Studies of the self-assembled growth mechanism on nanocrystalline silicon nanodots

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Received 23 February 2010, Revised 7 July 2010, Accepted 15 July 2010, Available online 8 October 2010

ABSTRACT

Nanocrystalline silicon (nc-Si) nanodots have been grown on corning glass (7059) substrate using a self-assembly VHF-PECVD method under the following experimental conditions: Fixed deposition temperatures of 300/400 °C, deposition times 30/60 s, plasma power of 10 W, silane gas flow rate of 10 sccm, as well as deposition pressure of 10^-2 torr. It is predicted that the onset of nucleation began immediately after the deposition and start to appear clearly after 20-60 s during which growth mechanisms occur. Essentially, the nanodots were formed onto the substrate in dome-like shapes by virtue of equilibrium surface energies, γ_Ls, γ_LS and γ_NS. The associated liquid/solid nucleation mechanism was then simulated and related parameters were obtained:

- Surface energies per unit area, \( \gamma_{LS} = 1.44 \text{ Jm}^{-2} \), \( \gamma_{LN} = 19 - 60 \text{ Jm}^{-2} \), \( \gamma_{NS} = 0.74 \text{ Jm}^{-2} \);
- Free energy change per unit volume \( \Delta G^* \sim 10^{-15} \text{ J} \);
- Critical radii \( r^* = 16 - 48 \text{ nm} \).

Other measurements were also carried out: PL analysis gave bandgap energies ~ 1.8-2.4 eV, whilst Raman spectra revealed the coexistence of nc-Si and amorphous Si. It is strongly suggested that, the nc-Si nanodot grown on glass substrate fulfills the Volmer-Weber growth mode with a minor modification.

1. INTRODUCTION

Nanodot is an artificial nanostructure having diameters between 10 - 100 nm beyond which quantum dot (<10 nm) and thin film (>100 nm) are dominant. It can fill up to more than 10^7 atoms, i.e. about ten times the number of atoms in a quantum dot. Recent studies on nanocrystalline silicon (nc-Si) revealed its potential applications in wide areas of interest, and it is the purpose of this paper to investigate the growth of nanodots both from the stand points of theoretical prediction and experimental evidence.

Nanocrystalline silicon is an allotropic form of silicon with paracrystalline structure, i.e., having short and medium range ordering in their lattice but lacking long range order. It has grains sizes less than 100 nm and normally exists in amorphous phase [1,2]. For nc-Si thin films fabricated using CVD methods varying grain structures are observed to be embedded in the amorphous matrix and its sizes are much dependent on the experimental conditions. The crystalline-amorphous changes are still the subject of interest, but generally can be envisaged as bonding rearrangement occurring in amorphous stage. nc-Si has many useful advantages over a-Si, one being that if grown properly it can have a higher electron mobility, due to the presence of the silicon crystallites. It also shows increased absorption in the red and infrared wavelengths, which make it an important material for use in a-Si solar cells [3]. One of the most important advantages of nanocrystalline silicon, however, is that it has increased stability over a-Si, one of the reasons being because of its lower hydrogen concentration. Although it currently cannot attain the mobility that poly-Si can, it has the advantage over poly-Si that it is easier to fabricate, as it can be deposited using conventional low temperature a-Si deposition techniques, such as PECVD, as opposed to laser annealing or high temperature CVD processes, in the case of poly-Si.

The main application of this novel material is in the field of silicon thin film solar cells. As nc-Si has about the same bandgap as crystalline silicon, which is ~1.12 eV, it can be combined in thin layers with a-Si, creating a layered, multi-junction cell called a tandem cell. The top cell in a-Si absorbs the visible light and leaves the infrared part of the spectrum for the bottom cell in nanocrystalline Si [3,4].

1.1. Nucleation and growth theory

An understanding of the growth mechanism can be dated back to the classical nucleation theory [5] and the three growth modes of nanofilm [6]. The diagram in Fig 1. distinguishes different growth mechanisms, including two-dimensional (a) layer-by-layer growth or Frank-van der
Merwe (F-M), (b) island growth or Volmer-Weber (V-W), and (c) mixed growth of Stranski-Krastanov (S-K). These growth modes generate stable islands, whereas three different kinds of ripening which may or may not include a two-dimensional layer (frequently called the wetting layer) are possible. The diagrams demonstrate that in order to achieve stable and controllable S-K growth, for which uniform small islands or nanodots may result, it is necessary to employ semiconducting crystal systems that include a large amount of strain resulting from structural misfits, ε (i.e., work far out along the horizontal axis in this diagram), and the deposited material must not exceed a certain critical coverage.

