Structural Properties and Sensing Performance of CeYₓOᵧ Sensing Films for Electrolyte–Insulator–Semiconductor pH Sensors

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In this study we developed CeYₓOᵧ sensing membranes displaying super-Nernstian pH-sensitivity for use in electrolyte–insulator–semiconductor (EIS) pH sensors. We examined the effect of thermal annealing on the structural properties and sensing characteristics of the CeYₓOᵧ sensing membranes deposited through reactive co-sputtering onto Si substrates. X-ray diffraction, atomic force microscopy, and X-ray photoelectron spectroscopy revealed the structural, morphological, and chemical features, respectively, of the CeYₓOᵧ films after their annealing at 600–900 °C. Among the tested systems, the CeYₓOᵧ EIS device prepared with annealing at 800 °C exhibited the highest sensitivity (78.15 mV/pH), the lowest hysteresis voltage (1.4 mV), and the lowest drift rate (0.85 mV/h). Presumably, these annealing conditions optimized the stoichiometry of (CeY)O₂ in the film and its surface roughness while suppressing silicate formation at the CeYₓOᵧ–Si interface. We attribute the super-Nernstian pH-sensitivity to the incorporation of Y ions in the Ce framework, thereby decreasing the oxidation state Ce(IV)→Ce(III) and resulting in less than one electron transferred per proton in the redox reaction.

Due to increasing demands in environmental protection, health care and bio-nano technology, chemical- and bio-sensors based on semiconductors attract much attention as they can be miniaturized and integrated into small size electronic equipment for real time detection with high sensitivity. A dramatic change in the miniaturization of ion-selective membrane technology occurred when Bergveld demonstrated an ion-sensitive field-effect transistor (ISFET) by modifying a metal-oxide-semiconductor field-effect transistor (MOSFET) to expose the gate oxide film to an electrolyte solution. The theoretical chemical response of an ISFET sensor is the same as that of an ion-selective electrode. This sensor is replaced the fragile glass electrode with various gate oxides to detect a variety of ions. Since then, ISFETs offered several major advantages, including small size, rapid response, low output impedance, multiple ion sensor capability, and high compatibility for integration with complementary metal-oxide-semiconductor (CMOS) systems. Apart from ISFET, electrolyte-insulator-semiconductor (EIS) sensors without source and drain electrodes have relatively simpler structures and are easier to fabricate than ISFET devices. It is based on a MOSFET device without a metal gate for direct immersion in a buffer solution, the gate insulator material is a critical part of an ISFET or EIS device. The first sensitive pH response was attained with a SiO₂ membrane. Later, several metal oxide materials (e.g. Al₂O₃, Ta₂O₅, SnO₂, WO₃) were used as sensing membranes because these materials possess large surface areas and good sensing performance. In addition to conventional metal oxides, a number of refractory oxides having high-dielectric constants (high-k), such as TiO₂, ZrO₂, HfO₂, and Y₂O₃, have been received considerable attention as pH-sensitive membranes for promising applications in ISFET and EIS devices due to their good sensing properties. Nevertheless, their structures are generally unstable and very reactive for long-term and higher temperature applications as a result of the formation of a interfacial layer between the high-k metal oxide and the Si substrate in their manufacture. The interface layer can affect not only the performance an EIS sensor but also the sensor inoperable. Consequently, it is needed to find new materials and alternative processes to diminish interfacial defects.

Rare earth (RE) oxide films are possible replacements for traditional SiO₂ as the gate dielectric in advanced CMOS devices because of their large values of k, lower leakage currents, high resistivities, and good thermal...

