The electrochemical functionalization of n-type InP porous nanostructures and their feasibility for biochemical sensor applications were investigated. The porous structures have extremely large surface areas, i.e., over 10 m²/cm³, and superior electrical properties with conductive semiconductor substrates. As a first attempt at electrochemical functionalization, we successfully deposited a glucose oxidase (GOD) membrane onto an InP surface under an applied anodic bias of 1.2 V. With the addition of glucose, the response currents on the porous electrodes increased compared to those on planar InP electrodes due to their enlarged surface area. The sensitivity curves of the porous electrodes we used showed good linearity between the response currents and concentrations in a range from 0 to 5 mM.

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In recent years, the demand for biological and chemical sensors has increased in biochemical field areas such as blood sugar monitoring, antibody and DNA testing, and virus screening. One present challenge in the biochemical sensor research field is to utilize conductive nanomaterials with a large surface area. Various nanomaterials have been investigated for this purpose, such as nanoparticles, carbon nanotubes, semiconductor nanowires, conducting polymers, and porous materials.

Of these, the semiconductor porous nanostructures formed by electrochemical anodization are one of the more promising candidates for use as the building blocks of biochemical sensors. The direct formation of porous structures has been investigated for various materials such as Si, GaAs, GaP, InP, GaN, and CdSe. Hamamatsu et al. succeeded in electrochemically forming arrays of straight nanopores without branches on n-InP(001) substrates. The self-assembled formation of hexagonal arrays of pores was later reported by Fujikura et al. and Langa et al., where the pores were laterally separated by InP nanowalls and were formed along a straight vertical direction over a distance of approximately 1–100 µm. Such close-packed nanostructures have extremely large surface areas (i.e., over 10 m²/cm³) compared with the semiconductor surfaces prepared by conventional methods used in the semiconductor industry such as lithography and dry etching, vacuum evaporation, and epitaxial growth. Merchant et al. recently reported that the InP nanowalls between each pore show proper conductivity and carrier mobility. These features reported for InP porous nanostructures are very attractive for biochemical sensor applications.

The purpose of this study is to investigate the feasibility of amperometric biochemical sensors based on InP porous nanostructures functionalized by enzymatic molecules. In terms of electron mobility, InP is a more suitable semiconductor material for both potentiometric and amperometric sensors than Si, which has been extensively studied using metal-oxide-semiconductor structures. Various gas sensors formed on n-type InP have recently been reported, such as H₂ gas sensors and NO₂ gas sensors. They have shown good performance with high sensitivity due to the large change in the conductance response of the target gases. Almost all of the sensors based on InP have utilized bare surfaces or Schottky interfaces to detect the sensing target. To create greater variation in sensing targets, the functionalization process of utilizing organic molecules is important. However, very few efforts have been made to functionalize InP surfaces. In this study, we first examined the electrodeposition of a glucose oxidase (GOD) membrane on InP planar substrates. We then applied the functionalization process to InP porous nanostructures and investigated their electrocatalytic activity for application to amperometric glucose sensors.

Experimental

Reagents and materials.— GOD (enzyme class 1.1.3.4, from Aspergillus niger, 200 units/mg) and β-D-(+) glucose were obtained from Sigma-Aldrich Inc. and were used as received. We obtained 0.1 M phosphate buffer solutions (PBSSs) with pH 7.0 and 7.4 from Načala Tesque Inc. and used them as solvents for the electrodeposition of GOD and for electrochemical measurements, respectively. We obtained 60% HNO₃ and 35% HCl solutions from Kishida Chemical Inc. and used them as electrolytes for the formation of porous structures, as described in the literature. All other solvents and reagents used were of analytical grade, and deionized (DI) water was used to prepare the solutions.

Sn-doped InP(001) substrates (n-type, n = 1 × 10¹⁸ cm⁻³) were obtained from Wafar Technology Ltd. The substrates were cleaved into square pieces and used as working electrodes (WEs). The geometric surface area of a WE was defined by the photos resist at 0.13 cm².

Fabrication of GOD-functionalized InP porous electrodes.— The procedures for preparing GOD-functionalized InP porous electrodes are shown in Fig. 1. Electrodes were first anodized at 4 V in an electrolyte comprising 1 M HCl (200 mL) with HNO₃ (3 mL). Here, pore diameter and wall thickness can be controlled by adjusting the electrochemical anodic and cathodic conditions. After that, the porous surface was photoelectrochemically etched in the same electrolyte under illumination to remove the disordered irregular layer formed at the initial stage of pore formation. Figure 2a and b shows typical scanning electron microscope (SEM) images of an InP porous sample after photoelectrochemical (PEC) etching for 200 s. Regular-sized pores of about 130 nm in diameter appeared on the surface over a wide area. The pore depth d was 3.2 µm from the SEM observations of the cross section. Finally, we conducted an electrodeposition of GOD in a 10 mg/mL GOD-containing electrolyte based on a PBS with pH 7.0. After electrodeposition, each sample was washed with DI water and stored under dry conditions in a refrigerator.

