Structural Study of Porous Si with Different Crystal Orientation Fabricated Using Two-Step Alternating Current Photo-Electrochemical Etching Technique

Fatimah Zulkifli, Rosfariza Radzali, Alhan Farhanah Abd Rahim, Aïnorkhilah Mahmood and Aslina Abu Bakar

Abstract— Two different crystal orientation of n-type Si substrates n(100) and n(111) were used to fabricate porous structure using two-step Alternating Current Photo-Electrochemical (ACPEC) etching technique. This research aims to investigate the difference of porous structure and properties for different orientation of n-type Si substrate. In this work, before alternating current (AC) was supplied, the samples were immersed in HF:Ethanol for 10 minutes. Then, each sample were etched at 20 mA/cm² current density for 30 minutes. The porous Si samples then were characterized for Field Emission Scanning Electron Microscopy (FESEM), Atomic Force Microscopy (AFM) and High Resolution X-ray Diffraction (HR-XRD) to study the surface morphology of the samples. The results obtained showed that porous Si with orientation of (100) has higher pore density formation, higher surface roughness value in RMS and deeper pore depth with improved crystalline quality compared to Si with orientation of (111).

Index Terms—Alternating Current, Crystal Orientation, Electrochemical Etching, Porous Si.

I. INTRODUCTION

Silicon (Si) is the second abundant material after carbon and is widely used in semiconductor technology. The discovery of porous Si efficient visible luminescence at room temperature has garnered many attentions from researchers [1, 2]. Numerous electronic devices, ranging from optoelectronic to chemical and biochemical sensors applications, have been developed using porous Si due to its unique features such as high surface area, band gap shift, and efficient luminescence properties [3-7]. Solar cells [8, 9] and hydrogen gas sensing [6] are some of the few examples of application built using porous Si. Photo-electrochemical (PEC) etching technique is an etching technique which uses a current supply, either direct current (DC) or alternating current (AC), along with the help of light source. This technique has its own advantages such as low processing temperature, low structural damage, simple and versatile, and low cost compared to other technique available [10]. By varying the etching parameters such as etching duration and current density [5, 11-13], the porosity formation is more or less can be easily controlled.

In this work, two different crystal orientation of n-type Si which are n(100) and n(111) were used to fabricate porous structure using two-step Alternating Current Photo-Electrochemical (ACPEC) etching technique. The effect of different orientation of Si substrate towards the formation and properties of porous structure were investigate.

II. EXPERIMENTAL PROCEDURE

Two types of N-type silicon substrates were used to fabricate porous sample in this research, namely n(100) and n(111). The techniques used to fabricate these samples is called two-steps ACPEC etching technique. Silicon wafer of both type were cut into smaller pieces approximately 1cm x 1cm to fit into Teflon cell. Then, Radio Corporation of America (RCA) cleaning method was used to clean the sample from any impurities before the etching process. It was important to ensure the sample was clean as the impurities could affect the etching process and ruin the results obtained from the sample.

Fig. 1 shows the experimental setup. A Teflon cell with O-ring and metal plate was used to hold the sample for etching process. The metal plate acted as anode during the process while platinum wire which was dipped into the electrolyte solution acted as cathode. The Si sample was fixed in between O-ring and metal plate with the front side was exposed to the electrolyte solution. The electrolyte solution used in this research was a mixture of hydrofluoric acid and ethanol with ratio of 1:4. This solution has been commonly used to fabricate porous silicon by other research groups [5, 14, 15]. Ethanol was used in the electrolyte solution to act as a surface-active agent which help to reduce surface tension during the etching process to achieve high density pore formation [16]. The ammeter was used to monitor the current supply.
Two-steps ACPEC etching technique consists of two etching steps. The first step was to immerse the Si sample in electrolyte solution for a period of time to form a high density of etch pits [17]. The second step was to anodize the sample in electrolyte solution by supplying alternating current (AC) with current density of 20 mA/cm² for 30 minutes under the illumination of incandescent light [14]. Anodization using AC help to reduce the hydrogen bubbles and allow the HF to react with the Si surface [17]. Once etching process was done, the samples were rinsed with DI water and dry under ambient air.

All of the etched samples were characterized for their structural properties using Field Emission Scanning Electron Microscopy (FESEM) (Model: Jeol JSM 7401F), Atomic Force Microscopy (AFM) (Model: Dimension EDGE, BRUKER), and High Resolution X-ray Diffraction (HR-XRD) (Model: PANalytical X’pert Pro MRD).

III. RESULTS AND DISCUSSIONS

A. Fields Emission Scanning Electron Microscopy Analysis

Fig. 2 shows the FESEM images of as grown and porous Si sample with different crystal orientation. ImageJ software was utilized to examine the average pore size and porosity for each samples.