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decreasing the oxidation state of Ce (Ce⁴⁺ → Ce³⁺) in a pH range of 5.5–7.5. We attribute this super-Nernstian response to the incorporation of Y ions into the Ce framework, thereby reducing the number of electrons transferred per proton. Several researchers in literature have reported the incorporation of other elements (e.g., Ti, TiO₂, Y) into RE oxide films to achieve lower pH sensitivities and excellent thermal stabilities. Among them, CeO₂ thin films have several advantages, including strong adhesion, high refractive index, good redox, high mechanical strength, excellent thermal stability, for applications in humidity sensors, fuel cells, and gas sensors. Ansari and co-workers also demonstrated that the nanostructured CeO₂ film was deposited on Au electrode using the sol-gel method for glucose sensor. Besides the above properties, they possess a high dielectric constant, a small lattice mismatch with silicon, and good thermodynamic stability in contact with silicon, making them particularly attractive materials for use as alternative gate dielectrics in nano-CMOS, flash memory, and EIS devices. The major concern when using RE oxide films as sensing membranes, however, is prone to moisture absorption due to their hygroscopic nature, which degrades permittivity through the formation of low-permittivity hydroxides. The moisture absorption of RE oxide films stems from the presence of oxygen vacancies in the films. Several researchers in literature have reported the incorporation of other elements (e.g., Ti, TiO₂, Y) into RE dielectric films, which suppress the moisture absorption of the RE oxide. In this concern, we have reported Tidoped RE oxide sensing membranes in our earlier works which addresses the high hygroscopic nature of the RE oxide. The structural and sensing properties of CeYₓO₂ films as sensing membranes for EIS pH sensors have yet to be studied. In this work, we propose a CeYₓO₂ sensing film displaying super-Nernstian pH-sensitivity for use in EIS pH sensors. We have investigated the effects of thermal annealing on the structural properties and sensing characteristics of CeYₓO₂ thin films deposited on Si substrates through reactive co-sputtering. Moreover, we have employed X-ray diffraction (XRD), X-ray photoelectron spectroscopy (XPS), and atomic force microscopy (AFM) to monitor the quality of the CeYₓO₂ sensing films and determine the optimal annealing conditions. We measured the sensing characteristics (pH sensitivity, hysteresis, and drift) of these CeYₓO₂ films. The sensitivity of one such CeYₓO₂ EIS sensor was higher than that expected from the Nernst law. We attribute this super-Nernstian response to the incorporation of Y ions into the Ce framework, thereby decreasing the oxidation state of Ce (Ce⁴⁺ → Ce³⁺) and forming a mixed state of oxidized and reduced CeO₂ in the solution; as a result, less than one electron per proton would be transferred in the redox reaction.

**Methods**

**Fabrication.** EIS device structures incorporating CeYₓO₂ sensing films were fabricated on 4-in p-type (100) Si wafers. Prior to deposition of the CeYₓO₂ film, the wafer was cleaned through standard RCA cleaning and then treated with 1% HF to remove the native oxide. Subsequently, ~60 nm CeYₓO₂ film was deposited on the Si substrate through rf co-sputtering from cerium oxide and yttrium targets with a mixture of Ar and O₂ (Ar:O₂ = 25:5) ambient during sputtering. During the sputtering of CeYₓO₂ film, the rf power of CeO₂ and Y targets was 150 and 150 W, respectively. The chamber pressure was maintained at ~10⁻² Torr and the deposition rate of the film was ~1.5 nm/min. The samples were subjected to rapid thermal annealing (RTA) in O₂ ambient for 30 s at 600, 700, 800, or 900°C to achieve a crystalline (CeYₓO₂) film. To form a backside contact, the backside SiO₂ layer was removed in a buffer oxide etchant (BOE) and then an Al film (400 nm) was deposited using a thermal evaporator. To define the sensing area of the deposited CeYₓO₂, an automatic robotic dispenser was used with an adhesive silicone gel (S181) acting as a segregating layer. The EIS device was assembled on a Cu wire on a custom-made printed circuit board by silver glue. To avoid leakage of the electrolyte, adhesive epoxy was encapsulated between EIS film and the Cu. The structure of the fabricated CeYₓO₂ EIS device is illustrated in Fig. 1.

