn, define the OECT capacitance Ce = CV × W × L × t, where CV is the volumetric capacitance—an intrinsic property of the polymer—and t the polymer thickness.

To gain more insight on the key design parameters, the system is modeled by considering the various components such as the Gsc gate electrodes, the electrolyte, and the OECT channel. The system is described by using a modular circuit approach displayed in Figure S3, Supporting Information. A single column of gates is considered and a Randles circuit (Csc in parallel to a resistance Rsc, with Csc and Rsc the gate capacitance and resistance) extracted from IS, is used to model each gate. The vertical gate-to-gate and gate-to-channel coupling is described by the ionic resistance RYY. Similarly, the horizontal gate-to-gate coupling is described by RXX. RYY and RXX are experimentally accessed by varying the distance between the gates, the electrolyte concentration, and geometries. Finally, the OECT channel is modeled as a transistor including a Randles circuit extracted from IS measurements, namely a capacitance, CCh, in parallel to a resistance, RCh. The CCh and RCh elements are partitioned at the source (C0 and R0) and drain (C0 and R0) side of the channel. In the case of the long-channel system, the PEDOT:PSS channel is divided into five subchannels (Lsc = L/5) which are spatially coupled to the Gsc gates and define a colunnar, OECTx. Figure S3, Supporting Information, also shows the circuit model of both the long- and short-channel systems.

The comparison of the fabricated system showed in Figure 1b with the circuit model, provides direct evidence of the role of the electrolyte and ionic–electronic interactions, which results in the various coupling elements. The spatiotemporal properties of the system depend on the balance between the gate capacitance, the distance between the various components (e.g., gate-to-gate and gate-to-channel), the electrolyte concentration, and the channel geometrical dimensions and capacitance. Focusing on the long-channel system, we used the model to investigate the impact of two key system parameters: the gate capacitance Cg and the vertical gate distance ΔY (here shown
Figure 5. Impact of the system parameters on the spatiotemporal response. Spatial mapping of the percentage peak current normalized to the global maximum \(\Delta I/\text{MAX} \) (%). Logarithmic scale is used. The gate capacitance \(C_G\), and distance between the gates along the vertical (Y) direction are varied (here shown as gate-to-gate resistance \(R_{YY}\)). By observing the subpanels across the horizontal direction, the vertical gate distance \(\Delta Y\) (or \(R_{YY}\)) is increasing, while moving across the subpanels in the vertical direction, \(C_G\) is increasing. a) \(C_G = 9.85 \times 10^{-6} \text{ F}, \Delta Y_1 = 175 + 7000Y \mu\text{m}\) (or \(R_{YY} = 10.3 \text{ Ohm}\), \(Y = 0\)). b) \(C_G = 9.85 \times 10^{-3} \text{ F}, \Delta Y_2 = 1750 + 7000Y \mu\text{m}\) (or \(R_{YY} = 103 \text{ Ohm}\)). c) \(C_G = 9.85 \times 10^{-3} \text{ F}, \Delta Y_3 = 17500 + 70000Y \mu\text{m}\) (or \(R_{YY} = 1030 \text{ Ohm}\)). d) \(C_G = 9.85 \times 10^{-6} \text{ F}, \Delta Y_1 = 175 + 7000Y \mu\text{m}\). e) \(C_G = 9.85 \times 10^{-6} \text{ F}, \Delta Y_2 = 1750 + 7000Y \mu\text{m}\), that is, parameters of the experimental systems. f) \(C_G = 9.85 \times 10^{-3} \text{ F}, \Delta Y_1 = 17500 + 70000Y \mu\text{m}\). g) \(C_G = 9.85 \times 10^{-3} \text{ F}, \Delta Y_2 = 17500 + 70000Y \mu\text{m}\). h) \(C_G = 9.85 \times 10^{-3} \text{ F}, \Delta Y_3 = 17500 + 70000Y \mu\text{m}\). i) \(C_G = 9.85 \times 10^{-3} \text{ F}, \Delta Y_4 = 17500 + 70000Y \mu\text{m}\). All the other geometrical and physical parameters are the same of those obtained from the experimental long-channel system, \(W = 500 \mu\text{m}, L = 15 \text{ mm} \mu\text{m}, \mu = 14 \times 10^{-5} \text{ A} \text{ V}^{-2}, \nu = 0.75 \text{ V}^{-1}\). In order to directly compare the various results, we calculated the percentage response of the output peak current, \(I_D/\text{MAX} \) (%), in respect to the global maximum peak, \(I_{\text{MAX}}\), obtained by applying the input signal at a \(G_n\) gate of the grid. The spatial mappings of \(I_D/\text{MAX} \) (%), by varying \(C_G\) and \(R_{YY}\), is shown in Figure 5. To highlight the impact of the parameters, both \(C_G\) and \(R_{YY}\) are increased/reduced by one order of magnitude with respect to the values obtained by modeling the experimental long-channel system while all the other system parameters are fixed (Table S1, Supporting Information).

