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RuO\(_2\) Supercapacitor Enables Flexible, Safe, and Efficient Optoelectronic Neural Interface

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Optoelectronic biointerfaces offer a wireless and nongenetic neurostimulation pathway with high spatiotemporal resolution. Fabrication of low-cost and flexible optoelectronic biointerfaces that have high photogenerated charge injection densities and clinically usable cell stimulation mechanism is critical for rendering this technology useful for ubiquitous biomedical applications. Here, supercapacitor technology is combined with flexible organic optoelectronics by integrating RuO\(_2\) into a donor–acceptor photovoltaic device architecture that facilitates efficient and safe photostimulation of neurons. Remarkably, high interfacial capacitance of RuO\(_2\) resulting from reversible redox reactions leads to more than an order-of-magnitude increase in the safe stimulation mechanism of capacitive charge transfer. The RuO\(_2\)-enhanced photoelectrical response activates voltage-gated sodium channels of hippocampal neurons and elicits repetitive, low-light intensity, and high-success photoelectrical response activates voltage-gated sodium channels of hippocampal neurons and elicits repetitive, low-light intensity, and high-success rate firing of action potentials. Double-layer capacitance together with RuO\(_2\)-induced reversible faradaic reactions provide a safe stimulation pathway, which is verified via intracellular oxidative stress measurements. All-solution-processed RuO\(_2\)-based biointerfaces are flexible, biocompatible, and robust under harsh aging conditions, showing great promise for building safe and highly light-sensitive next-generation neural interfaces.

1. Introduction

Bioelectronics uses electrical stimulation of neurons to treat a wide variety of diseases such as hearing loss, Parkinson’s disease, epileptic seizures, and depression without drug injection into the body.[1] In bioelectronic devices, electrical signals, which are converted to ionic currents at the abiotic–biotic interfaces, are generally carried to the targeted tissues via wires. Instead, light offers a wireless stimulation trigger[2] enabling minimally invasive implantation of the optoelectronic devices and operation at high spatiotemporal resolution.[3] Silicon-based optoelectronic devices are successfully used for photostimulation of retina,[4] but the low absorption coefficient of silicon due to its indirect bandgap necessitates thicker devices, which adds to the overall rigidity, volume, and weight of the devices.[5] Alternatively, organic optoelectronics shows high promise for next-generation neural interfaces due to the high absorbance and unique solution processability of organic semiconductors, which together enable low-cost and configurable device fabrication, lower thickness, and flexibility.[6]

For organic optoelectronic neural interfaces, stimulation mechanism and efficiency of conversion of light to ionic currents are important to ensure safe operation without damaging tissues and to enhance the dynamic range of the devices, respectively. Since irreversible faradaic reactions can change the pH and deteriorate homeostasis of the cellular environment, operation using double-layer capacitance or reversible faradaic reactions is critical for safe stimulation.[7] In fact, supercapacitors, which show high promise for energy storage applications, are based on fast and reversible reduction–oxidation (redox) reactions at the electrode–electrolyte interface, which yield notably large interfacial capacitance values.[8] This phenomenon can be fundamentally used for photostimulation of neurons by organic neural interfaces.[9] Among pseudocapacitive materials (e.g., PEDOT:PSS[10]), RuO\(_2\) is one of the most studied supercapacitor materials as it has the highest specific capacitance (=1000 F g\(^{-1}\)).[11] Although different ruthenium complexes (e.g., Ru\(^{16+}\)/Ru\(^{14+}\)) have been previously used as anticaner agents,[12] supercapacitor RuO\(_2\) has attracted little attention for biomedicine[13] and bioelectronics,[14] and has not been explored for organic optoelectronic neural interfaces yet.
Here, we demonstrate that RuO₂ supercapacitor significantly boosts photogenerated charge injection densities over 20-fold in biological fluid of artificial cerebrospinal fluid (aCSF) in comparison with the control devices without RuO₂. It enables safe, repetitive, and low-light intensity photostimulation of hippocampal neurons below 1 mW mm⁻². More interestingly, we developed an all-solution-processed fabrication (on a flexible substrate) of organic optoelectronic neural interfaces by sensitive incorporation of RuO₂ via electrochemical deposition that allowed high-level control and optimization of charge injection density. The results point out that RuO₂ is a powerful material option for flexible optoelectronic neural interfaces and its electrochemical integration enables biocompatible, robust, and highly light-sensitive devices for future retinal implants.