**Characterization.** After RTA, the structural and compositional properties of the CeYₓO₂ sensing films were investigated using a combination of XRD and XPS. The XRD patterns were recorded on a Bruker D8 discover diffractometer with Cu Kα radiation (λ = 0.1541 nm) in a scanning angle (2θ) range of 20–60° at a scanning speed of 1° min⁻¹, operated at 40 kV and 40 mA, respectively. XPS was conducted using a Physical Electronics Quantum 2000 scanning ESCA microprobe, with a focused monochromatic Al Kα X-ray (1486.7 eV) source for excitation and a spherical section analyzer. Curve fitting of XPS spectra was performed using a Shirley background and

![Schematic representation of the structure of a CeYₓO₂ EIS sensor.](image-url)
The intensity of the (CeY)O$_2$ (111) peak increased upon increasing the RTA temperature, consistent with a more crystalline Ce$_x$Y$_{1-x}$O$_2$ structure. A strong (CeY)O$_2$ (111) peak, (CeY)O$_2$ (200) peak, and two weak (CeY)O$_2$ (220) and (311) peaks. RTA improved the crystalline structure of the (CeY)O$_2$ films. The intensity of the (CeY)O$_2$ (111) peak increased upon increasing the RTA temperature, consistent with a more crystalline Ce$_x$Y$_{1-x}$O$_2$ structure. A strong (CeY)O$_2$ (111) peak, (CeY)O$_2$ (200) peak, and two weak (CeY)O$_2$ (220) and (311) peaks appeared in the patterns of the samples that had been annealed at 800 and 900 °C, suggesting the formation of a stoichiometric (CeY)O$_2$ film. We attribute this annealing behavior to the Y atoms reacting with CeO$_2$ to provide a more stoichiometric Ce$_x$Y$_{1-x}$O$_2$ structure.

The morphology of the Ce$_x$Y$_{1-x}$O$_2$ films prepared at various temperatures is revealed by AFM images. Figure 3 displays AFM surface images of Ce$_x$Y$_{1-x}$O$_2$ films that had been annealed at different temperatures. The surface roughness (rms) values of 1.77, 1.81, 1.92, and 252 nm were determined for the Ce$_x$Y$_{1-x}$O$_2$ films treated at 600, 700, 800, and 900 °C, respectively. It can be seen that the surface roughness value increased with increasing the annealing temperature. This behavior is attributed to the aggregation of the native grains into the larger clusters upon annealing. This different cluster size affects the surface morphology of the films. The surface roughnesses of the Ce$_x$Y$_{1-x}$O$_2$ films annealed at the various RTA temperatures were consistent with the intensities of the diffraction peaks in Fig. 2; therefore, the AFM and XRD data are in good agreement. Figure 3(d) depicts that the Ce$_x$Y$_{1-x}$O$_2$ film annealed at 900 °C had larger clusters and becomes rougher. We suspect that self-diffusion of Ce and Y atoms increased during high-temperature annealing, enhancing the clustering of grains and, thereby, increasing the surface roughness of the Ce$_x$Y$_{1-x}$O$_2$ film. An increase in grain size might improve the sensing performance when the film was used for the membrane. In contrast, Fig. 3(a) demonstrates that the Ce$_x$Y$_{1-x}$O$_2$ film at 600 °C featured smaller grains and showed smoother surface.

To further investigate the surface composition and chemical states, the Ce$_x$Y$_{1-x}$O$_2$ films after RTA at various temperatures were performed by XPS technique. The Ce 3d, Y 3d, and O 1s spectra of the Ce$_x$Y$_{1-x}$O$_2$ films prepared at different temperatures are shown in Fig. 4(a), (b) and (c), respectively. The Ce 3d$_{5/2}$ spectra in Fig. 4(a) can be deconvoluted into three peaks: $\nu$ (~882.6 eV), $\nu'$ (~884.7 eV) and $\nu''$ (~888.9 eV). The 3d$^{10}$4f$^0$ state of Ce$^{3+}$ species are labeled as $\nu$ and $\nu''$, whereas the 3d$^{10}$4f$^1$ state of Ce$^{4+}$ is represented as $\nu''$. The binding energy of the Ce 3d$_{5/2}$ peak at 882.1 eV is taken as energy reference, while at 885.3 eV is assigned to the Ce$_2$O$_3$ structure. Three $\nu$, $\nu'$ and $\nu''$ peak binding energies exhibit quite similar evolutions, indicating no samples