In the limit of densely spaced gate rows and low gate capacitance, \(C_G\), (upper left diagonal subpanel[s] of Figure 5, or Figure 5a,b,d) \(I_D/\text{MAX} \) (% is independent on Y-direction. The system practically shows only very limited spatiotemporal response due to the ultralow response time across the Y-direction in respect to the probing conditions. In contrast, the spatiotemporal response across the X-direction is attributed to the polymeric channel. Ions transported through the electrolyte reach the channel and are converted into bulk electronic current. This is a spatial phenomenon that takes place along various positions of the channel length, depending on the horizontal position of the gate. The electronic charge carriers that originate closer to the source electrode are transported for longer distances with respect to the ones originated closer to the drain, where the electronic current is measured. The spatiotemporal response is emerging when the row-to-row distance \(\Delta Y\) is increasing (across the horizontal subpanel direction). In the limit of large \(\Delta Y\) distances (Figure 5c,f,k) the spatiotemporal dependence is diminished for distant rows, from which the channel appears as a point device. The spatiotemporal response across Y-direction can also be recovered by increasing the gate capacitance \(C_G\) (across the vertical subpanel direction), and thus by increasing the response time across this direction. The analysis shows that optimal conditions are achieved in the case of Figure 5e (fabricated system) where 2D spatiotemporal response is displayed by all \(G_n\) gates. The calculations of the Figure 5 show that the spatiotemporal response (viz. OECT current) reduces by increasing the gate–channel distance (parameter \(R_{YY}\)) and this can be tuned by changing the size of the most distanced electrodes. More in detail, for a given \(C_G\), the spatiotemporal response increases by increasing the vertical distance but the current obtained from the most distanced gates vanishes and cannot be practically detected anymore (see e.g., Figure 5c,f,k). By increasing
the spatiotemporal response can be enhanced but an optimum response depends on the gate distance. At small distances (Figure 5a,d,g) the maximum spatiotemporal response is achieved with the maximum \( C_G \), while at larger distances (Figure 5b,e,h) larger capacitances could result in a limited response of the most distant gates (Figure 5h) and the optimum is obtained when \( C_G = C_{G2}, C_{G2} = 98.5 \times 10^{-6} \text{F} \). Further increasing the gate-to-channel distance results in limited current response of about half of the gates and the increasing \( C_G \) is counter-productive (Figure 5c,f,k).

A deeper analysis on the impact of the channel length and the gate capacitance on the spatiotemporal response is shown in Figure S4, Supporting Information. The impact of the \( W/L \) ratio on the spatiotemporal response is also studied in Figure S4, Supporting Information. The spatiotemporal response depends on two key channel parameters, namely the channel length and the channel area. A long channel is essential to obtain a spatially distributed ion transduction along the channel, which result in a spatial time response. The channel area is directly related to the channel capacitance which affects the amount of ions flowing through the electrolyte upon the applied gate pulse. As a result, the channel length, \( L \), and the channel area, \( W \times L \), are important device parameters while the ratio, \( W/L \), impact the magnitude of the drain current but has no impact on the spatiotemporal response of the device. Overall, optimal spatiotemporal response requires a balanced design between the electronic-to-ionic conversion at the gates, the ionic transport through the electrolyte, the ionic-to-electronic conversion along the channel, and the electronic transport through the channel. As shown from Figure S4, Supporting Information, the multiterminal device can be further scaled down. Apart from its applicability in the proposed system, it is noteworthy that the model can extend the proposed concept in a variety of materials/geometries and can be used as a universal tool for design and validation.

According to the aforementioned analysis, the large channel providing the optimized spatiotemporal response is used for the demonstration of ionic–electronic signal multiplexing. In Figure 6, the gates, \( G_n \) (\( n = 1–25 \)), are randomly addressed by applying square voltage pulses (input signals) at each \( G_n \) sequentially. Details about the custom-made, accessing/measurement set-up are presented in the Supporting Information, and measurement conditions are described in Experimental Section. The amplitude of the output drain current \( I_D \) is also defined for every addressing event. The random accessing process is monitored and represented across the two directions of the \( 5 \times 5 G_n \) grid in Figure 6a. The amplitude of \( I_D \) is represented by scanning across the rows of the \( G_n \) grid (Y-direction, for \( Y = 1–5 \) or row #). Despite the random \( G_n \) accessing process, Figure 6b shows that \( I_D \) amplitudes are accumulated in five distinct distributions that correspond to the rows of the \( 5 \times 5 \) \( G_n \) grid. The response of accumulated \( I_D \) amplitudes of each row (X-direction or column #, for \( Y = 1–5 \) or row #) is shown in Figure 6c. The iontronic multiplexer is able to discriminate random inputs across the X-direction, with \( I_D \) amplitudes to be accumulated in district ranges that increase across X-direction. For distant rows (row #5), however, the ability for X-axis discrimination is suppressed, due to the fact that the channel appears as a point device. The theoretical distribution of \( I_D \), as predicted by the modular model, is also indicated in Figure 6b,c (x symbols) highlighting the effectiveness of the circuit model to simulate and predict the experimental behavior of the iontronic multiplexer.

