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Hemmatian, Zahra Tunuguntla, Ramya H Noy, Aleksandr <u>et al.</u>

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Electronic control of H⁺ current in a bioprotonic device with carbon nanotube porins

Zahra Hemmatian¹, Ramya H. Tunuguntla², Aleksandr Noy^{2,3}, Marco Rolandi¹*

1 Department of Electrical and Computer Engineering, University of California Santa Cruz, Santa Cruz, CA, United States of America, 2 Physics and Life Sciences Directorate, Lawrence Livermore National Laboratory, Livermore, CA, United States of America, 3 School of Natural Sciences, University of California Merced, Merced, CA, United States of America

* mrolandi@ucsc.edu

Abstract

Hybrid biotic abiotic devices can be used to interface electronics with biological systems for novel therapies or to increase device functionality beyond silicon. Many strategies exist to merge the electronic and biological worlds, one dominated by electrons and holes as charge carriers, the other by ions. In the biological world, lipid bilayers and ion channels are essential to compartmentalize the cell machinery and regulate ionic fluxes across the cell membrane. Here, we demonstrate a bioelectronic device in which a lipid bilayer supported on H⁺- conducting Pd/PdH_x contacts contains carbon nanotubes porin (CNTP) channels. This bioelectronic device uses CNTPs to control of H⁺ flow across the lipid bilayer with a voltage applied to the Pd/PdH_x contacts. Potential applications of these devices include local pH sensing and control.

Introduction

Bioelectronic devices that interface with biological systems have many potential applications including new therapies and computational systems with functionalities beyond silicon [1, 2]. Examples include electroceuticals [3], wearables [4], electronic plants [5], and edible electronics [6]. In biological systems, membrane proteins and ion channels contribute to most of the communication between cells and their environments. Ion channels either passively allow or actively control the flow of ions, typically Na⁺, K⁺, Cl⁻, and Ca²⁺, and small molecules across the cell membrane [7]. Although protons are not directly involved in neuronal action potential generation and propagation, proton (H⁺) currents and concentration, [H⁺], gradients play essential physiological roles in a number of other processes [8]. The most striking example is oxidative phosphorylation in mitochondria in which proton gradients serve as a means to translate the energy from oxidation of glucose during the Kreb's cycle into ATP, the biological energy currency [9, 10]. Other examples include the light-activated H⁺ pumping by archaeal bacteriorhodopsins [11], the activation of bioluminescence from H⁺ in dinoflagellates [12], the bacterial flagellar motor activation [13], and the activity of the antibiotic Gramicidin [14].

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Recently, carbon nanotube porins (CNTPs) have emerged as artificial channels that can conduct protons across lipid bilayers [15]. In particular, narrow sub-1-nm diameter CNTPs that force water into a single-file wire conformation show very high H⁺ conductivity exceeding that of Gramicidin channel and even exceeding the intrinsic conductance of Nafion [16].

Man-made electronic platforms, which use electronic currents to carry charge have an intrinsic difficulty connecting to the biological systems that rely mostly on ionic currents [17]. To this end, many efforts in bioelectronics focus on strategies to interface ionic and electronic signaling. For example, researchers showed carbon nanotube, silcon nanowire, and organic field effect transistor devices that integrated gramicidin and rhodospins as gating elements [18–20]. Organic bioelectronics[21–22] with mixed ionic [23–25] and electronic conductivity enables devices that can both record and stimulate physiological function, and can be assembled into logic circuits [26]. We have recently demonstrated bioprotonic devices that control the flow of H⁺ in field effect transistors (H⁺-FETs) [27–30] and memories [31], and integrated these devices with enzymes to create logic gates [28]. H⁺ conducting transistors with squid reflectin proteins have also been described [32–34]. All of these H⁺ conducting devices incorporated Pd/PdH_x contacts, which translated an H⁺ current into an electrical response [30,35].

We have previously demonstrated the integration of gramicidin, alamethicin, and deltarhodpsin with Pd/PdH_x contacts and created devices that control H⁺ currents and modulate pH gradients across phospholipid membranes [36, 37]. Here, we expand these types of devices to a fully synthetic platform by using CNTPs as H⁺ channels mimics (Fig.1). These devices comprise a supported lipid bilayer (SLB) that mimics the function of a cell membrane at the Pd/solution interface and acts as a self-sealing support for the insertion of the CNTP channels.