Characterization and amperometric measurements.— A series of electrochemical functionalization and amperometric measurements was conducted using a standard cell with three electrodes: an InP porous electrode used as a WE, a Pt counter electrode, and a saturated calomel electrode (SCE) used as a reference. The WE potential with respect to SCE was precisely controlled with a potentiostat (Princeton Applied Research 273 A).

First, we attempted to optimize the conditions for the electrodeposition of GOD onto the InP surfaces. For this purpose, an InP(001) planar electrode was used as a WE that was dipped into a...
GOD-containing electrolyte. The applied bias for the electrodeposition of GOD was varied over a range from 0 to 1.2 V, and the deposition time was set at 60 s. The GOD membrane formed on the InP surface was analyzed by X-ray photoelectron spectroscopy (XPS) using a Perkin-Elmer PHI 1600 with a monochromatic Al Kα source at 1486 eV.

We then investigated the sensing properties of the GOD-functionalized InP porous electrodes used as an amperometric glucose sensor. The electrodes were dipped into a PBS with pH 7.4, and a sensing bias \( V_s \) was set at +0.65 V under darkroom conditions. Mild magnetic stirring was maintained throughout the measurements. After the background current stabilized, amperometric measurements were carried out by successively adding a glucose solution into the PBS and recording the current signals.

**Results and Discussion**

Electrodeposition of the GOD membrane onto an InP surface.— Figure 3a and b shows the C 1s XPS spectra obtained from the reference InP bare surface and the GOD-deposited surface at \( V_d \) of 1.2 V. As shown in Fig. 3a, a sharp peak was observed at 285 eV in the hydrocarbon (C–H) region, and a small and broad oxide peak was observed at a higher energy. A more complicated spectrum was observed from the GOD-deposited InP surface, as shown in Fig. 3b. Simple apportionment of the C 1s spectrum was enabled by the amic characteristics of GOD with two specific peaks at 288 and 286 eV, corresponding to those of carbonyl and N–C–CO groups, respectively. These groups were clearly absent in the C 1s spectrum obtained from the InP bare surface. These results confirmed that a GOD membrane was formed on the InP surface by the electrodeposition process.

To optimize the electrochemical conditions, XPS analysis was performed on the deposited GOD membranes by changing the anodic bias applied to the InP substrates. Figure 4 compares the ratios...
of the XPS peak areas of various elements (C 1s, N 1s, and O 1s) to P 2p$_{3/2}$ that originated in the InP substrates. For the sample prepared with an anodic bias of 0.4 V, the XPS ratios showed no significant changes from those obtained with the reference sample. However, as the anodic bias was increased to 0.8 V, the ratios of GOD-related elements such as C, N, and O increased. As shown in Fig. 4, the sample prepared with an anodic bias of 1.2 V had a more pronounced tendency to show increased ratios of GOD-related elements. These results suggest that a GOD membrane was efficiently deposited on the InP surface at a higher anodic bias. The bias-dependent XPS spectra can be explained as follows. In this study, the electrolyte pH was adjusted to 7.0, which is higher than the isoelectric point for GOD (pI = 4.2). Because the GOD molecules were negatively charged in the electrolyte, the electric force that draws the GOD toward the InP surface was increased under the anodic bias applied.

InP surfaces are readily oxidized in an aqueous electrolyte under an anodic bias applied even under darkroom conditions. A higher anodic bias increases the risk of inactivating the GOD molecules due to the protein denaturation caused by the high electric field at the electrolyte/InP surface. For the same reason, Matsumoto et al. reported that the anodic bias of 1.3 V (vs AgCl/Ag electrode) is suitable for the GOD deposition onto the Pt surface. Thus, we decided to use the anodic bias of 1.2 V for our next experiments on the electrodeposition of GOD onto an InP porous surface.

Glucose detection by InP porous electrode — The electrocatalytic activity of InP porous electrodes was investigated after the electrodeposition of the GOD membrane at $V_d$ of 1.2 V. Figure 5a compares the current response of the porous electrodes with a pore depth $d$ of 3.2 or 1.9 μm and that of a reference planar electrode ($d = 0$ μm) to which glucose solution was added to the PBS with pH 7.0. The surface area used to calculate the current density comprises only the geometric surface area and does not include the wall inside. For planar electrodes with a GOD membrane, almost no response current was observed. However, anodic response currents were clearly observed on InP porous electrodes, as shown in Fig. 5a. As the amount of glucose solution increased as indicated by the arrows, the anodic current increased. The current level obtained from the sample with a $d$ of 3.2 μm was larger than that obtained from the sample with a $d$ of 1.9 μm. This suggests that the electrocatalytic activity of the InP porous electrodes was enhanced due to the increased surface area with pore depth $d$.