### 3. Discussion

In this work, we demonstrated a bio-inspired iontronic multiplexer based on spatiotemporal dynamics of multiterminal OECTs. Inspired by biological systems, an electrolyte is the communication medium of the iontronic multiplexer. Electrolyte gating of the channel with a grid of gates allows for the formation of “virtual” wires in the electrolyte continuum that exhibit an Ohmic-like behavior. The configuration of the proposed iontronic multiplexer defines an all-to-one connectivity between the inputs and output, with inputs that are channelized though the spatiotemporal device dynamics and are readily differentiated at the output. The system exhibits a well-defined spatiotemporal response and predictable behavior as confirmed by the physical-based iontronic circuit model. Although the proposed model has been validated for a specific set of materials and electrolytes, it also has great predictive value and can be used universally in a greater family of materials consisting of mixed-conductors and electrolytes (e.g., organic or inorganic materials and solid or liquid electrolytes). Owing to the spatiotemporal response of the output current, the intrinsic properties of the system, namely the distance-dependent \( G_n \)-channel coupling, open possibilities for blind and local multiplexing of ionic–electronic signals. The term “blind” means that the output inherently depends on the spatial arrangement of the source of the electrical perturbation, without a-priori knowledge of the address of a gate, \( G_n \). Multiplexing is also local, meaning that it is readily incorporated in the spatiotemporal properties, and it does not result as a complex access circuitry, as for example in current passive or active matrices.

The blind and local iontronic multiplexing can be used for applications when simplicity and compactness is preferred against circuit complexity, as it decreases significantly the complexity in access and reading processes of spatially distributed signal sources (i.e., [bio]sensors or biological entities). Therefore, our approach finds relevant application in various emerging application fields including bioelectronics, wearables, e-skins, and neuromorphic sensors, where minimization of physical wiring and peripheral circuitry is essential for compact communication between the sensing and processing modalities. In addition, the proposed iontronic system can be extended for multiplexing of temporally sparse or even asynchronous, impulse-like input signals of fixed amplitude, such as in the case of spiking neural networks and time/frequency-domain coding of information. The reverse process of spatial reconstruction requires the detection of output current levels, which are directly related to the position of a signal source.

We also note that complementary information can be retrieved by gating multiple channels in perpendicular orientations with a grid of gate electrodes, thus enhancing fidelity and error-tolerance in input signal detection. Any limitations of the output current detection for long gate–channel distances can be mitigated by modifying the gate capacitance. Soft, solid-state
electrolytes for gating OECTs, including polyelectrolytes and ionic gels are readily available,
[60–62] enabling potentially conformal multiplexing media. Electrolytes also exhibit volumetric conductivity, therefore even allow for 3D multiplexing when used as shared medium. Finally, the electrochemical nature of the shared medium of communication offers degrees of freedom that are inaccessible when multiplexing with conductive/metal tracks. As an example, the concept can be further extended beyond ionic to biochemical multiplexing, as the system’s various terminals can be selectively functionalized aiming for different sensing modalities in a host environment.

4. Experimental Section

Device Fabrication: The devices were fabricated using standard microfabrication techniques. Source, drain, and gate electrodes were patterned on 26 mm × 76 mm microscope glass slides which were assiduously cleaned first in a soap (Micro-90) and then in a 1:1 (v/v) solvent mixture (acetone/isopropanol) sonication bath. The gold (Au) electrodes were patterned via photolithography and the use of photoresist (S1813). Two layers of Parylene C were subsequently used to electrically insulate the device. Between these layers an anti-adhesive layer (Micro-90 soap solution, 1% v/v in deionized water) was deposited in order to facilitate a later peel-off step. Contrariwise, an adhesion promoter (silane A-174 [gamma-methacryloxypropyltrimethoxysilane]), was employed between the substrate and the first Parylene C layer in order for the adhesion to be enhanced. A different photoresist (AZ 9260) was afterward spun on top of the two Parylene C layers and a second photolithography/development step defined window openings on the photoresist. Reactive ion etching with O2/CF4 plasma was used to remove Parylene C under those window openings and to create both the transistor channel and their corresponding gates. PEDOT:PSS was afterward spun creating a thin film of a conducting polymer that connected source and drain and coated the gate’s active area, resulting in a PEDOT:PSS film of thickness ≈500 nm. The subsequent peel-off step defined both the transistor’s channel and the gate electrodes. The external metal lines were protected under the insulating Parylene C layer. Finally, the devices were hard baked for an hour at 140 °C and placed in deionized water over night for the removal of the excess of