2. Results

The solution-processed fabrication of the optoelectronic neural interface started with the sequential spin-coating of ZnO nanoparticles (NPs) and P3HT:PCBM bulk heterojunction (BHJ) thin films on fluorine-doped tin oxide (FTO)/indium tin oxide (ITO) substrates (Figure 1a). During these steps, a part of the substrate was covered with a mask to prevent the deposition of ZnO NPs and P3HT:PCBM BHJ to the area where RuO₂ return electrode will be formed. Then, after removal of the mask, RuO₂ layer was coated on the back electrode (FTO/ITO) via cyclic voltammetric deposition.\[15\]

We investigated the structural properties of RuO₂ films having different number of electrodeposition cycles (5, 15, 25, 40, 60). The atomic force microscopy (AFM; Figure S1, Supporting Information) revealed surface morphology of RuO₂ coatings, which have root-mean-square (RMS) surface roughness of 15.8, 15.6, 18.4, 22.9, and 23.1 nm for 5-, 15-, 25-, 40-, and 60-cycle RuO₂ coatings, respectively. Considering the maximum diffusion length of protons in RuO₂ (≈20 nm), those roughness values are at desirable levels for facilitating ion transportation during fast redox reactions.\[16\] X-ray photoelectron spectroscopy (XPS) analysis of Ru 3d\[3/2\] spectrum (Figure S2, Supporting Information) indicates a high amount of Ru(IV) species due to the peak at 281.4 eV corresponding to RuO₂.xH₂O,\[17\] which are possibly due to their higher film thickness and narrower pore sizes (Table S1, Supporting Information). In the low-frequency region, imaginary impedance was gradually decreasing as the frequency region with a resistance–constant phase element (R–Q) (Figure S4, Supporting Information). In the high-frequency region, 5-, 15-, and 25-cycle RuO₂ coatings exhibited similar resistances, while 40- and 60-cycle RuO₂ coatings display increased resistances possibly due to their higher film thickness and narrower pore sizes (Table S1, Supporting Information). In the low-frequency region, imaginary impedance was gradually decreasing as the cycle number increased (Table S1, Supporting Information), indicating higher capacitance values for thicker films as we observed in CV measurements. Even though the capacitance value from the CV followed the trend of the number of deposition cycles, the capacitance measured using EIS showed 25 cycles as an outlier, which is probably because of the increased resistance and thus lower porosity.

High CSC and capacitance of RuO₂ indicate its potential for achieving improved charge injection levels for the optoelectronic neural interface. To experimentally test that, we studied the effect of RuO₂ on the photosresponse of a control device. In our device architecture, ZnO/P3HT:PCBM photovoltaic heterostructure serves as the photocurrent-generating electrode. P3HT:PCBM BHJ absorbs the impinging light and afterward electron–hole pairs are dissociated at the donor–acceptor interface. Free electrons are transferred toward ITO through ZnO NPs. ZnO layer serves as the hole blocker due to its valence band energy level and electron transporter owing to its high electron mobility.\[20\] The device design consists of ZnO/P3HT:PCBM layer over 1 cm² area of the ITO-coated substrate and electrodeposited RuO₂ film over the remaining 0.5 cm² area of the same substrate (Figure 2a). The control devices are bare ITO as the return electrode without RuO₂ coating.

We analyzed the photoelectrical performance of the control and RuO₂ biointerfaces by comparing their charge-injection performances in a wireless, free-standing measurement setup (Figure 2b). Devices were placed in aCSF and injected charge was measured in voltage-clamp mode with a patch pipette that is positioned close to the biointerface surface (<5 μm). This measurement configuration demonstrates the ability of biointerfaces to perturb the local electric equilibrium at the device–electrolyte interface, revealing the potential of the biointerface to stimulate...
neurons. The photoresponse of the control devices was characterized by an initial capacitive peak triggered by light onset and a second peak with a smaller magnitude taking place with the end of illumination (Figure 2c). The capacitive onset peak decays rapidly to its ground level, meaning no significant capacitive current is flowing under steady illumination condition because of the double-layer capacitance with minimal Faradaic reactions under steady photovoltage. This considerably limits the amount of injected charge required for effective stimulation of neurons.

After RuO₂ deposition, the reversible surface redox processes induced by the coating lead to a remarkable amount
of additional charge injection into the electrolyte due to the enhancement of the return electrode capacitance (Figure 2c).[22] The steady photovoltage during illumination evokes reversible Faradaic reactions on the RuO₂ surface, predominantly redox reactions of Ru⁴⁺/Ru³⁺ couple,[14a,18] leading to formation of a redox capacitance, which is embedded in the redox impedance of the interface.
of return electrode, in addition to the double-layer capacitance (Figure S5, Supporting Information). Considering that the double-layer capacitance of typical organic electrode–electrolyte interfaces is on the order of 1 to a few tens of μF cm\(^{-2}\),\(^{6,9}\) the double-layer capacitance of RuO\(_2\)–electrolyte interface is expected to be on the similar order. However, the redox impedance, which includes the pseudocapacitance of RuO\(_2\) and is in parallel to the double-layer capacitance, significantly boosts the interfacial capacitance of the return electrode, which is counter-balanced by faradaic reactions at P3HT:PCBM–electrolyte interface (Figure S6, Supporting Information). As a result, the onset photocurrent peak increased more than two-fold (Figure S7, Supporting Information). Noticeably, the amount of the total charge injection substantially increased, which can be controlled with the deposition cycles. While the charge injection capacity of control devices was 2.5 ± 1.3 pC (mean ± SD), for 5-cycle-RuO\(_2\) biointerfaces it jumped to 29.8 ± 8.1 pC (mean ± SD), indicating more than an order of magnitude increase. For 60 cycles, the injected charge reached 57.1 ± 13.7 pC (mean ± SD), corresponding to over 20-fold enhancement compared to control devices (Figure 2d). If the counter-balancing faradaic reactions at P3HT:PCBM–electrolyte interface are suppressed, the charge injection performance of RuO\(_2\)-based biointerfaces notably decreases (Figure S6, Supporting Information), which confirms that the observed improvement in photoresponse is due to the redox impedance of RuO\(_2\). As a further control experiment, we checked whether RuO\(_2\) generates photocurrent by itself, and verified that RuO\(_2\) layer alone produces only a marginal photocurrent, which is at least two orders of magnitude lower than the photovoltaic part (Figure S8, Supporting Information). Since the best performing device is 60 cycle RuO\(_2\), the rest of the experiments were conducted with 60 cycle RuO\(_2\) biointerfaces (referred as “RuO\(_2\) BI” in the rest of the text).

