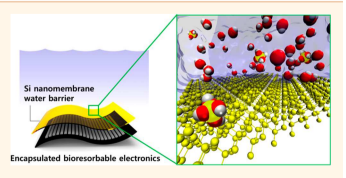


Dissolution of Monocrystalline Silicon Nanomembranes and Their Use as **Encapsulation Layers and Electrical Interfaces** in Water-Soluble Electronics

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Supporting Information

ABSTRACT: The chemistry that governs the dissolution of device-grade, monocrystalline silicon nanomembranes into benign end products by hydrolysis serves as the foundation for fully eco/biodegradable classes of high-performance electronics. This paper examines these processes in aqueous solutions with chemical compositions relevant to groundwater and biofluids. The results show that the presence of Si(OH)₄ and proteins in these solutions can slow the rates of dissolution and that ion-specific effects associated with Ca²⁺ can significantly increase these rates. This information allows for effective use of silicon nanomembranes not only as active layers in eco/biodegradable electronics but also as



water barriers capable of providing perfect encapsulation until their disappearance by dissolution. The time scales for this encapsulation can be controlled by introduction of dopants into the Si and by addition of oxide layers on the exposed surfaces.

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The former possibility also allows the doped silicon to serve as an electrical interface for measuring biopotentials, as demonstrated in fully bioresorbable platforms for *in vivo* neural recordings. This collection of findings is important for further engineering development of water-soluble classes of silicon electronics.

KEYWORDS: bioresorbable electronics, transient electronics, silicon dissolution, encapsulation, water barrier

ne of the most attractive features of modern electronic devices is their ability to operate in an almost perfectly reliable fashion, with little to no degradation over exceptionally long periods of operation. Transient electronics is a class of technology designed to offer the opposite behavior, in which the constituent materials disappear, disintegrate, or otherwise change in some controlled physical or chemical way after a period of stable function. A subset of transient technology involves biologically and environmentally resorbable devices that dissolve in biofluids or groundwater into benign end products. Although certain classes of passive biomedical devices such as resorbable sutures and polymer matrices for drug release adopt such transient characteristics, it is only through recent research that similar modes of operation are now possible with active electronic systems, including those that use high-performance semiconductors, conductors, and dielectrics.^{6–8} Envisioned applications include bioresorbable diagnostic/therapeutic devices that avoid long-term adverse effects and secondary surgical procedures and eco-resorbable consumer gadgets that minimize the costs and health risks associated with waste disposal and recycling.

Of the various biodegradable electronic materials that now exist as options in transient electronics, device-grade, monocrystalline silicon is notable because of its potential to support high-performance devices that align, at least partly, with conventional, nontransient technologies. Recent studies of dissolution of monocrystalline silicon nanomembranes (Si NMs) in aqueous environments provide critical information on rates of hydrolysis in physiological and environmental conditions. Experimental and theoretical efforts offer insights into important characteristics and mechanisms of this chemistry. Specifically, the data suggest that silicon undergoes hydrolysis with catalyzing effects of ions present in the surrounding water, to generate silicic acid (Si(OH)₄) and hydrogen as final products. 1,3,6,9 Direct measurements of H₂ evolution and the Si dissolution rates support the stoichiometry of the proposed reaction (Si + $4H_2O \rightarrow$ $Si(OH)_4 + 2H_2$). The process proceeds without reactive diffusion of water into the silicon or of water permeation through defect sites, as indicated by the linear dependence of the thicknesses of Si NMs on reaction time. Rates of 1-100 nm/day occur at near neutral pH in saline solutions, depending on detailed experimental conditions, consistent with historical studies of silicon etching in alkaline and HF-containing etchants, where variations in step/defect densities, distributions of interface energy states, surface oxides, and metallic/organic contamination can affect the observations. ^{10–15} Additionally, doping with boron and phosphorus dramatically decelerates the dissolution at concentrations above 10¹⁹ cm⁻³, consistent with etching behavior in high-pH solutions.9 Here, we extend these collective studies to examine effects of proteins and divalent cations (Ca²⁺ and Mg²⁺) and the role of silicic acid (Si(OH)₄) as a product of dissolution and a ubiquitous component of natural and biological fluids, commonly at concentrations between 10⁰ and 10¹ ppm.

The slow rates of dissolution and the perfect monocrystalline nature of the Si NMs also enable their use as bioresorbable water barriers to encapsulate implantable electronic systems.

Compared to alternatives based on spin-cast polymers and inorganic materials (oxides, nitrides, etc.) formed by physical or chemical vapor deposition, 1,16,17 Si NMs ensure zero water transmission but retain bioresorbable forms, where submicron thicknesses are sufficient for device lifetimes in the range of several days to years. Additional results presented here demonstrate these capabilities, along with two strategies for extending the lifetime: decelerating the dissolution rate by use of doping at high concentrations and delaying the initiation of dissolution by use of thin layers of SiO₂ grown on the silicon surface. Reactive molecular dynamics simulations provide insights into the reactions that occur at the Si/water and SiO₂/water interfaces. Experiments show the ability to form flexible electrode arrays using thin patterned Si NMs on Mo, in fully bioresorbable formats for in vivo neural recordings.