Figure 6. Local multiplexing of ionic–electronic signals. a) The gates, Gn (n = 1–25), of the iontronic multiplexer are randomly addressed by applying square voltage pulses (input signals) at each Gn sequentially (pulse amplitude: 500 mV, pulse width: 100 ms, in total ≈1000 access events), and the amplitude of the output current, I0, is defined by measuring the drain current (VDS = -100 mV). I0 during the random Gn accessing is represented across X- and Y-direction. b) Distribution of I0 (amplitude of I0 and the corresponding Gaussian fit) by scanning events across the rows of the Gn grid (Y-direction, row #). c) Distribution of I0 (amplitude of I0 and the corresponding Gaussian fit) of each row (X-direction, column # for each row #). Position on the 5 × 5 Gn gate grid and I0 are color-coded accordingly. The theoretical distribution of I0 as predicted by the modular model is also indicated (x symbol). The system is able to discriminate random input events in the 2D space. NaCl electrolyte at concentration c = 100 mm is used.
any low molecular weight molecules. Following the fabrication protocol described above, both large (W × L = 500 µm × 15 000 µm) and small (Width × Length, W × L = 20 µm × 20 µm) multiterminal devices were fabricated. The gate grid was the same in both cases, with dimensions for each gate electrode (2000 µm × 2000 µm) and center-to-center spacing, 8000 µm. Each gate row was at a distance from the channel that equals to 1750 + 7000·Y (µm), with Y = 2–5, the number of gate row (detailed schematic of the dimensions is depicted in Figure S1, Supporting Information). PEDOT:PSS formulation: 38 mL of PEDOT:PSS aqueous dispersion (Clevios PH-1000), 2 mL of ethylene glycol (conductivity enhancement), 50 µL of 4-dodecylbenzenesulfonic acid (film formation), and 0.4 mL of 3-methacryloxypropyl-trimethoxysilane (surface adhesion promoter and polymer cross-linking agent).

Device Characterization: All measurements of the multiterminal OECT were performed with an aqueous NaCl electrolyte, at various concentrations (c = 5–500 mM). Electrochemical IS measurements were realized with an impedance analyzer (PalmSens4). Typically, IS spectra were acquired in a frequency range = 50 mHz–100 kHz and with an amplitude of the AC signal = 25 mV. During the experiments, polydimethylsiloxane wells were used to confine the electrolyte. Source and drain electrodes were shorted and the PEDOT:PSS film was operated as the working electrode in a three electrode configuration set-up, where S and D electrodes were on the same potential and the channel was effectively operated as an electrode. A thin platinum (Pt) sheet was the reference electrode. Each of the 25 gold gate electrodes, covered with PEDOT:PSS, was the counter electrode. In the multiterminal OECT configuration, each device with 25 gates resulted in 25 impedance spectra in total. Low frequency I–V characteristics of the multiterminal OECTs, were performed with a semiconductor analyzer (Keithley 4200A-SCS) and source measure units. High-frequency pulsed measurements were performed with a custom-built setup (refer to Supporting Information): a custom-made, electronic board with 25 pulsing units was used for applying pulses at the gate electrodes (inputs: I), while resulting output current (output: O) was measured, a semiconductor analyzer (Keithley 4200A-SCS), at the drain electrode (D) with the source electrode (S) being grounded. The electronic board was synchronized with the semiconductor analyzer with time triggering. Blind multiplexing measurements were performed with this configuration and a software to control the electronic board that was randomly and sequentially assigning gate addresses and applying random input pulses at each gate electrode. As a result of this random gate addressing, the output current was measured with the semiconductor analyzer (Keithley 4200A-SCS).

Analysis: Data plotting, as well as the corresponding analysis (peak analysis of the output current, equivalent circuit modeling of the IS spectra) was performed in OriginPro 2016. Peak analysis was performed with a threshold of 2–5% of the maximum for output current peak detection, after subtracting the background current (2–5% threshold was found to be adequate for reliable detection of output peaks from background noise). Circuit modeling of the IS spectra was performed with a series RC circuit. Additional equivalent circuit modeling was performed with the impedance analyzer software (PSTrace). Modeling of the I–V and pulsed I–t characteristics of the multiterminal OECT was performed with MATLAB. Circuit simulations were performed by including the custom models in Cadence Virtuoso software.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest
The authors declare no conflict of interest.

Data Availability Statement
Data available on request by the authors.

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