Temporal characteristics of the generated photocurrent provide information about the maximum achievable stimulus frequencies. Figure 2e shows photocurrents of RuO\(_2\) BI under pulsed illumination for pulse-widths of 10 ms, 5 ms, 1 ms, 500 μs, and 200 μs. The capacitive onset peak was preserved even for 200 μs pulse, implying that high-frequency stimulus, e.g., 100 Hz, can be applied via RuO\(_2\) BI. For example, when we applied 1 ms 100 Hz pulsed excitation, the photocurrent peak and shape were well-preserved throughout the repeated photoexcitation cycles (Figure S9, Supporting Information). Lastly, we evaluated the spatial distribution of the photoresponse by measuring the photovoltage while moving the patch pipette in horizontal and vertical directions, starting from the center of the illumination spot (radius R = 160 μm; Figure 2f inset). Charge injection decays in an exponential fashion in both directions as the pipette was gradually moved away from the spot center (Figure 2f), suggesting a localized charge injection primarily governed by illumination spot area. In horizontal direction, the measured charge was still over 90% at the spot edge (1R away from the center) and it drops to 62% 2R away from the center. This represents a relatively slower decay compared to a previous study,\(^{6,9}\) most likely due to the device geometry with large return electrode.

To quantify the short-circuit photoelectrical response of RuO\(_2\) BI, we measured its photoresponse via a three-electrode potentiostat/galvanostat system (Figure 3a). Working electrode was connected to the ITO back electrode, while the counter and reference electrodes were floating in the ionic aCSF medium.\(^{14}\) The intensity-dependent photocurrent and photovoltage measurements in Figure 3b,c showed the maximum achievable current density of 6 mA cm\(^{-2}\) and 290 mV photovoltage for RuO\(_2\) BI under 445 nm 85 mW cm\(^{-2}\) LED illumination, respectively. Integrating the area under the photocurrent–time traces corresponds to the injected charges for each illumination intensity (Figure 3d) and RuO\(_2\) BI has a charge injection density of more than 10 μC cm\(^{-2}\) for intensities greater than 30 mW cm\(^{-2}\). This charge injection level is sufficiently high according to the reported threshold charge density values for stimulation of human epiretinal, cortical, auditory brainstem, and subthalamic nucleus neurons.\(^{19}\) In addition, RuO\(_2\) BI has more than an order of magnitude higher charge injection capacity compared to control devices (Figure 3d).

Previously we showed that high-frequency stimulus (e.g., 100 Hz) via short pulses (e.g., 1 ms) can be applied with RuO\(_2\) BI, Now, we demonstrate that even for such short pulse durations, the charge density of RuO\(_2\) BI is in the μC cm\(^{-2}\) range. For 1 ms pulse, injected charge density is 2.4 ± 0.6 μC cm\(^{-2}\) (mean ± SD), while it is 1.6 ± 0.7 μC cm\(^{-2}\) (mean ± SD) for 500 μs pulse (Figure 3e inset), which are still near the thresholds for stimulation of certain neural tissues (e.g., subthalamic nucleus, auditory brainstem).\(^{19}\) Moreover, for lower stimulus frequencies 20 ms pulse would result in a high charge density of 20.2 ± 2.4 μC cm\(^{-2}\) (mean ± SD; Figure 3e).