RESULTS AND DISCUSSION

Figure 1 summarizes the rates of dissolution of Si NMs in phosphate buffered saline (PBS, 1x) with different concentrations of protein, Si(OH)₄, and cations. Figure 1a describes reductions in thickness of Si NMs (p-Si, boron, 8.5−11.5 Ω·cm (10¹⁵ cm⁻³), (100), SOITEC) by hydrolysis in PBS solutions with different albumin concentrations at 37 °C. The experiments involve 0.25 mm² pieces of silicon-on-insulator (SOI) wafers that have a top silicon layer (i.e., the Si NM) with 200 nm thickness immersed in 30 mL of solution immediately after exposure to UV ozone and buffered oxide etchant, to form a hydrogenterminated surface. Addition of bovine serum albumin into PBS provides access to a wide range of protein concentrations relevant to those in natural and biological fluids; the latter have concentrations between 10⁻² and 10² g/L (~80 g/L in blood, \sim 20 g/L in interstitial fluids, 0.1–1 g/L in saliva and aqueous humor, 0.01–0.1 g/L in urine). 18,19 Analysis of spectroscopic reflectance data collected from the SOI wafer determines the thickness of the Si NM (Figure S1a). Figure 1b compares rates of dissolution of Si as a function of albumin concentration in PBS. The values fall between 59 ± 0.5 (error defined as error in slope in linear fitting of thickness average at times) and 35 ± 0.4 nm/day. The deceleration likely follows from adsorption of protein on the silicon surface (0.5 to 1.5 nm, slightly varying with a kind of protein) and its resulting effects on limiting diffusive access of water to the surface. ^{20–22} Although the current studies focus on albumin, which is the most abundant protein in blood plasma, protein-specific effects on the deceleration of Si dissolution may exist due to the differences in their interaction to the Si surface.

Figure 1c compares the rates of dissolution as a function of $[Si(OH)_4]$ in PBS with three different albumin concentrations at 37 °C. Here, the $Si(OH)_4$ results from a Si standard solution (Si 1000 ppm, Sigma-Aldrich) prepared by dissolving high-purity silicon metal in 2% NaOH solution. After dilution to 100 ppm of Si, adjusting the pH and salt composition returns these values to those of the original PBS solution. In contrast to extensive data on the role of $Si(OH)_4$ in dissolution and condensation of silica, $^{23-26}$ its influence on the dissolution of Si has not been examined previously. The data in Figure 1c show reductions in the rates with increasing $[Si(OH)_4]$ at all protein concentrations,

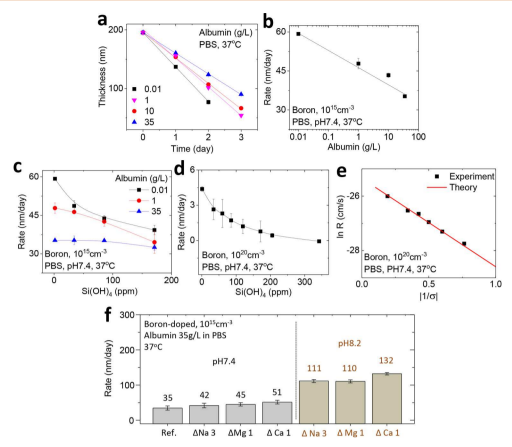


Figure 1. Dependence of the rate of dissolution of Si on the presence of protein, silicic acid, Ca^{2+} and Mg^{2+} in phosphate buffered saline (PBS, pH 7.4). (a) Change in thickness of p-Si NMs (boron, 10^{15} cm⁻³) as a function of time during dissolution. (b) Dependence of the dissolution of p-Si on the presence of protein. (c) Dependence of the dissolution of p-Si on the presence of $Si(OH)_4$ with different protein concentrations. (d) Dependence of the dissolution of p⁺-Si (boron, 10^{20} cm⁻³) on the presence of $Si(OH)_4$. (e) Polynuclear dissolution model for p⁺-Si dissolution. (f) Cation-specific effects on the rate of dissolution of p-Si.

qualitatively consistent with mechanistic reasoning based on the chemistry of Si hydrolysis (Si + $4H_2O \rightleftharpoons Si(OH)_4 + 2H_2$). Also, an increased dependence of the rate on [Si(OH)₄] follows reduced protein concentration, that is, for (0.01g/L), 59 ± 0.5 to $39 \pm 1.9 \text{ nm/day}$; for (35g/L), 39 ± 0.4 to $33 \pm 1.1 \text{ nm/day}$. Notably, the rate is non-negligible even in solutions at the saturation level of $Si(OH)_4$ (~185 ppm of $Si(OH)_4$ at 37 °C),²⁷ where this species starts to polymerize to SiO2, due to rates of Si hydrolysis that are faster than those associated with conversion of Si(OH)₄ to Si or SiO₂ with 10^0-10^2 ppm of Si(OH)₄ concentrations. By comparison, the rates of dissolution of SiO₂ have enhanced dependence on [Si(OH)₄], with values that approach zero at the same Si(OH)₄ saturation level.²³⁻²⁶ (Experiments in protein-free, purified PBS solutions show large variations and spatial nonuniformities in the dissolution rates depending on detailed conditions, likely due to uncontrolled levels of organic contaminants in ambient conditions. 13,28 The addition of small amounts of organic molecules such as proteins or phenol red as a pH indicator yields reproducible rates across individual samples and between experiments.)