Materials and methods

Devices

We fabricated bioprotonic devices as previously described [36,37]. In brief, we defined a Pd contact area ranging from $10 \ \mu\text{m}^2$ ($2 \ \mu\text{m} \times 5 \ \mu\text{m}$) to $200 \ \mu\text{m}^2$ ($2 \ \mu\text{m} \times 100 \ \mu\text{m}$) by evaporating 100nm Pd with a 5nm Cr as an adhesive layer onto a Si wafer with a 100 nm SiO₂ insulating layer. A photolithographically patterned SU-8 photoresist defined a microfluidic channel over the device. A polydimethylsiloxane (PDMS) well was placed on top of the microfluidic channel to confine the solution and to provide enough room to insert the Ag/ AgCl electrode that acts as a counter and a reference electrodes. The channel was filled with a buffered solution of different strength and composition as described in the Results section.

Liposome and CNTPs Liposome

Liposomes were prepared using a 1,2-Dioleoyl-sn-glycero-3-phosphocholine (DOPC, Sigma Aldrich Lipids). We followed a hydration- dehydration protocol to form liposomes. The lipids were dissolved in chloroform into glass vials and became dehydrated by evaporation of the solvent under a stream of argon gas. To dry the lipids further we stored them in a vacuum desic-cator chamber overnight. Liposomes were prepared by hydrating the lipids with a buffer solution containing 10 mM HEPES, 150 mM NaCl, 30 mM KCl pH = 7.0 to the dried lipid film to obtain a final lipid concentration of 2 mg ml⁻¹. This solution was hydrated at room temperature and bath-sonicated for 30 min. Liposomes were then extruded 20 times through a 200 nm (LUVs) pore-sized polycarbonate membrane using a mini-extruder (Avanti Polar Lipids). We used Dynamic Light Scattering (DLS) to characterize the size of liposomes.

To incorporate 1.5 or 0.8 nm diameter CNTPs into the liposomes, we first dissolved appropriate amount of DOPC–CNTP complex into 2 ml solution. We then kept the mixture in a vacuum desiccator overnight to remove the solvent. The dried DOPC–CNTP complex was



Fig 1. A bioprotonic device with integrated carbon nanotube porins (CNTPs) supports proton current across the SLB through the CNTPs when a negative voltage (-*V*) is applied on the Pd contact. When H⁺ reach the surface of the Pd contact, they are reduced to H by an incoming electron and diffuse into the Pd to form palladium hydride (PdH_x) . The current density at the contact ($-i_{H+}$), measures the rate of H⁺ flux along the CNTPs.

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hydrated with 10 mM HEPES, 150 mM NaCl and 30 mM KCl at pH 7.0 and bath-sonicated until completely solubilized. We prepared a dried lipid film in a separate glass vial, hydrated it using the solubilized DOPC-CNT complex, and bath-sonicated the solution to obtain a final lipid concentration of 2 mg ml⁻¹. The mixture was extruded through a 200 nm polycarbonate filter using a mini-extruder (Avanti Polar Lipids). The vesicle size was characterized using a DLS instrument (Malvern).

Supported Lipid Bilayers (SLB)

We deposited DOPC or DOPC-CNTP liposomes solution onto the Pd contact and formed an SLB over it using vesicle fusion. This SLB mimics a cell membrane, electrically insulates the Pd contact ($\rho \sim 3 \text{ G} \Omega \text{ cm}^{-1}$) and divides the solution into two volumes. We refer to the larger volume containing the Ag/AgCl electrode as the bulk solution (B). We refer to the small volume between the SLB and the Pd contact as the isolation layer (IL).

Atomic Force Microscopy (AFM)

We used an MFP-3D OriginTM (Oxford Instruments- Asylum Research) operating in air at room temperature, in conductive atomic force microscope (c-AFM) mode, which measures the current through the tip as a function of applied voltage, V. To preserve the integrity of the SLB coating we maintained high relative humidity of 75%. At this high relative humidity and without drying the sample, we postulate that a thin layer of water is still present on the SLP and keeps it stable. For sample preparation, excess buffer solution was gently removed from the surface of the Pd contact. The sample then was placed into a conductive sample holder to complete the electric circuit. The tip position was controlled using a custom-made LabView program that converts graphic information to voltage commands, which are sent to the AFM scanner. Current measurements were performed using a dual-phase amplifier with the amplitude of -250 mV and frequency of 0.99 Hz.

Results and discussion

In our bioprotonic devices, the Pd contact acts as a working electrode and the Ag/AgCl acts as a counter and reference electrode. For a bare Pd contact, when we apply a DC voltage difference between the Pd contact and the Ag/AgCl electrodes (V), we measure the resulting proton current (i_{H+}). For -V applied to the Pd contact, H⁺ flow from the solution onto the Pd surface, where they are reduced to H by an e⁻. These H⁺ leave behind a OH⁻ ions that make the pH of the solution increase for large enough currents as we have previously demonstrated. H then absorb into the Pd to form PdHx. Conversely, for +V, the H in PdH oxidizes into H⁺ at the Pd/solution interface and are released into solution.