We tested the stability and biocompatibility of RuO\(_2\) BI and checked whether repeated photoexcitation of RuO\(_2\) BI causes reactive oxygen species (ROS) intake by neurons or not. Stability was studied by investigating the charge injection performance of RuO\(_2\) BI after detailed tests including low-temperature sterilization, accelerated passive and reactive aging, and repeated photoexcitation experiments. Low-temperature hydrogen peroxide (H\(_2\)O\(_2\)) plasma sterilization is an FDA approved technique for testing medical devices.\(^{25}\) After application of low-temperature H\(_2\)O\(_2\) plasma sterilization, the charge injection levels of RuO\(_2\) BI reduced to 78% ± 9% (mean ± SD) of its pre-sterilization value (Figure 4a). The sterilized RuO\(_2\) BI samples were further subjected to accelerated reactive aging test, where we kept the samples in H\(_2\)O\(_2\)-added aCSF at 87 °C oven and quantified their charge injection performance after every 48 h, which corresponds to a simulated aging period of 2 months. After an aging period of 12 months, RuO\(_2\) BI preserved 76% ± 8% of its post-sterilization and 60% ± 6% of its pre-sterilization charge injection values indicating robust operation of RuO\(_2\) BI under harsh aging conditions (Figure 4b). At this point, we wondered whether the reduction in charge injection values of RuO\(_2\) BI is due to the degradation of ZnO/P3HT:PCBM active electrode or RuO\(_2\) coating. The EIS measurements of active electrode and RuO\(_2\) coating before and after plasma sterilization and reactive aging tests showed that the impedance-frequency behavior of RuO\(_2\) is preserved after the tests, while the response of active electrode deviated from its pre-sterilization behavior (Figure S10, Supporting Information). This implies that performance drop is primarily due to the degradation of the photoactive electrode, which can be prevented via gold layer deposition on top of the electrode.\(^{23c}\) and RuO\(_2\)
remains intact under harsh aging conditions. Furthermore, RuO₂ BI exhibits excellent stability under passive accelerated aging test, showing no significant decrease in the charge injection performance after 24 months (Figure 4b). Moreover, the charge retention performance of RuO₂ BI was tested under repeated 10,000 cycle pulsed photoexcitation. After 10,000 illumination cycle, 84% ± 5% of injected charge and the capacitive current profile (Figure 4c inset) are preserved, implicating the retained functionality of the biointerfaces (Figure 4c).

To evaluate the cytotoxicity of RuO₂ BI, we conducted in vitro biocompatibility test with primary hippocampal neurons via MTT toxicity assay and immunofluorescence imaging before and after 14 days of neuron culture. MTT analysis showed that neurons on RuO₂ BI demonstrate a statistically nonsignificant difference of cell viability compared to biocompatible ITO samples after 48 h incubation period (Figure 4d). This result is supported by immunofluorescence images of ITO and RuO₂ samples. Day 0 and day 14 images of ITO and RuO₂ BI demonstrated preserved cell viability and morphology for both sample types, suggesting low cytotoxicity of RuO₂ BI for in-vitro primary hippocampal neurons (Figure 4e).

Despite the reversibility of RuO₂-induced redox reactions, we wanted to reassure that RuO₂ BI does not generate ROS in the electrolyte upon photoexcitation. To validate that, we quantified the amount of ROS intake of hippocampal neurons cultured on RuO₂ BI by measuring the fluorescence intensity of H₂-DCFDA agent after subjecting the biointerfaces to 10,000-cycle photoexcitation (Figure 4g). We used neurons on bare FTO as negative control, on bare FTO treated with 100 μM H₂O₂ as positive control, and on RuO₂ BI before photostimulation as control groups. Expectedly, neurons on positive control samples show high fluorescence intensity due to the intake of H₂O₂ in the extracellular medium. On the other hand, there is no statistical difference between the fluorescence intensities of neurons grown on bare FTO, RuO₂ BI before photostimulation, and RuO₂ BI after photostimulation (Figure 4f). Hence, repeated photoexcitation of RuO₂ BI does not cause intracellular oxidative stress for neurons, which indicates the absence of ROS formation in the aCSF. This suggests a safe photostimulation mechanism for RuO₂ BI. This is further supported by the pH measurement of extracellular medium during 40000-cycle photoexcitation of
Figure 4. Stability, biocompatibility, and intracellular oxidative stress analysis. a) Charge injection performance of RuO₂ BI before and after low temperature H₂O₂ plasma sterilization (mean ± SD for n = 3). The sum of three RuO₂ BI before the sterilization is normalized to 1. Inset shows photocurrent traces before and after the sterilization. b) Variation of RuO₂ BI charge injection levels during the passive and reactive accelerated aging tests (mean ± SD for n = 3). Reactive aging test was applied to the biointerfaces that underwent plasma sterilization. c) Photocyclic stability of RuO₂ BI showing the variation of injected charge during the application of 10 000 photoexcitation pulse (mean ± SD for n = 4). Illumination: 5 Hz stimulus frequency, 10 ms pulse-width, 2 mW mm⁻² optical power density. Inset shows photocurrent traces before and after 10 000 photocycles. d) MTT assay analysis for quantifying cell viabilities of primary hippocampal neurons cultured on RuO₂ BI (mean ± SD for n = 4). Bare ITO sample, which is known to be biocompatible, was used as control. The level of significance was calculated using an unpaired, two-tailed t-test; *p < 0.05 was evaluated as statistically significant. e) Immunofluorescence images of ITO and RuO₂ BI samples obtained on day 0 and day 14 to observe the cell viability and morphology of hippocampal neurons after 14 days of culture. f) Quantification of ROS intake, i.e., intracellular oxidative stress, of neurons cultured on RuO₂ BI after subjecting the biointerfaces to 10 000 photoexcitation cycle (mean ± SD for n = 3). Illumination: 5 Hz stimulus frequency, 10 ms pulse-width, 2 mW mm⁻² optical power density. RuO₂ before stimulation samples are control, bare FTO is negative control, bare FTO with H₂O₂ is positive control. The fluorescence intensity of negative control is normalized to 100%. g) Fluorescence microscope images of FTO, RuO₂ before stimulation, RuO₂ after stimulation and FTO with H₂O₂ samples stained with Hoechst and H2DCFDA, which were used to calculate the relative intracellular fluorescence intensities.
RuO$_2$ BI (Figure S11, Supporting Information). pH values measured after each 10 000-cycle photoexcitation do not show significant differences, implying that the chemical composition of aCSF is not altered by irreversible faradaic processes during repeated photoexcitation.