Figure 1d summarizes the dependence of the rate of dissolution of highly doped Si (p⁺-Si, boron, 10^{20} cm⁻³) on Si(OH)₄ as an additional point of comparison. The rates decrease rapidly with increasing [Si(OH)₄] and virtually cease at ~200 ppm of Si(OH)₄, similar to the behavior of SiO₂. The slow rates of dissolution and the strong dependence on Si(OH)₄ for this case can be explained by the presence of a surface layer of SiO₂, as

supported by previous ellipsometric measurements that show a stable oxide on highly doped Si (\sim 1 nm) and barrier-less oxidation pathways in the presence of both boron and phosphorus impurities. Lightly doped Si, by comparison, has a porous passivation layer that is permeable to water molecules and that continuously grows in parallel with Si dissolution. Figure 1e presents fits of experiment data to a polynuclear dissolution model previously used to describe the dissolution of amorphous SiO₂ in the presence of Si(OH)₄. This model explains the dependence of SiO₂ dissolution on [Si(OH)₄] by reduced step propagation rates and increased activation energy for the formation of vacancy (or highly reactive) sites where dissolution starts to spread. According to the model, the rate can be expressed as

$$\ln R = \ln(hs \cdot A) - \frac{\pi \alpha^2 \omega h}{(kT)^2} \cdot \frac{1}{|\sigma|}$$

where R is the volume of dissolution nuclei formed per unit area of surface and time, h is the height of the vacancy, S is the surface area affected by each nucleation event, A is a pre-exponential factor for nucleation rate, ω is the specific volume of the molecule, α is the local interfacial free energy, k is Boltzmann constant, and T is the temperature. The degree of undersaturation is $\sigma = \ln(C/Ce)$, where C and Ce (185 ppm) are experimental and equilibrium concentrations of $Si(OH)_4$, respectively. The equation indicates that the logarithm of the rate of removing a unit of material from the surface has a squared dependence on the interfacial free energy (α) to create a new surface with highly reactive

Si atoms exposed. Considering the presence of a surface oxide layer on silicon and dissolution of this oxide, 32 we used parameters of $\omega_{\rm SiO_2}=4.54\times10^{-23}\,\rm cm^3/molecule$, height $h_{\rm SiO_2}=2.17\,\rm Å$, and $T=310.15\,\rm K$ in simulations for the dissolution of amorphous SiO₂. 26 Fitting experimental data for highly doped Si, as shown in Figure 1e, to this model yields an interfacial free energy $\alpha=40\,\rm mJ/m^2$, slightly lower than that of amorphous colloidal/biogenic silica and approximately a factor of 2 smaller than that for high-density SiO₂ such as quartz or thermally grown oxide. 26 By comparison, using parameters of Si crystal structures $(\omega_{\rm Si}=1.99\times10^{-23} \rm cm^3/molecule$, $h_{\rm Si}=1.36\,\rm Å)$ in the fit yields an unlikely large value of 84 mJ/m².

Accelerated dissolution occurs at high pH and in the presence of divalent cations (Ca²⁺ and Mg²⁺), as in Figure 1f. Experiments to define the latter effects involve 1 mM of CaCl₂ and MgCl₂ separately added to protein-containing PBS solutions from their 1 M standard solutions. A concentration of 1 mM is comparable to that found in naturally occurring biofluids. Figure 1f describes the accelerating effect of Ca²⁺ at pH values between 7.6 and 8.2. Solutions containing CaCl₂ dissolve Si faster than the solutions with NaCl or MgCl₂ at the same ionic strength. Since the [Cl⁻] in CaCl₂ and MgCl₂ solutions is lower than that in a NaCl solution at the same ionic strength, the observed acceleration in CaCl₂ solution can be attributed to the cation. The results follow similar trends reported for dissolution of quartz and silica, where Ca²⁺ accelerates the hydrolysis of SiO₂ more effectively than Na⁺ at near neutral pH. 25,26 Here, Ca²⁺ may have an increased ability to deprotonate surface silanol groups formed by absorption of water to Si surface and/or to enhance the reactivity of water and siloxane groups as suggested in previous studies of SiO2 dissolution.^{34–37} A discussion of the process of forming silanol and siloxane groups on the Si surface appears subsequently.