After we formed a supported lipid bilayer (SLB) membrane with integrated CNTPs on the Pd contacts, applied voltage to the contact and measured i_{H+} . As control, with the Pd contact protected by the SLB in the absence of CNTPs, applied voltage of V = -250 mV resulted in a negligible $i_{H+} = -0.06 \pm 0.01$ nA (Fig 2A and 2D). This amount of i_{H+} indicates that the SLB creates an effective barrier that minimizes transport of H⁺ to the Pd contact surface. To confirm this result, we set V = 20 mV after applying V = -250 mV for 10 minutes. If any H⁺ crossed the SLB with V = -250 mV, they will reduce onto the surface and diffuse into the Pd to form PdH_x [28]. This PdH_x has a higher protochemical potential (μ_{H+}) than the pH = 7.0 solution at V = 20 mV. As a result of this higher μ_{H+} for the PdH_x contact, H would oxidize at the



Fig 2. (a) Pd contact with SLB. The SLB blocks H⁺ from transferring from the solution to the Pd contact even with V = -250 mV (vs. Ag/AgCl). (b) Pd contact with SLB incorporating 0.8 nm diameter CNTPs is semipermeable to H⁺, with CNTPs facilitating the rapid flow of H⁺ to the Pd/solution interface. (c) Upon addition of Ca⁺² to the bulk solution, H⁺ current through CNTPs becomes partially blocked. (d) i_{H+} vs. time plots recorded at V = -250 mV and V = 20 mV. Blue trace: SLB, red trace: SLB with CNTPs, gray trace SLB with CNTPs in presence of Ca⁺² ions in the bulk solution. (The data are collected from 3 different devices with different dimensions: Pd / SLB: 3 different devices of 2×50 µm, Pd/SLB+CNTPs: 3 different devices of 2×50 µm. The error bars are the root mean square of the displacement of the data from the average value).

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Fig 3. (a) Pd contact with SLB incorporating 0.8 nm diameter CNTPs with K-HEPES buffer at pH = 6.0, is semipermeable to H^+ , with CNTPs facilitating the rapid flow of H^+ to the Pd/solution interface. (b) Pd contact with SLB incorporating Narrow CNTPs with K-HEPES buffer pH = 7.0, is still semipermeable to H^+ but facilitating lower flow of H^+ to the Pd/solution interface. (c) i_{H+} versus time plot for V = -250 mV and V = 50 mV. Gray trace SLB, red trace SLB+ CNTps (K-HEPES, pH = 6.0), black trace SLB+ CNTPs (K-HEPES, pH = 7.0). The i_{H+} for measurements K-HEPES pH = 6.0 is higher than K-HEPES pH = 7. We can hypothesize that at pH = 6.0 we have a driving force due to the lower pH across the membrane in addition to the applied voltage that expedite the flow of H^+ while at pH = 7.0 we have only the applied voltage as a driving force to transport the H^+ across. We did not observe any significant different between the i_{H+} at pH = 8.0 as compare to pH = 7.0 which might be due the buffer capacity of HEPES at different pH condition (Fig A in <u>S1 File</u>). (The data are collected from 3 different devices with different dimensions: SLB- K-HEPES pH = 7.0 : 3 different devices of $2 \times 50 \mu m$. The error bars are the root mean square of the displacement of the data from the average value).

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 PdH_x contact solution interface and a proton current would flow from the PdH_x contact into the solution at the IL. This current would be measured as $i_{H+}[28]$ with a positive value. The blue trace in Fig 2D shows that this is clearly not the case.

In contrast, when we inserted CNTPs in the SLB, i_{H+} was much larger with V = -250 mV and i_{H+} continually increasing with $i_{H+} = -1.12 \pm 0.05 \text{ nA}$ at t = 10 min. (Fig 2B and 2D). This large i_{H+} confirms our conjecture that the CNTPs inserted in the SLB support the H⁺ current. To verify this conjecture, we added 1 mM Ca⁺² ions, which were previously demonstrated to block H⁺ from entering CNTPs [16] and carrying current across the SLB (Fig 2C) [19]. As expected, i_{H+} values recorded at V = -250 mV is i $_{H+} = -0.47 \pm 0.03 \text{ nA}$ (Fig 2D, grey trace).