Fig. 2(a) and (c) show the surface morphology of as grown Si n(100) sample and Si n(111). Both images showed that the surface were very smooth with no pores or crack present. From Fig. 2(b) and (d), it can be seen that the formation of porous structure on Si n(100) sample was more uniform compared to porous structure on Si n(111) as there were less formation of pores can be seen for porous Si n(111) sample. On top of that, porous Si n(100) was observed to has higher density of porous structure compared to porous Si n(111) sample. Similar result obtained by Abd Wahab et al. where Si n(100) formed more uniform circular pore structure and dense compared to Si n(111) [18]. The shape of pores that formed on porous Si n(100) were mostly crisscross in shape with some of them contain square shape pores in the middle of the crisscross pores. Meanwhile, the shape of pores that formed on porous Si n(111) sample were mostly irregular with circle and square like pores. The pore length (L), width (W) and diameter (D) were measured as indicated in the inset image of Fig. 2. The pore length is the length of the pore and the pore width is width of pore from the crisscross shaped pore. While, the pore diameter is the diameter of either the circular or square shaped pore. The average length of pores for porous Si n(100) sample was ~3.18 µm with average width of ~0.45 µm and average diameter of the square shaped pores was ~0.62 µm. As for porous Si n(111) sample, the average diameter of the irregular shaped pores was ~0.73 µm.

From Fig. 2, the estimation of porosity percentage for each sample was determined using ImageJ by a proportion of sample surface involved with pores to the entire surface of the sample [19]. It can be seen that sample porous Si n(100) sample has higher porosity percentage compared to sample porous Si n(111) sample which are 24.43% and 16.88%, respectively. The lower porosity percentage of sample Si n(111) compared to Si n(100) was due to lower etching rate of Si n(111). The lower etching rate for Si n(111) could be related to its atomic structure which was densely packed compared to Si n(100) [20, 21]. On top of that, the lower etching rate of Si n(111) is also probably due to the energy required to remove atoms from the surface was relatively much higher in Si n(111) compared to Si n(100).

This is because each atom of Si n(111) surface has one dangling bond and three back bonds while each atom of Si n(100) surface has two dangling bond and two back bonds only. Therefore, the
activation energy to remove an atom from the surface of Si n(111) is higher which it needs to break three back bonds compared to only two back bonds in the case of Si n(100) surface [20, 21]. Based on these results, it can be concluded that pore formation was better on sample Si n(100) compared to sample Si n(111). Therefore, FESEM results indicated that Si n(100) was more suitable to fabricate porous structure with high density and uniformity pores. Table I shows the summary of pore size for each porous sample with their respective porosity.

B. Atomic Force Microscopy Analysis

Fig. 3 shows the AFM measurement of as grown and porous Si sample with different orientation. The average surface roughness in root mean square (RMS) was determined using the NanoScope Analysis Software and estimated pore depth of each sample were determined using the line scan. The data obtained was calculated and tabulated in Table II.

The low surface roughness of as grown Si n(100) and n(111) with RMS value of 0.30 nm and 0.19 nm, respectively showed the absence of pores on the surface. On the other hand, porous Si n(100) and n(111) samples have higher RMS value compared to as grown sample, thus showing that there were pores present on the surface as the structure of the porous samples have been altered during the etching process [22].

The AFM results also shows that porous Si n(100) sample has higher average pore depth of ~260.83 nm compared to average pore depth of porous Si n(111) which was ~86.78 nm. This result is supported by FESEM images in Fig. 2(b) where porous Si n(100) showed deeper pores (visible as darker area) compared to porous Si n(111) in Fig. 2(d). Porous Si n(100) also has higher RMS value which was 215.0 nm compared to porous Si n(111) which has RMS value of 161.0 nm. The lower surface roughness of Si n(111) was associated to its lower etch rate compared to Si n(100) [20, 21] as explained previously. Similar results were observed from previous research where Si n(100) sample has higher RMS value compared to Si n(111) sample [18]. Therefore, porous Si n(100) sample has more pores formed compared to porous Si n(111) that has widely spread pores formation.

NanoScope Analysis Software

Table II

<table>
<thead>
<tr>
<th>Sample</th>
<th>Average Pore Depth (nm)</th>
<th>Surface Roughness in RMS (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>As grown Si n(100)</td>
<td>0.262</td>
<td>0.296</td>
</tr>
<tr>
<td>Porous Si n(100)</td>
<td>260.832</td>
<td>215.0</td>
</tr>
<tr>
<td>As grown Si n(111)</td>
<td>0.213</td>
<td>0.194</td>
</tr>
<tr>
<td>Porous Si n(111)</td>
<td>86.775</td>
<td>161.0</td>
</tr>
</tbody>
</table>

Fig. 3. AFM measurement of as grown and porous Si samples with different orientation (a) as grown Si n(100), (b) porous Si n(100), (c) as grown Si n(111) and (d) porous Si n(111).