Figure 2 summarizes results of Si NM dissolution in various types of saline solutions. Environmental and biological fluids cover a range of pH values and concentrations of ions and

organic compounds. For example, the pH of common biological and environmental solutions can be as low as 4.5 as in urine and sweat and as high as 8.4 as in carbonate-containing solutions such as seawater. In terms of salinity, rivers and oceans contain 0.1 g to >50 g/L solid contents mostly associated with dissolved cations (Na⁺, Ca²⁺, and Mg²⁺) and anions (Cl⁻, HCO₃⁻, SO₄²⁻, and silica). In extracellular biofluids, the concentrations are strictly regulated to within 8-20 g/L, mostly composed of NaCl. Biofluids also contain high-energy nutrients such as glucose and proteins for cell metabolism. Figure 2a,b shows changes in color and thickness associated with 100 μ m² patterns of Si NMs (boron, 10¹⁵cm⁻³) with thicknesses of 200 nm in different saline solutions. Here, artificial perspiration has lower pH and higher NaCl concentration than the other fluids (pH 4.5). Bovine serum contains salt compositions similar to those of PBS, but with additional bicarbonate (15-30 mM) divalent cations (~1 mM Mg^{2+} , Ca^{2+}) and various proteins (56 g/L in total) (see Table S1). Hank's balanced salt solutions (HBSS) at two different pH values (7.6 and 8.2) and in formulations with and without divalent cations (Ca²⁺, Mg²⁺) provide additional points of comparison. Figure 2b indicates that the lowest rates occur in perspiration due primarily to its low pH; the highest rates occur in pH 8.2 HBSS with Ca2+ and Mg2+, consistent with previous observations of enhanced dissolution by Ca²⁺. Figure 2c summarizes the rates in various types of environmental and simulated body fluids at room temperature and 37 °C. Water purified by reverse osmosis, free from ions and larger molecules, shows the lowest rates (0.01 nm/day at 25 °C, 0.2 nm/day at 37 °C), while tap water, known to contain 30-40 ppm of Na⁺ and Ca²⁺ and organic/ inorganic contaminants, leads to higher rates $(0.7 \pm 0.1 \text{ nm/day})$ at 20 °C, 2.6 \pm 0.4 nm/day at 37 °C). Artificial sweat shows slower rates of dissolution than tap water $(0.5 \pm 0.1 \text{ nm/day at})$ 37 °C) due to its low pH despite its high NaCl concentration. Serum dissolves Si even at higher rates (3.5 \pm 0.5 nm/day at 20 °C, 21 \pm 0.4 nm/day at 37 °C) than these water samples

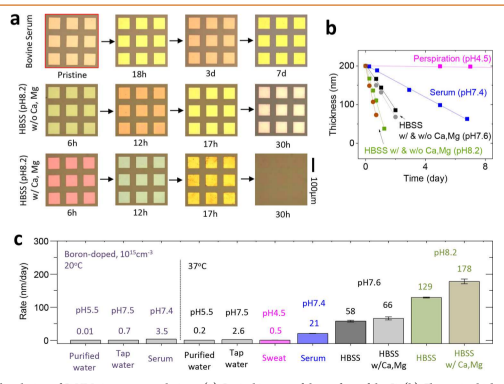


Figure 2. Rate of dissolution of Si NMs in aqueous solutions. (a) Optical images of the surface of the Si. (b) Change in thickness of p-Si NMs in simulated biofluids. (c) Rates of dissolution in different aqueous solutions.

due to its high salinity but slower rates than those of HBSS due to its protein content.

The observation that hydrolysis of Si NMs proceeds by surface reactions only, in a predictable manner without diffusion through defects or pinholes, creates the possibility for their use as temporary water barriers in bioresorbable electronic systems. Here, the defect-free, monocrystalline nature of the Si NMs leads to zero water transmission even for submicron thicknesses. Knowledge of the rates for dissolution allows design selection to match lifetime requirements for targeted applications. Soak tests using setups that incorporate thin films of magnesium (Figure 3a) underneath Si NMs facilitate evaluation of barrier properties owing to the visible corrosion of the Mg that follows contact with water. 38 Results obtained with spin-cast layers of poly(lactic-coglycolic acid) (PLGA, 5 μ m) and electron beam deposited films of SiO₂ (200 nm), instead of Si NMs, indicate almost immediate failure (<10 min) due to soaking in PBS (Figure S2) as a result of water permeation through the materials themselves and through pinhole defects, respectively. Figure 3b shows the rates of dissolution of Si, where a 1.5 μ m thick Si membrane survives for \sim 10 months with a dissolution rate of 4.8 \pm 0.2 nm/day at 37 °C. Figure 3c presents an image of an intact Mg pad captured after 2 months of soaking in PBS. Accelerated tests performed at 70 °C (Figure 3d) yield dissolution rates of 67 ± 0.2 nm/day and corrosion of the Mg pad abruptly at 23 ± 4 days.

Figure 4 presents two strategies to modulate the lifetime of this Si NM encapsulation without changing the thickness, the first of which exploits high doping concentrations (Figure 4a, p $^+$ -Si, boron, 10^{20} cm $^{-3}$). The dissolution of p $^+$ -Si follows the same dependence on salinity, Ca $^{2+}$ concentration, and pH as that of p-Si, but with overall slower rates. Significant deceleration (\sim 40×) in bovine serum compared to that in p-Si suggests that the p $^+$ -Si can enable encapsulation lifetimes of several years even with submicron thicknesses (\sim 180 nm/year). The second strategy

involves chemically functionalizing the Si surface to reduce its reactivity. Options in Si–C and Si–O chemistries provide a wide range of barrier lifetimes by delaying initiation of the Si dissolution. 39,40 A simple approach exploits growth of surface oxides at low temperatures by exposure to UV ozone (UVO) or O_2 plasma. The oxide thickness saturates at \sim 20 nm after 20 min exposure to O_2 plasma and at \sim 3 nm after 2 h exposure to UVO (Figure S3). Results of studies summarized in Figure 4b demonstrate a delayed initiation of Si dissolution by \sim 10 and \sim 30 days, with 3 nm UVO-grown oxide and 20 nm plasma-grown oxide, respectively. Relatively large variations in dissolution rate after initiation may originate from nonuniform thicknesses of the oxides, consistent with the surface morphologies observed by atomic force microscopy (Figure 4c).