Figure 2. XRD patterns of Ce$_x$Y$_{1-x}$O$_2$ films that had been annealed at various temperatures.
From Fig. 4, it is found that the main chemical valence of cerium for +4 oxidation state and a small amount of +3 oxidation state co-existed in the film. The effective ionic radii of Ce$^{4+}$, Ce$^{3+}$, and Y$^{3+}$ are 0.97, 1.14 and 1.01 Å, respectively. The +3 oxidation state of cerium was introduced by the spontaneous transformation of smaller radius Ce$^{4+}$ due to larger radius Ce$^{3+}$, the addition of smaller radius Y$^{3+}$ can compensate the lattice expansion induced by larger Ce$^{3+}$ ions. A shift of the v peak position toward lower binding energy, from 882.6 to 882.3 eV, prepared upon increasing the RTA from 600 to 900 °C, indicative of reactions of the Ce and Y atoms with O atoms to form a stoichiometric (CeY)O$_2$ structure. For the film annealed at 600, 700 and 800 °C, the $v'$ and $v''$ peak positions of were fixed at 884.7 and 888.9 eV, but 900 °C-annealed sample suddenly shifted toward lower binding energy by 0.3 and 0.2 eV, respectively. It was thus interpreted that the deposited CeY$_x$O$_y$ film undergoes solid-phase intermixing with Si from the substrate during high-temperature annealing to form a Ce-silicate layer. The silicate layer is speculated to result from reactions between silicon and CeO$_x$ formed at the surface of oxide film and Si substrate. The Y 3d$_{3/2}$ and 3d$_{5/2}$ double peaks of the Y$_2$O$_3$ reference appeared at 158.5 and 156.7 eV, respectively. Figure 4(b) displays these signals for the film annealed at 600 °C, at 158.9 and 157 eV, respectively, suggesting a poor (CeY)O$_2$ structure incorporating Y in the form of CeY(OH)$_x$, which probably formed at the surface of the sample during its exposure to air. The positions of the Y 3d$_{3/2}$ and 3d$_{5/2}$ peaks shifted to higher binding energies upon increasing the annealing temperature. For the sample subjected to RTA at 800 °C, the binding energies of the Y 3d$_{3/2}$ and 3d$_{5/2}$ orbitals were 159.1 and 157.2 eV, respectively, reflecting the presence of (CeY)O$_3$ structures. The intensities of the Y 3d$_{3/2}$ and 3d$_{5/2}$ split peaks provided by CeY$_x$O$_y$ remained quite constant for annealing temperatures of up to 800 °C, but the former increased suddenly while the latter decreased after annealing at 900 °C. This phenomenon is consistent with the reactions of Ce atoms with O and Si atoms to form a thicker Ce-silicate layer. Figure 4(c) presents the O 1s spectra for the annealed CeY$_x$O$_y$ films, with appropriate curve-fitting of the peaks. The O 1s peak of the Ce$_2$O$_3$ reference appeared at 530.6 eV. In the three spectra, the O 1s peaks at 533.4, 531.5 (531.4 eV for 700 and 800 °C, 531.6 eV for 900 °C), and 530.6 eV (530.7 eV for 800 and 900 °C) represent the Ce–OH, Ce–O–Y, and Ce–O bonds, respectively. For 600 °C-annealed sample, the O