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Table III

<table>
<thead>
<tr>
<th>Sample</th>
<th>2θ (°)</th>
<th>FWHM (°)</th>
<th>Crystallite Size (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>As grown Si n(100)</td>
<td>69.125</td>
<td>0.11</td>
<td>91.58</td>
</tr>
<tr>
<td>Porous Si n(100)</td>
<td>69.175</td>
<td>0.10</td>
<td>100.76</td>
</tr>
<tr>
<td>As grown Si n(111)</td>
<td>28.475</td>
<td>0.14</td>
<td>61.12</td>
</tr>
<tr>
<td>Porous Si n(111)</td>
<td>28.425</td>
<td>0.16</td>
<td>53.48</td>
</tr>
</tbody>
</table>

Fig. 4. The HR-XRD of as grown and porous Si samples with different orientation.

TABLE II

SUMMARY OF AFM MEASUREMENT OF AS GROWN SI N(100), POROUS SI N(100), AS GROWN SI N(111) AND POROUS SI N(111)

TABLE III

SUMMARY OF DATA OBTAINED FROM HR-XRD MEASUREMENT
C. High Resolution X-Ray Diffraction Analysis

Fig. 4 shows the HR-XRD measurement of as grown and porous Si samples with different orientation. Results obtained from HR-XRD measurement can be used to evaluate the crystalline properties of the samples.

From Figure 4, the peak position for as grown Si n(100) and porous Si n(100) occurred at about 69.1° represented by (400) reflection of cubic Si phase. There was no peak shift occurred between porous Si n(100) and as grown Si n(100) which indicate no stress was experienced during the etching process [15]. In addition, the peak intensity for porous Si n(100) is lower compared to as grown Si n(100). This is probably due to the formation of pores which influence the etching process [22]. Similar result could be observed from previous research work where the peak intensity for porous Si n(100) was lower than the as grown Si n(100) [23]. For as grown Si n(111) and porous Si n(111), the peaks appeared at about 28.4° which indicate the presence of cubic Si phase by (111) reflections. Similar trend has been observed for Si n(111) samples in which there were no peak shift and lower peak intensity of porous Si n(111) compared to as grown Si n(111). This result is similar to some previous research where the peak intensity for porous Si n(111) was lower than the as grown Si n(111) [14, 22]. Therefore, the formation of pores do not change the peak position of the samples. Table III tabulates the peak position, FWHM and the crystallite sizes of all the samples.

The average crystallite size of the samples can be obtained by using the Debye-Scherrer equation (1) as shown below:

\[ D = \frac{0.9 \lambda}{\beta \cos \theta} \]  

where D is the average crystallite size, \( \lambda \) is the X-Ray wavelength (0.15406 nm), \( \beta \) is the FWHM in radians, and the \( \theta \) is the diffraction angle in radians. From Table 3, the Full-Width-Half-Maximum (FWHM) value for porous Si n(100) is narrower compared to as grown Si n(100) with average crystallite size of \( \sim 100.76 \) nm. The lower value of FWHM indicated better crystalline quality due to the uniform pores formation of porous Si n(100) compared to as grown Si n(100) [17]. Similar result could be observed in research by [23] where porous Si n(100) has lower FWHM value and larger crystallite size. Meanwhile, porous Si n(111) has higher FWHM value which resulted with lower crystallite size value of \( \sim 53.48 \) nm compared to as grown Si n(111). A possible explanation for this was the formation of a few irregular pores with small diameter that were widely spread on the surface of porous Si n(111). The increase in FWHM value is probably due to the growth of native oxide on the surface of porous Si n(111) [17]. This result is in line with the result obtained by [14] that showed smaller crystallite size of porous Si n(111) compared to as grown Si n(111). From this result, different crystal orientation of Si resulted in different surface morphology.

IV. CONCLUSIONS

Porous Si was successfully fabricated by using two-step ACPEC etching technique for different crystal orientation of Si substrate, namely n(100) and n(111). Based on results shown by FESEM, porous Si n(100) produced a higher density of pores and porosity percentage compared to porous Si n(111). Results from AFM also supported the data from FESEM as porous Si n(100) has higher surface roughness in RMS value and higher average pore depth compared to porous Si n(111). HR-XRD results showed that there was improvement in crystalline quality for porous Si n(100). In conclusion, based on all the data obtained, porous Si n(100) was more suitable to be used to fabricate porous sample as it formed higher density of pores with higher average pore depth than porous Si n(111). The crystalline quality for porous Si n(100) sample was also improved after the etching process.

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