We finally explored the light-induced effect of RuO$_2$ BI on primary hippocampal neurons by measuring single-cell intracellular membrane potential and current with respect to a distant Ag/AgCl electrode. Figure 5 illustrates the measurement configuration, where RuO$_2$ BI is...
operating wirelessly (i.e., electrically floating in aCSF) to photo-stimulate the cultured neurons. First, we checked if photoexcitation of RuO$_2$ BI can open voltage-gated sodium channels and induce inward sodium current, which is essential for eliciting action potential. As an initial control experiment, we measured the current–voltage characteristics of hippocampal neurons under dark conditions by applying different membrane holding voltages via the amplifier (Figure S12, Supporting Information). Rapid negative inward currents, which originate from voltage-gated sodium channels, start to appear at holding voltage of $-50$ mV. These sodium inward currents for $-50$, $-30$, and $-10$ mV holding potentials are followed by slower outward potassium currents (Figure S12a, Supporting Information). When we add voltage-gated sodium channel blocker QX-314 chloride$^{[26]}$ into the intracellular solution, negative inward currents are no longer observed, confirming that they are due to voltage-gated sodium channels (Figure S12b, Supporting Information). Under light conditions without any external stimulus from amplifier, photoexcitation of RuO$_2$ BI with 5 ms pulses of 2.4 mW mm$^{-2}$ illumination intensity induced fast sodium inward currents followed by a slower outward potassium current, a typical current profile observed during an action potential (Figure 5b)$^{[27]}$. Lower light intensity, 0.6 mW mm$^{-2}$, did not result in the same response, which possibly led to a sub-threshold stimulation. As a control experiment, we added QX-314 chloride to the intracellular solution. As in the previous experiment, since photoexcitation of RuO$_2$ BI leads to rapid sodium inward current and slower outward potassium current for neurons under standard conditions, the inward sodium current is nearly vanished in the existence of QX-314 chloride (Figure 5c). These experiments denote that RuO$_2$ BI activates voltage-gated sodium channels of hippocampal neurons upon photoexcitation, which is expected to elicit action potentials.

To verify the firing of neurons, we conducted current-clamp recordings to examine their intracellular membrane potential with respect to a distant Ag/AgCl electrode. The recordings demonstrate that light-induced photoresponse of RuO$_2$ BI reproducibly stimulates neurons under 5 ms pulsed excitation with stimulus frequencies of 5, 10, and 20 Hz (Figure 5d). Evoked action potentials are temporally precise and some peaks are followed by afterdepolarizations, which frequently occur during firing of hippocampal neurons.$^{[28]}$ The spikes with 5 ms pulses are induced by minimum 2.4 mW mm$^{-2}$ light intensity. Figure 5e displays that longer pulses with lower illumination intensities, 10 ms at 1.25 mW mm$^{-2}$ and 20 ms at 0.65 mW mm$^{-2}$, can also elicit action potentials. As pulse-width increases, the minimum required power for photoactivation decreases. However, the minimum total energy for evoking action potentials stays nearly constant as expected for extracellular stimulation electrodes.$^{[29]}$ The resultant action potential success rates show similar fashions for 5, 10, and 20 ms pulses, showing slight decrease as the stimulus frequency is increased. The success rates stay above 80% for all pulse-widths and stimulus frequencies, except for 5 ms 20 Hz pulse, for which it is 69% ± 3 (mean ± s.d.) (Figure 5f). Higher success rates for longer pulse-widths are ascribed to continuous charge injection during light-on periods due to RuO$_2$ return electrode. On the other hand, action potential latencies are directly proportional with the pulse-widths (Figure 5g). Overall, the electrophysiology recordings signify the effective photostimulation performance of RuO$_2$ BI indicated by temporal precision, rapid reversibility, and short latencies.