![Figure 3. AFM surface images of CeY$_x$O$_y$ sensing films that had been annealed at (a) 600, (b) 700, (c) 800, and (d) 900 °C.](image-url)
1 s peak intensities provided by Ce(OH)ₓ was larger compared to other annealing temperatures, respectively. This result may be attributed a poor CeYₓOᵧ film to the hydroxide formation after exposure to an air ambient. The moisture absorption of RE oxide film may be attributed to the metal defects or oxygen vacancies in the films. The intensity of the O 1 s peak corresponding to (CeY)O₂ increased upon increasing the RTA temperature—except at 900 °C, where it decreased relative to the signal for Ce₂O₃. This behavior suggests that Ce and Y atoms reacted with O atoms to form a stoichiometric (CeY)O₂ film. For the CeYₓOᵧ component of the sample annealed at 900 °C, the O 1 s peak shifted to higher binding energy by 0.1 or 0.2 eV relative to those of the samples subjected to the other annealing temperatures. In addition, the intensity of the O 1 s peak for the Ce₂O₃ component of the sample annealed at 900 °C was weaker than other RTA temperatures. During high-temperature annealing, the Ce atoms diffused readily from the CeOₓ film to form a thicker Ce-silicate layer at the CeYₓOᵧ–Si interface.

**Sensing Characteristics.** The metal gate of the MOSFET is replaced by a reference electrode, while the liquid in which this electrode is present makes contact with the gate oxide. In case of an ISFET the gate voltage is the voltage at the reference electrode, usually grounded reference electrode, but the threshold voltage comprises also terms which reflect the interfaces between the liquid and the gate oxide on the one side and the liquid and the reference electrode at the other side. The threshold voltage (V_{th}) of an ISFET device is defined by the equation

\[
V_{th} = E_{ref} - \psi + \chi^{\text{int}} - \frac{\Phi_{Si}}{q} - \left( \frac{Q_{SS} + Q_{SC}}{C_{OX}} \right) - 2\phi_{fp}.
\]

where \(E_{ref}\) is the reference electrode potential relative to vacuum, \(\psi\) is the surface potential, which results from a chemical reaction, usually governed by the dissociation of oxide surface groups, \(\chi^{\text{int}}\) is the surface dipole potential of the solution, \(\Phi_{Si}\) is the Si work function, \(q\) is the elementary charge, \(Q_{SS}\) is the surface state density at the Si surface, \(Q_{SC}\) is the depletion charge in Si, \(C_{OX}\) is the gate oxide capacitance per unit area, and \(\phi_{fp}\) is the Fermi potential of semiconductor (p-type for the n-channel MOSFET considered here). All terms usually refer to constant but does not include \(\psi\), because this term makes the ISFET sensitive to the electrolyte pH, which is controlling the dissociation of the oxide surface groups. Designing a pH sensitive ISFET with a maximum sensitivity needs therefore a detailed study of the oxide–electrolyte interface to choose the best oxide film, which is not a priority for SiO₂ in MOSFETs. In an ISFET device, the change in the surface potential at the oxide–aqueous electrolyte interface will shift the threshold voltage. It is mainly related to the charge concentration, which can be interpreted using the site-binding model to characterize the properties of an oxide–electrolyte interface. The surface of metal oxide usually contains hydroxyl groups. These groups can donate or accept a proton from the solution, leaving a negatively charged or a positively charged surface group, respectively. The equilibrium reactions can occur between protons in the solution and the hydroxyl groups formed at the oxide–solution interface. The mechanism responsible for the oxide surface charge can be explained by the site-binding model, which describes the equilibrium between the so-called amphoteric Si-OH surface sites and the H⁺–ions in the solution. In the site-binding model, the terminal OH groups on the surface of the gate oxide can be neutral in the form (OH⁻), protonated (OH₂⁺; i.e., accepted a proton), or deprotonated (O⁻; i.e., donated a proton) at the oxide–electrolyte interface. At low pH (high proton concentration in solution), the equilibrium is shifted toward a protonated (i.e., positively charged) surface, which becomes a positive site. As a result, the C–V curve is shifted to the right direction. Around the point of zero charge, a general expression for the pH sensitivity of an ISFET sensor is given by