Furthermore, the response current reached peak values at the very beginning of the glucose addition process and decreased thereafter. Figure 5b shows the transient curves of anodic currents obtained on a porous electrode with a $d$ of 3.2 μm by adding different concentrations of glucose at a time. As expected, the measured current density was strongly dependent on the glucose concentration. After the currents reached peak values at the very beginning, they then decreased rapidly and approached various saturation values, as shown in Fig. 5b. This kind of transient current behavior has not been previously observed in planar electrodes on a time scale of seconds to minutes. The current response we observed is most probably due to the diffusion-limited process of reactants inside nanopores with a small diameter.

Glucose in electrolytes can be detected by a GOD-functionalized electrode with the following reactions

$$\text{glucose} + \text{O}_2 \rightarrow \text{glucono-} \text{lactone} + \text{H}_2\text{O}_2 \quad [1]$$

$$\text{H}_2\text{O}_2 \rightarrow \text{O}_2 + 2\text{H}^+ + 2\text{e}^- \quad [2]$$

GOD catalyzes the conversion of a glucose molecule, as described in Eq. 1, where one glucose molecule reacts with one water-dissolved oxygen molecule generating one molecule of hydrogen peroxide ($\text{H}_2\text{O}_2$). When an anodic bias was applied to a sensor electrode large enough to induce the reaction described in Eq. 2, anodic currents were observed as response currents. We have recently reported that $\text{H}_2\text{O}_2$ can be detected by an InP porous elec-

![Figure 4](image-url) 

**Figure 4.** (Color online) XPS peak area ratio of C 1s, N 1s, and O 1s to P 2p$_{3/2}$ on GOD-deposited surfaces as a function of the anodic bias applied.

![Figure 5](image-url) 

**Figure 5.** (Color online) (a) Current response of the porous and reference planar electrodes to the addition of glucose. Arrows indicate the timing of the addition of glucose into the PBS electrolyte. (b) Transient curves of anodic currents obtained on the porous electrode with different concentrations of glucose only added once.
difference is that the response speed was limited by the glucose reaction described in Eq. 2, whose reaction rate was much lower than that described in Eq. 1.

The sensitivity for detecting glucose obtained in this study was higher than 0.023 $\mu$A/mM cm$^2$ that was previously obtained for carbon-paste electrodes. Carbon-related materials have been investigated earlier as a sensor material due to their high potential abilities in terms of both surface area and electron mobility. In consideration of this point, our data obtained for the InP porous nanostructures was not bad for an initial attempt. However, our sensor structure has not been optimized as compared with the recent progress on carbon nanotubes. Xie et al. reported that a higher sensitivity of 52.7 $\mu$A/mM cm$^2$ was obtained for Pt-coated multi-walled carbon nanotube electrodes. Improved sensitivity of the InP porous electrodes can be possibly obtained by optimizing such structural properties as pore depth and pore diameter and by using Pt nanoparticles to enhance functionalization inside the pores.

The data shown in Fig. 6 were obtained by using the same electrode repeatedly while changing its electrolyte. The results indicate that the present InP porous electrodes are stable enough to obtain reproducibly increased currents in response to the glucose concentration. However, the performance of the electrodes degraded for a few weeks during which the response currents decreased by about 30% compared to the data obtained at the first measurement. This is probably due to the deactivation of the GOD membrane, where we had not considered surface protection as a first step for the functionalization of the InP porous nanostructures with a GOD membrane.

We suppose that InP itself is chemically stable in terms of biochemical applications. We have recently reported that InP-based ion-sensitive field effect transistors showed a stable operation in a pH range from 3.0 to 12.0. This covers the optimum pH range for various enzymatic reactions, such as lipase (pH 4.0–5.0), amylase (pH 6.7–7.0), and trypsin (pH 7.8–8.7). From our present results described here, we consider that InP porous electrodes functionalized by electrodeposition have considerable potential for practical application to various amperometric biochemical sensors.

**Conclusion**

We investigated the functionalization process of n-type InP porous electrodes and their feasibility for biochemical sensor applications. As a first attempt at achieving electrochemical functionalization, a GOD membrane was successfully deposited onto an InP surface under an applied anodic bias. The response currents of the glucose sensors increased on the porous InP electrodes due to their enlarged surface area. The functionalized porous electrodes exhibited good sensitivity with respect to detecting glucose concentrations in the electrolyte, where the response currents detected were attributed to H$_2$O$_2$ anodization. We consider that the present porous electrodes functionalized by electrodeposition have considerable potential for practical application to various amperometric biochemical sensors.

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