Figure 5 presents results of reactive molecular dynamics (RMD) simulations of NMs of Si and SiO₂ (\sim 1.2 nm thickness) designed to reveal atomic scale interactions between Si atoms and water molecules. Here, the Si(100) NM consists of 6 atomic layers of Si with 100 Si surface atoms (atomic density 2.33 g/cm³, Figure 5a). The SiO₂ NM consists of 283 SiO₂ molecules in total (60 surface Si atoms) with amorphous structure (atomic density 2.33 g/cm³). Si atoms on the surface of each NM are initially terminated with dangling bonds without bonding to hydrogen or hydroxide and with occasional Si-O⁻ termination for the SiO₂ NM (16%). Figure 5b describes the water permeation event, as defined by transmission of a water molecule through the NM from one side of the water bath to the other side (see Figure 5a). No permeation is observed in Si NM over the simulation time of \sim 12 ns in 10 simulation replicas owing to the hermetic, crystalline structure of the material. Si surface atoms, initially with dangling bonds, form chemical bonds with OH- from water molecules, as shown in Figure 5b. Formation of Si-O-Si bonds occurs due to the nucleophilic attack of O from OH-terminated Si to nearby surface Si atoms with dangling bonds. Roughly 30%

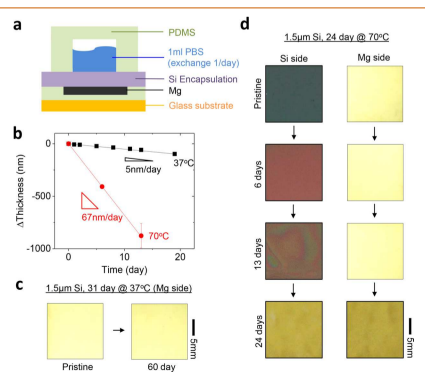


Figure 3. Si NMs as water-soluble water barriers. (a) Experimental setup in which a thin Mg pad allows direct imaging of the penetration of water through an overlying Si NM barrier layer. (b) Change in thickness with the operational condition. (c) Optical images of the Mg at various times at 37 °C. (d) Optical images of Si and Mg pad at 70 °C.

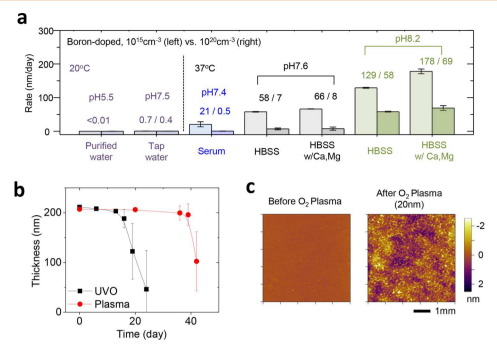


Figure 4. Strategies for controlling the lifetime of a Si NM barrier layer. (a) Rates of dissolution of p⁺-Si (boron, 10²⁰ cm⁻³). (b) Change in thickness of a p-Si NM in PBS. (c) Atomic force microscopy images of the surface of the p-Si.

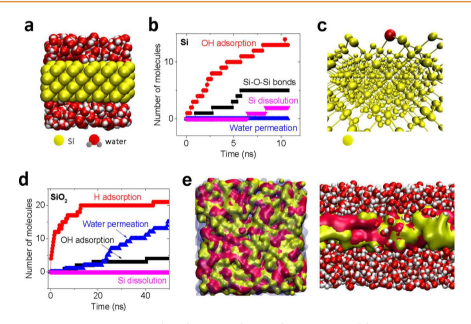


Figure 5. Reactive molecular dynamics simulation of Si (1 nm) and SiO₂ (1.25 nm) NMs in water. (a) Initial structure of the Si NM. (b) Reaction products for the Si NM. (c) Si–O–Si formation on the Si surface. (d) Reaction products for the SiO₂ NM. (e) Water permeation through the SiO₂ NM. Top view (left) and cross-sectional view (right).

of OH^- adsorbed on the Si surface creates Si-O-Si bonds, as shown in Figure 5c. The formation of $Si(OH)_2$ on the surface of the Si is also possible, which accounts for 42% of the OH^- adsorption. Dissociation of three Si atoms from the Si NM occurs from these $Si(OH)_2$ as a form of $Si(OH)_2^{2+}$, which eventually converts to $Si(OH)_4$ by interactions with surrounding water. Notably, no dissolution is observed from the Si atoms in Si-O-Si bonds, as expected based on the higher stability of SiO_2 against hydrolysis compared to Si. Concurrent oxidation and dissolution is consistent with a previous experimental report result on lightly doped $Si.^{32}$ Figure 5d compares the same type of dynamic changes in SiO_2 NMs. Due to the amorphous structure of

the slab, water begins to permeate through the SiO_2 NMs after a couple of nanoseconds. Water channels (Figure 5e) spontaneously form at 6 and 23 ns, as indicated by a sharp increase in water permeation in Figure 5d. The presence of initial voids in the SiO_2 structure, as distinct from the crystalline order in the Si NMs, leads to the channel formation. The average time for a single water molecule to permeate through the SiO_2 is 7.23 ns. Adsorption of H on the SiO_2 occurs due to rapid bond formation between H⁺ and oxygen-terminated Si atoms.