3. Discussion

This study demonstrated the potential of pseudocapacitive material of RuO$_2$ to realize highly light-sensitive organic optoelectronic biointerfaces. One major obstacle for organic optoelectronic biointerfaces, especially capacitive ones, has been the difficulty of achieving high photogenerated charge densities,$^{[15]}$ and the proposed strategy in this report can be an important step toward realizing much-elevated charge injection densities. Although other alternatives such as dielectric SiO$_2$ coating and pseudocapacitive PEDOT:PSS coating were used to enhance the interfacial capacitance,$^{[10,30]}$ our findings show that RuO$_2$ distinctively outperforms those materials in terms of their contribution to charge injection densities. Moreover, the increased charge injection efficiency allows for neurostimulation with shorter pulses and latencies that can enable improved temporal control.

Advantageously, RuO$_2$ can be directly integrated into a wide variety of device architectures$^{[5,6]}$ to improve the device performance levels via the low-cost electrodeposition technique and even on non-planar substrates. Comparatively, iridium oxide in the form of SIROF was shown to perform well in a silicon-based planar photodiode architecture in terms of its contribution to charge injection performance.$^{[31]}$ However, spattering is a relatively complex and expensive method compared to solution-processing techniques. Recently it was demonstrated that sputtered RuO$_2$ exhibits similar charge storage/charge injection performances with SIROF, indicating its high potential for electrical stimulation/recording electrodes.$^{[32]}$ Here, electrodeposition of RuO$_2$ also performs at similar levels of SIROF and sputtered RuO$_2$, together with the advantage of simple device fabrication. Stability, bio-compatibility, and safe stimulation pathway through highly reversible charge transfer reactions demonstrated by intracellular oxidative stress measurements of RuO$_2$-based biointerface show promise for in vivo applications. Moreover, the effective performance of RuO$_2$ can motivate the investigation of alternative pseudocapacitive coating materials for neural interfaces. Carbon materials (e.g., carbide-derived carbon), metal oxides (e.g., manganese oxide), and conducting polymers (e.g., polypyrrole) can be effective due to their high capacitance, low toxicity, and reversible charge injection mechanisms.$^{[114]}$

In conclusion, our findings pave the way toward building safe and highly efficient optoelectronic biointerfaces by using pseudocapacitive RuO$_2$. The biointerfaces showed improved photoelectrical performance parameters compared to control samples, i.e., increased capacitive photocurrent peak and higher charge storage/injection efficiencies. Detailed electrochemical analysis of RuO$_2$ coating revealed the reversible nature of redox reactions, indicating a safe charge injection mechanism for RuO$_2$-based optoelectronic biointerfaces, which was further supported by intracellular ROS measurements. Excitation of RuO$_2$-based biointerfaces via short light pulses can activate voltage-gated sodium channels, which leads to reproducible action potential firing on primary hippocampal neurons.
Consequently, the safe and efficient stimulation mechanism obtained by simple solution-processed RuO₂ integration shows great promise for building flexible, low-cost, biocompatible, stable, and highly light-sensitive optoelectronic biointerfaces for future retinal prosthesis.

4. Experimental Section

Biointerface Fabrication: ITO/POET (Sigma Aldrich, 639303) and FTO/glass substrates (Ossila, TEC 10) were cleaned by sonication in detergent solution, deionized water, acetone, and isopropanol consecutively for 15 min each, followed by 20 min UV ozone treatment. Poly(3-hexylthiophene-2,5-diyl) (P3HT) with regioregularity of 95.7% and a molecular weight of 75467 g mol⁻¹ (Ossila) and [6,6]-PPhenyl-C61-butyric acid methyl ester (PC61BM) with a molecular weight of 1031 g mol⁻¹ were supplied by Ossila Ltd. 1,2-dichlorobenzene solutions of P3HT (20 mg mL⁻¹) and PCBM (12 mg mL⁻¹) were prepared and stirred at 70 °C overnight. P3HT and PCBM solutions were mixed (1:1 volume ratio) and stirred for further 2 h using magnetic stirrer. ZnO precursor sol–gel solution (0.45 M) was prepared by mixing 219.3 mg Zinc acetate dehydrate (Zn(CH₃CO₂)₂·2H₂O), 2 mL of 2-Methoxyethanol (C₃H₅O₂) and 73 mg of Ethanolamine (HOCH₂CH₂NH₂). Ruthenium (III) chloride hydrate (RuCl₃·H₂O) with molecular weight of 207.43 g mol⁻¹ was purchased from Sigma–Aldrich. Aqueous solution (0.01 M) of RuCl₃·H₂O was prepared for electrochemical deposition of ruthenium oxide (RuO₂).

For the fabrication of RuO₂-based biointerfaces, first, a certain area of FTO/ITO substrate, where RuO₂ will be coated, was covered with a mask. Then, ZnO precursor sol–gel solution was spin-coated on the remaining area of the substrate at 2000 rpm, followed by 15 min annealing at 290 °C for FTO and 200 min at 200 °C for ITO. On ZnO layer, P3HT:PCBM solution was spin-coated at 2000 rpm, followed by 15 min annealing at 150 °C. Then, the mask was removed. The area under the mask was immersed into 0.01 M aqueous solution of RuCl₃·H₂O and RuO₂ layer was formed via cyclic voltammetric deposition by cycling the potential between −0.2 and 1.2 V at a scan rate of 50 mV s⁻¹.