**Figure 4.** XPS spectra displaying the (a) Ce 3d, (b) Y 3d, and (c) O 1 s energy levels in CeYₓOᵧ films that had been annealed at various temperatures.
\[
\frac{d\psi}{dpH} = 2.303 \frac{kT}{q} \frac{\beta}{\beta + 1}
\]  
(2)

where \( k \) is Boltzmann’s constant, \( T \) is the absolute temperature, and \( \beta \) is a parameter that denotes the chemical sensitivity of the gate oxide and is proportional to the density of reactive sites, given by

\[
\beta = \frac{2q^2 N_s K_a K_b}{kT C_{DL}}
\]  
(3)

where \( N_s \) is the total number of surface sites per unit area, \( K_a \) and \( K_b \) are the equilibrium constants at the acid and base point, respectively, and \( C_{DL} \) is the double-layer capacitance derived from the Gouy–Chapman–Stern model. In this model, the double-layer capacitance consists of a series network of a Helmholtz-layer capacitance (the Stern capacitance) and a diffuse-layer capacitance. The Helmholtz layer model indicate that the ions in the solution have a finite size and the centers of the ions cannot approach the surface any closer compared with the ionic radius (including a possible water layer). The diffuse layer comprises the same amount of charge (of opposite sign) as the oxide surface charge, since the Helmholtz layer is by definition no containing any charge. If \( \beta \) parameter approaches infinite, the ISFET has a theoretical maximum value of accurately 59.6 mV/pH at 300 K, which is also called the maximum Nernstian sensitivity.

To achieve the high sensitivity of an EIS sensor, we evaluated the sensing performance of the CeY\(_x\)O\(_y\) EIS devices annealed at various temperatures. Figure 5(a–d) display the pH-dependence of a group of C–V curves for the CeY\(_x\)O\(_y\) EIS devices prepared with annealing at 600, 700, 800, and 900 °C, respectively, and measured at values of pH in the range from 2 to 12. The change in the number of hydrogen ions on the sensing films led to shifts in the normalized C–V curves. We attribute the kinks in the C–V curves to the presence of interface traps between the CeY\(_x\)O\(_y\) film and Si substrate. Here, the reference voltage (\( V_{REF} \)) for each pH buffer solution was the voltage required to obtain a normalized capacitance of 0.5. The insets to Fig. 5 (a–d) display the \( V_{REF} \) as function of pH values for the CeY\(_x\)O\(_y\) EIS devices that had been prepared with the various annealing temperatures. The

![Graphs showing pH-dependence of C–V curves for CeY\(_x\)O\(_y\) EIS devices](image-url)
pH-sensitivities of the CeY$_2$O$_3$ films after RTA at 600, 700, 800, and 900 °C were determined to be 59.98, 62.77, 78.15, and 73.68 mV/pH, respectively. Thus, among these tested systems, the CeY$_2$O$_3$ EIS device annealed at 800 °C exhibited the highest sensitivity, possibly because of the higher surface roughness (AFM) of its CeY$_2$O$_3$ sensing film. A higher surface roughness will increase the surface site density and, therefore, improve the detection sensitivity, as previously discussed for the site-binding model. Nevertheless, the sensitivity of the CeY$_2$O$_3$ film annealed at 900 °C was lower than that of the film annealed at 800 °C, presumably because of a thicker silicate layer formed at the CeY$_2$O$_3$–substrate interface after treatment at such a high temperature. Moreover, the pH-responses for the CeY$_2$O$_3$ EIS sensors were higher than that expected (59.6 mV/pH) from the Nernst law, possibly because the addition of Y ions into the Ce framework enhanced the decrease in the Ce oxidation state (Ce$^{4+}$ → Ce$^{3+}$). The wide application of ceria (CeO$_2$) is driven by its ability to store and release oxygen (by creating oxygen vacancy sites) due to the easily accessible redox cycle of cerium ions (Ce$^{5+}$ ↔ Ce$^{4+}$).