In addition to use as a bioresorbable barrier layer, Si NMs can simultaneously serve as the foundation for thin, flexible

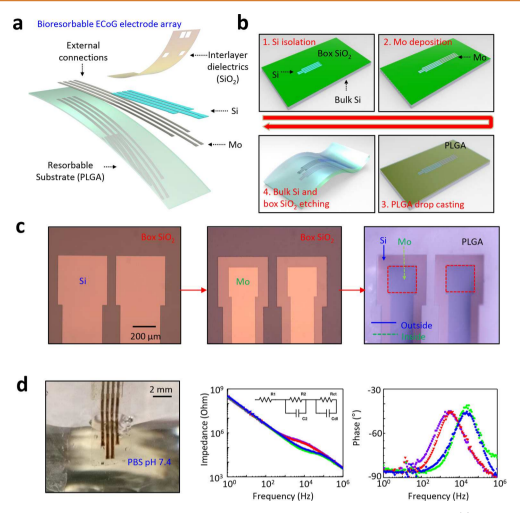


Figure 6. Thin, flexible neural electrode arrays with fully bioresorbable construction by use of doped Si NMs. (a) Schematic exploded-view illustration of a passive array for electrocorticography. (b) Fabrication process. (c) Optical images of the device. (d) Electrochemical impedance spectrum of four electrodes in PBS solution.

biosensing electrodes, as illustrated in Figure 6, with a demonstration that uses coatings of Mo underneath patterned Si NMs $(10^{15} \,\mathrm{cm}^{-3}, 1.5 \,\mu\mathrm{m})$. Mo metal without the Si NM on top undergoes comparatively fast dissolution, with a tendency to crack, fragment, and flake in biofluids (Figure S5). Figure 6a presents an exploded schematic diagram of the device structure. Here, a layer of SiO2 defines openings to the Si for four measurement sites for electrocorticography (ECoG) on rat cortex. A flexible sheet of the bioresorbable polymer (PLGA) serves as the substrate. The fabrication processes appear in Figure 6b. The first step involves isolation of the Si layer on an SOI wafer, followed by deposition of Mo electrodes. After drop-casting PLGA on the electrodes, back-etching of the bulk Si and buried SiO2 yields a flexible and bioresorbable neural sensing array. A final layer of 50 nm of SiO₂ defines the sensing areas (250 \times 250 μ m). Optical micrographs in Figure 6c highlight the structure of the electrodes. The area of Si NM encapsulation is slightly larger than that of the Mo electrodes to ensure protection against water penetration from the side edges. Electrochemical impedance spectra (EIS) collected at room temperature in PBS solutions simulate the interface characteristics between neural electrodes and brain tissues. The magnitude of the impendance and its phase angle for the four electrodes are in Figure 6d. At frequencies most relevant for neural sensing (near 1 kHz), the magnitude varies between 380 and 620 k Ω . The phase angle suggests capacitive (~90°)

behavior at low frequencies (1–10 Hz), with less capacitive behavior (~45°) at fequencies near 1 kHz. An equivalent Randles circuit with the addition of a resistor in parallel with a capcitor captures the impedance spectra of Si/Mo electrodes (Figure S4b). The additional capacitor might suggest a Schottky barrier at the Mo/Si interface. The interface charge transfer resistance, $R_{\rm ct}$ and double layer capacitance, $C_{\rm db}$ are found, from fitting, to be 2.2 G Ω and 0.45 nF, respectively. The lifetime of the current Si electrodes (10 15 cm $^{-3}$, 1.5 μ m) is expected to be ~50 days at 37 °C based on the abrupt change in the electrode EIS in accelerated tests at 70 °C at 4 days (Figure S5).

In vivo neural recordings with these types of bioresorbable neural sensors in adult rat animal models demonstrate good performance. The electrode array (4 sites) and a control electrode rest on the right barrel cortex of an anesthetized rat, as illustrated in Figure 7a. Representative 5 s segments of ECoG recordings of sleep spindle activity are shown in Figure 7b. The up—down dynamics at ~1 Hz with high-frequency oscillations occurring during the up states is typical of the sleep-like state induced by ketamine—dexmedetomidine anesthesia. Consistently, the power spectra of these recordings (Figure 7c) show two clear physiological peaks, one at 0.5–2 Hz (up—down state) and the other at 30–90 Hz (gamma-frequency oscillations in the up states), without discernible difference or signal attenuation in any frequency band between the bioresorbable and control electrodes.