To confirm that H⁺ are indeed the carriers for the observed for i_{H+} , we performed the experiments with different buffers that maintain pH = 7.0 or pH = 6.0 (Fig 3A, Fig 3B). Using the same V sequence that we used in Fig 2, we consistently measured i_{H+} to be higher at pH = 6.0 (i_{H+} = -2.08 ± 0.02 nA nA) (Fig 3C, red trace) than at pH = 7.0 (i_{H+} = -1.58 ± 0.04 nA nA) (Fig 3C, black trace). This is expected because lower pH value corresponds to higher H⁺ concentration for H⁺ current to flow. To exclude the possibility of the current being caused by the ionic flux through the CNTPs, we also performed experiments with buffer with and without K⁺ ions. At the same pH, we observed very little difference between i_{H+} whether K⁺ ions were present, i_{H+} = 1.10 ± 0.05 nA or not i_{H+} = 1.16 ± 0.04 nA (Fig A in S1 File)., indicating





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that K^+ ions do not significantly contribute to i_{H^+} . Note that if K^+ ion current through CNTPs is, then even larger ions would provide an even smaller contribution to the device current.

To confirm the presence of CNTPs in the lipid bilayer, we performed conductive AFM imaging to generate a current map (Fig 4A). In this map, green spots correspond to areas where the current measured by the conductive AFM tip is larger than the background, shown in purple. When the AFM tip is placed on top of a CNT indicated by a hot spot shown in green, the measured current is as high as i = 1.78 nA \pm 0.09 nA with V = 250 mV applied between the tip and the Pd contact (Fig 4B, red trace (A)). When the tip is placed on the top of the lipid bilayer in a region associated with purple on the map, minimal current of i = 5.86 pA \pm 0.98 pA is measured (Fig 4B, black trace (B)) confirming that the supported lipid bilayer is a good insulator for the Pd contact. The current measured in Fig 4B is likely electronic current due to the direct contact of the CNTPs with the tip, which by pressing on the CNTP also creates a contact with the CNTP and the Pd/ PdH_x contact. There is likely a component of H⁺ current but we estimate that this component is negligible. From these images, we estimate 1,000 CNTPs per contact, thus the H⁺ current per CNTP as measured with the bioprotonic device is ~ 1 pA per CNTPs, which is much smaller than the one recorded in Fig 4B. Nonetheless, the proton current measured with the CNTPs integrated in the SLB is comparable with what have measured for gramicidin [36].

Conclusions

In conclusion, we have demonstrated that CNTPs can act as gramicidin mimics when integrated in a bioprotonic device formed by Pd contacts with a single lipid bilayer membrane. Using buffers with different H⁺ concentration and K⁺ concentration we have confirmed that CNTPs in our devices predominantly conduct a current of H⁺. Given the ability of CNTPs to penetrate cells, and their high intrinsic proton conductance CNTPs may be used to connect bioprotonic devices directly with cells to modulate intracellular pH or to create localized pH gradients in solution.

Supporting information

S1 File. Electronic control of H+ current in a bioprotonic device with carbon nanotubes. Fig A. (a) Pd contact with SLB incorporating Narrow CNT (0.8nm) with HEPES buffer at pH = 7.0, is semipermeable to H+, with CNT facilitating the rapid flow of H+ to the Pd/solution interface. (b) Pd contact with SLB incorporating Narrow CNT with K-HEPES buffer pH = 7.0, is still semipermeable to H+ and facilitating flow of H+ to the Pd/solution interface. (c) iH+ versus time plot for V = -250 mV and V = 50 mV. Gray trace SLB, red trace SLB + CNT (K-HEPES), black trace SLB+ CNT (HEPES). The change in iH+ for measurements with potassium (K-HEPES) and without potassium (HEPES) is negligible. (The data are collected from 3 different devices with different dimensions: SLB- K- KEPES pH = 7.0: 3 different devices of 2×50 µm, SLB/ CNTPs- K- KEPES pH = 6.0: 3 different devices of 2×50 µm, SLB/ CNTPs- K- KEPES pH = 7.0: 3 different devices of 2×50 µm. The error bars are the root mean square of the displacement of the data from the average value). (DOCX)

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Author Contributions

Conceptualization: Zahra Hemmatian, Ramya H. Tunuguntla, Aleksandr Noy, Marco Rolandi.

Funding acquisition: Marco Rolandi.

Investigation: Zahra Hemmatian, Ramya H. Tunuguntla.

Methodology: Zahra Hemmatian, Ramya H. Tunuguntla, Marco Rolandi.

Project administration: Marco Rolandi.

Supervision: Aleksandr Noy, Marco Rolandi.

Writing - original draft: Zahra Hemmatian, Marco Rolandi.

Writing - review & editing: Zahra Hemmatian, Aleksandr Noy, Marco Rolandi.

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