Photoresponse Analysis: Interfacial (open-circuit) photocurrents were recorded by means of a three-electrode setup consisting of Ag/AgCl as the reference electrode, platinum rod as the counter electrode, and the thin film samples as the working electrode in aCSF solution. Light pulses were applied via Thorlabs M450LPI LED with 445 nm nominal wavelength and the LED spectrum was provided in the previous study.²¹ The blue light was driven with Thorlabs DC2200 – High-Power 1-Channel LED Driver. Newport 843-R power meter was used to measure the optical power of incident light on the biointerfaces.

Chronopotentiometry and chronocamperometry measurements were performed via Autolab Potentiostat Galvanostat PGSTAT302N (Metrohm, Netherlands) using a three-electrode setup consisting of Ag/AgCl reference electrode, platinum rod counter electrode, and the thin-film biointerface samples as working electrode. The EIS was performed in frequency response analysis (FRA) potential scan mode while varying the frequency between 1 Hz and 10 kHz at 10 mV (RMS) AC voltage perturbation. The fitting of the high-frequency resistance and low-frequency capacitance was performed in NOVA software to extract the corresponding circuit parameters. A more detailed fit can be performed via a 3-RC circuit, however the extra detail would be a distraction and not add to the main argument of the paper. Therefore, only the high-frequency resistance and the low-frequency capacitance are considered.

Primary Neuron Isolation: All experimental procedures have been approved by the Institutional Animal Care and Use Committee of Koç University (Approval No: 2019.HADYEK.023) according to Directive 2010/63/EU of the European Parliament and of the Council on the Protection of Animals Used for Scientific Purposes. Hippocampal regions were extracted from decapitated E15-E17 Wistar Albino rat embryos and were placed immediately in ice-cold Hank’s Balanced Salt Solution (HBSS, Thermo Fisher Scientific, MA, USA). The hippocampi were incubated in 0.25% Trypsin-EDTA solution (Thermo Fisher Scientific, MA, USA) with 2% DNase-I supplement (NeoFroox, Eugene, Germany) for 20 min in a 37 °C incubator. Then the cells were centrifuged, and the supernatant was changed with Dulbecco’s Modified Eagle Medium/Nutrient Mixture F-12 (DMEM/F12 Thermo Fisher Scientific, MA, USA) supplemented with 10% fetal bovine serum (FBS, Heat Inactivated, GE Healthcare, IL, USA) and 1% penicillin/streptomycin (Thermo Fisher Scientific, MA, USA). DMEM/F12 was removed and Neurobasal Medium (NBM, Thermo Fisher Scientific, MA, USA) supplemented with B27, l-glutamine, β-mercaptoethanol, glutamate (Thermo Fisher Scientific, MA, USA) was added to the cell pellet. The cells were triturated and were passed through a 70 μm cell strainer. The homogenous cell solution was seeded in poly-d-lysine (PDL, Sigma–Aldrich, MO, USA) coated substrates. After 3-days incubation of cells on substrates in a 37 °C incubator with 5% carbon dioxide, the media of the cells on substrates were changed with NBM supplemented with cytosine arabinoside (Sigma–Aldrich, MO, USA) to inhibit the growth of glial cells. After 24-h incubation with cytosine arabinoside, the media were changed with NBM and the substrates with the hippocampal neurons used for the experiments.

Biocompatibility Assay: MITT viability assay was applied to investigate the cell viability of primary hippocampal neurons on the biointerfaces. The neural growth medium was prepared by using B27 supplemented Neurobasal medium. MITT cell viability assay (Abcam, ab211091) was utilized to evaluate biocompatibility of our biointerface. The devices were sterilized first by cleaning with 70% ethanol followed by air-drying. The surface was further sterilized under UV irradiation for 30 min. Substrates were placed in wells of the six-well plates. Primary hippocampal neurons were seeded (5 × 10⁵ cells per sample) on the substrates in B27-supplemented Neurobasal medium as described above and incubated in the neuron growth medium for 48 h after cytosine arabinoside supplemented neurobasal medium removal. After 48 h incubations, the media were replaced with 1 mL of MITT solution (5 mg mL⁻¹ in PBS, pH 7.4) and 4 mL of NBM mixture per well. Then, for an additional 4 h, the cells were incubated at 37 °C and 5% CO₂ atmosphere. The medium was vacuumed from each well and substrates were transferred to empty six-well plates. In each well, 1:1 mixture of DMSO and ethanol was added to dissolve the formazan crystals. The solution was transferred to a 96-well plate and the absorbance was measured at 570 nm light with Synergy H1 Microplate Reader (Bio-Tek Instruments). The relative cell viability was calculated as follows: Viability = (ODsample/ODcontrol) × 100. The optical density (OD) of the sample was obtained from the cells grown on a photoelectrode, and the OD of control was obtained from the cells grown on the ITO substrates.