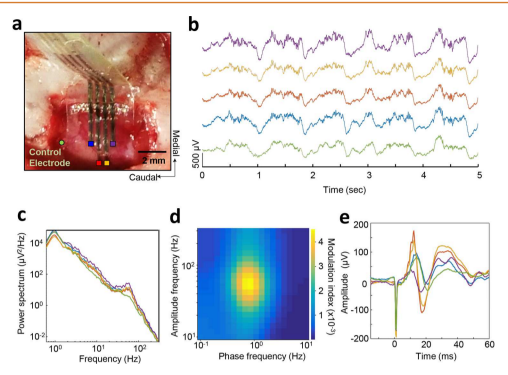


Figure 7. In vivo neural recordings in rats using a passive, bioresorbable electrode array. (a) Photograph of the electrode array on the right barrel cortex of an anesthetized rat. The colored boxes highlight the electrode contacts. A control electrode (green circle; electrode not present in photo) resides next to the next to the bioresorbable array. (b) Representative segment of cortical recordings from the four electrodes and the control electrode. (c) Power spectra of the recordings. (d) Comodulogram quantifying strong phase amplitude coupling between different neural oscillations. (e) Recorded cortical evoked response to electrical stimulation of a focal portion of the contralateral whisker pad. Sharp negative potential near 0 ms is the stimulus artifact.

The comodulogram in Figure 7d shows strong coupling between the phase of the up—down state and the amplitude of the gamma oscillations. The bioresorbable electrode can also be used for cortical recordings of somatosensory-evoked potentials (EPs). Figure 7e shows the cortical response to a focal electrical stimulation of the contralateral whisker pad recorded with the bioresorbable and the control electrodes. The response consists in a stimulus artifact near 0 ms followed by biphasic EPs at $\sim\!7$ ms latency, consistent with previous reports. Collectively, all recordings with the bioresorbable electrodes are comparable to those of the control electrode.

CONCLUSION

The results presented here reveal the dependence of the rate of dissolution of Si NMs on protein, Si(OH)₄ and Ca²⁺ in surrounding aqueous solutions. The results indicate a decelerating effect of Si(OH)₄ and proteins, and ion-specific accelerating effects of Ca²⁺. Uniform, controlled dissolution over large areas enables the use of Si NMs as bioresorbable water barriers, of utility in temporary electronic implants and environmental monitors. Strategies for extending the lifetimes of such types of barriers include the addition of dopants into the silicon or the growth of oxides onto its surface. The mechanical flexibility of doped Si NMs allows additional use as a bioresorbable electrode interfaces for biopotential measurements, as demonstrated through *in vivo* measurements of ECoG signals in rodent models. These collective findings have relevance to the further development of bio- and eco-resorbable forms of electronics.

EXPERIMENTAL METHODS

Dissolution of Si Nanomembranes. The time-dependent changes in the thickness of Si nanomembranes (200 nm initial thickness) on SOI

wafers (p-type (boron), 8.5–11.5 Ω -cm, (100), SOITEC) defined the dissolution rates. UVO cleaning (10 min) followed by removal of the surface oxide by immersion in buffered oxide etchant (2 min) prepared the silicon surface for the dissolution tests. ¹⁵ Soaking tests used various types of simulated biofluids including PBS (Corning cellgro), Hank's balanced salt solution (Thermo Fisher Scientific), bovine serum (RMBIO), and artificial perspiration (Pickering Laboratories) at room temperature (~20 °C) or at 37 °C. All solutions were stored in PETE bottles as purchased, and experiments were conducted in high-density polyethylene bottles (United States Plastic Corp.). The thickness of the Si NM was determined either by reflectometry (MProbe, Semicon Soft) or profilometry (Veeco/Sloan Dektak 3 ST, Plainview) after the soaking tests. Dissolution tests with highly doped silicon and with silicon coated with surface oxide followed the same procedures except for the silicon preparation steps. Highly doped silicon (p-type, boron, 10²⁰ cm³) was prepared by solid state diffusion (1050 °C, 15 min). Surface oxides were grown through exposure to O2 plasma (200 mT, 250 W, 20 min) or UV-induced ozone (10 min). Thickness of oxide was measured by spectroscopic ellipsometry (J.A. Woollam Co. Inc., USA). Atomic force microscopy (Asylum Research MFP-3D) defined the surface morphology of the SiO₂ grown by O₂ plasma before and after dissolution.

Si NM Encapsulation in Biofluids. Photolithographic patterning of electron beam evaporated layers of Ti/Mg (50/200 nm) defined traces on SOI wafers with top Si layers with thicknesses of 1.5 μ m. Casting a layer of polyimide (PI-2545, HD MicroSystems; 3.5 μ m) on the front side of such a substrate, laminating it against a glass substrate coated with poly(dimethylsiloxane) (PDMS; 10 μ m), and then inductively coupled plasma reactive ion etching (ICP-RIE, Surface Technology System) with gas flow of SF₆/O₂ with 40/3 sccm at a pressure of 50 mTorr removed the silicon wafer. Eliminating the buried oxide layer with HF left only the 1.5 μ m silicon membrane. A PDMS chamber bonded to the Si served to confine the PBS solutions. The solutions were replaced every day to avoid changes in dissolution rate associated with buildup of reaction products.