During the oxygen release process, the volume of the Ce compound increases in proportion to the change in the Ce oxidation state from Ce$^{4+}$ to Ce$^{3+}$. The stress energy resulting from this volume increase would enhance any further valence change of the Ce oxidation state. The introduction of Y ions into the Ce framework would compensate for the increase in volume and ease the change in valence of the Ce oxidation state. The potential response with respect to pH is related to the equilibrium between the oxidized and reduced forms of CeO$_2$. The hydroxyl groups can be considered as amphoteric sites, according to the site-binding theory, which can get or release a proton, for example:

$$\text{CeO(OH)} \rightleftharpoons \text{CeO}^- + \text{H}^+ \quad (4)$$

$$\text{CeO(OH)} + \text{H}^+ \rightleftharpoons \text{CeO(OH)}_2^+ \quad (5)$$

$$2\text{Ce(OH)}_3^- + \text{H}_2\text{O} \rightleftharpoons \text{Ce}_2\text{O(OH)}_6^- + 2\text{H}^+ + 2e^- \quad (6)$$

In Eqs (4–6), these reactions do not need electron transfer because the number of electrons transferred is equivalent to that of the protons (H$^+$). As a result, the pH-sensitivity is estimated to be 59 mV/pH when considering the redox potential. Nevertheless, pH sensitivities of over 59 mV/pH have been measured for other electrodes (e.g., IrO$_2$, Ta$_2$O$_5$). We suspect that a change in the Ce ion valence occurs during the reversible reaction from Ce$^{4+}$ to Ce$^{3+}$. The film can be considered only as an acidic oxyhydroxide, which the amphoteric character of the hydroxyl groups can be neglected, each state of the film with its own degree of proton dissociation:

$$\text{Ce(OH)}_3 \rightleftharpoons \text{Ce(OH)}_2^+ + \text{H}^+ \quad (7)$$

$$\text{Ce}_2\text{O(OH)}_6^- \rightleftharpoons \text{Ce}_2\text{O(OH)}_3^- + 3\text{H}^+ \quad (8)$$

Substitution of redox reactions (7) and (8) into reaction (9) yields

$$2\text{Ce(OH)}_2^- + \text{H}_2\text{O} \rightleftharpoons 2\text{Ce}_2\text{O(OH)}_3^- + 3\text{H}^+ + 2e^- \quad (9)$$

For this reaction, only two electrons are transferred for every three protons; in this case, the maximum slope for a Nernst equation could reach as high as 88 mV/pH. Thus, experimental observations of pH sensitivities of greater than 59 mV/pH were explained by considering mixed oxidized and reduced CeO$_2$ states that lead to less than one electron per proton being transferred in the redox reaction. In contrast, for pH-sensitivities of less than 59 mV/pH, more than one electron would be transferred per proton. In this study, our CeY$_2$O$_3$ films exhibited sensitivities to pH that are superior to those of sensing materials commonly used for ISFET and EIS sensors such as, Nb$_2$O$_5$ (52.12 mV/pH), Y$_2$O$_3$ (54.5 mV/pH), CeO$_2$ (58.76 mV/pH), ZnO$_2$ (58 mV/pH), Pr$_2$O$_3$ (52.9 mV/pH), Nd$_2$O$_3$ (56.01 mV/pH), Sm$_2$O$_3$ (56.2 mV/pH), Er$_2$O$_3$ (57.58 mV/pH), and Eu$_2$Ti$_2$O$_7$ (59.03 mV/pH).