Immunofluorescence Staining and Imaging: Primary hippocampal neurons (5 × 10⁵ cells per sample) were seeded as explained above on ITO control substrate and the biointerface. The samples with neurons were fixed by 4% paraformaldehyde immediately after primary hippocampal neuron isolation protocol or incubated for 14 days with regular medium changes at 37 °C in the cell culture incubator. After 14-day incubation, the primary hippocampal neurons were also fixed by 4% paraformaldehyde and washed three times with PBS-T (Phosphate Buffered Saline, 0.1% Triton X-100). Cells were blocked in PBS solution then stained.
containing 5% BSA (Bovine Serum Albumin) and 0.1% Triton X-100. Samples with primary hippocampal neurons were incubated with rabbit anti-NeuN antibody (ab177487, Abcam, Cambridge, UK) overnight, for neuron characterization, and washed three times with PBS-T. Then, samples with primary hippocampal neurons were incubated with goat anti-rabbit IgG H&L Alexa Fluor 555 (ab104139, Abcam, Cambridge, UK) to observe nuclei. Finally, immunofluorescence imaging was done using a fluorescence light microscope (Axio Observer Z1, ZEISS, Oberkochen, Germany).

Intracellular Oxidative Stress Measurement: The intracellular oxidative stress level was measured by 2’,7’-dichlorodihydrofluorescein diacetate (H$_2$DFCDA) (D399, Molecular Probes, Invitrogen). Primary hippocampal neurons (5 × 10^5 cells per sample) were seeded on FTO control substrates and the biointerfaces. The H$_2$DFCDA agent is nonfluorescent but it becomes intensive fluorescent agent after oxidation with H$_2$O$_2$. Thus, as a positive control group, neurons grown on the control substrates were treated with 100 × 10$^{-6}$ M H$_2$O$_2$ for 30 min. In the experimental group, neurons grown on the biointerfaces were exposed to 450 nm LED for 10,000 cycles with 5 Hz frequency. Cells were washed once with PBS and then incubated with 20 × 10$^{-6}$ M H$_2$DFCDA in aCSF solution for 45 min prior to imaging at 37°C, 5% CO$_2$ to allow H$_2$DFCDA to fully enter the cells. Hoechst (0.6 mg mL$^{-1}$) was used to stain the nucleus for 15 min at 37°C, 5% CO$_2$. To avoid remaining dye residues, cells were washed with PBS. To avoid photobleaching, all steps were carried out in the dark. All immunofluorescence images were taken in the same light intensity, exposure, time point, and magnification with live-cell fluorescence microscope (Axio Observer Z1, ZEISS, Oberkochen, Germany) and average fluorescence intensity was measured by ImageJ software (Image) Fiji, NIH, MD, USA). For image analysis, images were converted to 8-bit black and white image and background was subtracted. To measure number and area of cells, maximum entropy threshold was applied to the image and the particles were analyzed for integrated density. By this way, both area of neurons and intensity of fluorescence were measured in acquired images from randomly selected ten different regions of neural cell culture. Relative fluorescence intensity was calculated with negative (neurons on FTO substrate) and positive (neurons on FTO with H$_2$O$_2$ addition to induce ROS production) controls. Normal ROS production in FTO control substrates was accepted as 100% in relative fluorescence intensity comparison.

Electrophysiology Recordings: Single-cell electrophysiology experiments were performed using EPC 800 Heka Elektronik patch-clamp amplifier in whole-cell configuration. Biointerfaces were electrically floating in aCSF without any wire connection. Whole-cell transmembrane voltage and current recordings, which refer to intracellular membrane potential and current recordings with respect to a distant Ag/AgCl electrode, were taken in current-clamp and voltage-clamp modes, respectively. Downsampling was applied to the current-clamp data to obtain a feasible computational complexity for making statistical analysis of action potentials without causing loss of any meaningful data. The patch pipette resistance of 6–8 MΩ was used for the recordings. The biointerfaces were placed in aCSF and patch pipettes were filled with the intracellular medium. To block the voltage-gated sodium channels, 5 × 10$^{-3}$ M QX-314 chloride was added into the intracellular solution. Patch pipette and cells were monitored through a digital camera integrated with the Olympus T2 upright microscope.

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.

Author Contributions
O.K. and S.N. designed the experiments. O.K. fabricated and characterized the biointerfaces, performed photoresponse measurements and electrophysiology experiments. E.Y. performed primary hippocampal neuron isolation, biocompatibility assay, immunofluorescence staining, and imaging of primary hippocampal neurons. H.N.K conducted intracellular oxidative stress measurements. B.U. supervised the electrophysiology experiments and interpreted the data. A.S. supervised the cell culture, biocompatibility, immunofluorescence imaging, and intracellular oxidative stress experiments and interpreted the data. O.K. and S.N. wrote the manuscript with input from all authors. All authors contributed to the article and submitted the article to publication.

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

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