Reactive Molecular Dynamics Simulation. To model atomic scale interaction between Si/SiO₂ and water, RMD simulations were

performed with the Reaxff potential, 43 integrated in the large-scale atomic/molecular massively parallel simulator (LAMMPS) package.4 Previous studies show that the reactive potential developed for the Si/ SiO₂ interaction with water can reproduce the macroscale experimental properties of these interfaces.⁴⁵ The elements in our simulations are silicon, oxygen, and hydrogen. Two initial systems for Si and SiO2 NMs were created: Si NM with 602 silicon atoms (1.2 nm thickness) solvated with 3130 water molecules; SiO₂ NM with 283 SiO₂ molecules (1.25 nm thickness) solvated with 3119 water molecules. Periodic boundary conditions were used in all directions. Energy minimization of the system was performed for 1000000 steps. To ensure the capture of chemical reaction events between different atomic species, the time step was 0.1 fs. 46 The Nosé-Hoover thermostat maintained the temperature at T = 300 K. The simulation was run for 50 ns, and data were collected with the frequency of 0.1 ps. To compute the adsorption of H⁺ or OH⁻ groups in the simulations, we computed the root-mean-square displacement (rmsd) of H⁺ or OH⁻ between the simulation frames. At the adsorption event, the rmsd drops sharply from the average of 13.5 to 0.92 Å. Only H⁺ or OH⁻ with distance less than 2.8 Å from surface silicon atoms were considered adsorbed. To count the number of dissociated Si from the crystal structures, the rmsd of each Si is monitored in each simulation step. The rmsd of the silicon atom in the crystal structure is 1.1 Å. When a Si atom is dissociated, its rmsd sharply increases to >10 Å.

Fabrication of ECoG Arrays. The process began with thinning the handle substrate of an SOI wafer (1.5 μ m top Si layer) by mechanical grinding to a thickness of 150 μ m. The top Si on the SOI wafer was then patterned by photolithography (AZ 5214) and reactive ion etching (RIE) with SF₆ gas, to define the encapsulation layer. A thin film of Mo (300 nm) was deposited on the membrane by magnetron sputtering (AJA Orion 8 evaporation system, N. Scituate, MA) and patterned by photolithography (AZ nLOF 2070). Next, PLGA (~30 μ m, lactide/glycolide ratio of 75:25) was drop-cast on the top surface, and then the wafer and buried oxide were removed by etching with XeF₂ (XACTIX system) and buffered oxide (BOE, 6:1), respectively. Finally, a layer of SiO₂ was deposited by electron beam evaporation and patterned to define the sensing area for the neural electrodes (250 × 250 μ m). An ACF cable bonded to the terminal regions of the neural electrodes served as connection interfaces to external data acquisition (DAQ) systems.

EIS Measurements. Electrochemical impedance spectra of the ECoG neural electrodes were measured in PBS solutions to define the interface impedance characteristics (1-1 MHz, 10 mV amplitude, 0 V bias from open circuit voltage, Gamry Reference 600 potentiostat, EIS 300 frequency analyzer). Long-term accelerated soaking tests were also performed at 70 °C in PBS solutions to examine the effectiveness of Si NM encapsulation. Neural electrodes with Si NM encapsulation were fabricated through similar fabrication steps, except that glass substrates with PDMS adhesive instead of PLGA films were bonded to SOI surface, to avoid possible deformation induced by PLGA polymer at elevated temperature. For Mo-only neural electrodes, Mo was deposited directly on glass substrates. Parylene (1 μ m) instead of SiO₂ was deposited as a final layer to define the electrode sensing area, to minimize water permeation to the other areas of the electrodes. ACF cable was bonded to the terminal regions of the neural electrodes to connect to the potentiostat. Optical micrographs and EIS were collected at different soaking times, to compare the lifetime of neural electrodes with and without Si encapsulation.

In Vivo Neural Recording. The following procedures were approved by the Institutional Animal Care and Use Committee of the University of Pennsylvania. One 225 g Sprague—Dawley rat was anesthetized with a ketamine (60 mg kg $^{-1}$) and dexdomitor (0.25 mg kg $^{-1}$) solution and placed in a stereotaxic frame. A craniotomy was performed to expose the right barrel cortex. A skull screw was placed in the left parietal bone to serve as the reference electrode for the recordings. The recording array was placed over the exposed cortical surface. A control electrode (200 μ m diameter, tungsten) was placed on the cortical surface caudal to the array. Simultaneous neural recordings from the five electrodes were performed with a commercial electrophysiology system (Tucker-Davis Technologies) at a sampling frequency of 25 kHz. To elicit cortical evoked

responses, a pair of needle stimulating electrodes were inserted into the left mystacial pad. Brief electrical pulses ($500\,\mu\text{A}$, 1 ms, monophasic) were used to activate the intrinsic muscles of the vibrissae, causing a visible protraction of a whisker.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.7b06697.

Figures that summarize the chemical composition of the simulated biofluids; data on the spectral reflectance and surface profile of the SOI wafer; the encapsulation of the Mg pad with PLGA and electron-beam deposited SiO₂; the SiO₂ growth by exposure to UV ozone and O₂ plasma; EIS and surface morphologies of Mo and Mo/Si electrodes (PDF)

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Notes

The authors declare no competing financial interest.

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