The hysteresis and drift phenomena of a gate insulator influence the performance of ISFET and EIS sensors. A surface-site model can describe hysteresis characteristics when assuming that a fraction of the sites will respond slowly to changes in pH as a result of imperfections in the insulator film; these imperfections lead to porosity that produces interior sites that react with the ions in the solution. We tested our CeY$_2$O$_3$ EIS capacitors through a pH loop of 7 → 4 → 7 → 10 → 7 over a period of 25 min. We define the hysteresis voltage as the difference in reference voltage between the initial (first pH 7) and terminal (last pH 7) voltages in the pH loop. Figure 6 presents the hysteresis voltages of CeY$_2$O$_3$ EIS devices that had been prepared with annealing at various temperatures; they were immersed in buffer solutions having various values of pH in an alternating cycle (pH 7, pH 4, pH 7, pH 10, pH 7), for 5 min in each solution. The hysteresis voltage of the CeY$_2$O$_3$ EIS devices annealed at 600, 700, 800, and 900 °C were 126.3, 5.9, 1.4, and 40.9 mV, respectively. Thus, the EIS sensor annealed at 800 °C exhibited the smallest hysteresis voltage (1.4 mV), while that annealed at 600 °C had the highest (126.3 mV). Defects and vacancies in the films may have caused ions to attach to the surface and, thereby, delay the response to the reference voltage. Annealing at 800 °C could diminish the extent of crystal defects and, thereby, minimize the number of attached ions on the surface during the test.

According to the physical model for gate voltage drift proposed by Jamash et al., the overall capacitance of gate insulator (the series combination of the capacitance of the modified surface layer and the underlying insulator) should exhibit a slow, temporal variation. Therefore, the amount of inversion charge stored in the semiconductor at a given gate bias would slowly change over time, leading to a monotonic temporal change in the threshold voltage and, thus, in the drain current; this phenomenon is commonly known as “drift.”
modification of the insulator surface can lead to a variation in the number of surface sites over time, thereby resulting in a monotonic temporal increase in the threshold voltage. Figure 7 displays the drift characteristics of the CeY$_x$O$_y$ EIS devices prepared with annealing at the various RTA temperatures, measured in solution at pH 7 for 12 h. Here, the change in the reference voltage ($\Delta V_{REF}$) is defined as

$$\Delta V_{REF} = V_{REF}(t) - V_{REF}(0)$$  \hspace{1cm} (10)

The degradation slope of the reference voltage represents the EIS stability. The drift rates of the CeY$_x$O$_y$ EIS devices after RTA at 600, 700, 800, and 900 °C were 7.40, 1.76, 0.85, and 3.05 mV/h, respectively. Thus, the CeY$_x$O$_y$ EIS device annealed at 800 °C had the best long-term stability, while the device annealed at 600 °C exhibited a serious drift rate. The drift in the reference voltage might result from lattice defects (e.g., vacancies or dangling bonds) capturing clusters of ions. We suspect that the less-pronounced drift behavior after RTA at 800 °C arose from a lower density of crystal defects; that is, these annealing conditions could effectively remove such defects. In addition, a higher drift rate may be due to a larger number of crystal defects producing buried surface sites and/or traps.

**Conclusion**

We have observed a super-Nernstian response to pH from EIS devices incorporating CeY$_x$O$_y$ sensing membranes grown on Si substrates through reactive co-sputtering. XRD, XPS, and AFM confirmed the presence of (CeY)O$_2$ structures in these EIS devices. The pH-sensing performance of the EIS device incorporating the CeY$_x$O$_y$ film that had been annealed at 800 °C was superior to those of the devices subjected to other annealing
temperatures, including a higher sensitivity (78.15 mV/pH), a smaller hysteresis voltage (1.4 mV), and a lower drift rate (0.85 mV/h). This enhanced performance resulted from the formation of a stoichiometric (Ce,Y)O$_2$ film, a high surface roughness, and a low number of crystal defects after applying these annealing conditions. The super-Nernstian pH-response appears to have resulted from the Y component incorporated within the Ce framework enhancing the valence change of the Ce oxidation state, thereby causing less than one electron to be transferred per proton in the redox reaction, due to the mixed oxidized and reduced CeO$_x$ states in the solution.

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Author Contributions
TMP conceived experiments and data analysis and wrote manuscript. CWW and CYC assisted with experiment design and performed experiments and analyzed data. All authors discussed and commented on the manuscript.

Additional Information
Competing Interests: The authors declare that they have no competing interests.

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