For the case of V-W growth, the contribution of misfits at the material-substrate interface is no longer significant, except for certain semiconductors with suitable confirmation of phase diagram. There is a possibility that, V-W growth can occur in amorphous and non-crystalline semiconductors if the surface parameters are taken into consideration. These normally refer to the classical theory of heterogeneous nucleation, as it only depends on $\Delta T$ (undercooling temperature), $r$ (radius in heterogeneous nucleation is the same as in substrate surface is called the contact angle. The critical radius in heterogeneous nucleation is the same as in homogeneous nucleation, as it only depends on $\Delta G_v$ (change in energy of solidification). The substrate surface is usually considered homogeneous, although there is a possibility of some non-uniform surfaces. In this investigation planar substrate surface geometry is treated.

From the thermodynamic point of view, a change in the free enthalpy, $\Delta G$ is associated with nucleation via the following four processes: Firstly, creation of new volume of the nucleus (negative); secondly existence of surface area of liquid-nucleus (positive), thirdly existence of surface area of liquid-nucleus (positive) and lastly the energy lost at liquid-substrate interface (positive). Thus [8],

$$\Delta G = \beta_3 r^3 L_f \frac{\Delta T}{T_m} + \beta_1 r^2 \gamma_{LN} + \beta_2 r^2 \gamma_{NS} - \beta_3 r^2 \gamma_{LS}$$

(2)

where, $\beta_3 = 2 \pi (1 - \cos \theta)$, $\beta_2 = \pi \sin^2 \theta$, $\beta_1 = \pi (2 - 3 \cos \theta + \cos^3 \theta) / 3$, $\Delta G_v = L_f (\Delta T/T_f)$ is the energy change per unit area of the nucleus and has the values depending on the undercooling temperature, $\Delta T (T_m - T)$; positive if $T > T_m$, with $T_m$ being the temperature of the transition (solidification temperature), $L_f$ is enthalpy of fusion. Under these conditions, applying the Gibbs-Duhem principle, the new phase is not stable. Since $\gamma$ is a positive quantity, $\Delta G_v$ is then positive: the number of nuclei of the new phase are not stable and do not tend to grow. On the contrary, if $T < T_m$, negative, the new phase (solid) is stable. There are values of $r$ for which $\Delta G_v < 0$, and the corresponding nuclei will tend to stabilize the new phase since their formation reduces the free enthalpy of material. When $T < T_m$ ($\Delta T > 0$), $\Delta G_v$ has a
maximum for \( r = r^* \). For \( r > r^* \), the formation of nuclei of increasing size can result in stabilization of the solid phase since \( \Delta G \) decreases and even becomes negative. Both the critical energy, \( \Delta G^* \) and critical radius, \( r^* \) are obtained as,

\[
\Delta G^* = \frac{16\pi r^3_{NS}T_m^2}{3L_f\Delta T^2} \left(2 - 3 \cos \theta + \cos^3 \theta\right)
\]

(3)

\[
r^* = \frac{-2\gamma_{NS}}{\Delta G^*} = \frac{-2\gamma_{NS}T_m^2}{L_f^2\Delta T^2}
\]

(4)

\( \Delta G^* \) is equivalent to the potential barrier that must be lowered for nuclei of critical size \( r^* \) to be formed more easily. It seems that from equation (2) the more important supercooling is, the more this barrier is lowered. The Volmer thermodynamic model shows that a complete phase transition by nucleation is only possible if the liquid is supercooled. In fact, nucleation is initiated with lower supercooling (larger \( \Delta T \)) because the impurities present in the liquid phase, even in low concentration, induce nucleation, normally referred to heterogeneous nucleation.

2. EXPERIMENTAL

The samples were fabricated using a home-made very high frequency (150 MHz) plasma enhanced chemical vapour deposition system (VHF-PECVD) at a chamber pressure \( 10^{-2} \) Torr, substrate temperatures 300 °C and 400 °C, fixed argon flow rate of 10 sccm, and deposition times varied between 20-60 seconds. The principal process was based on flowing a known rate of silane gas (silane10%/argon90%) in a specially designed chamber, and allowing 150 MHz RF supply to effectively enhance the chemical processes whereby the silicon particles produced is buried on the substrate (corning glass) and other by-products are released to the atmosphere. Selected samples fabricated at 300°C/25W/15 mins were analysed using photoluminescence (PL) and XRD in order to determine the bandgap energy, \( E_g \) and contents, respectively. From PL spectrum shown in Fig. 5, \( E_g \) were found to lie between \( ~1.8 - 2.4 \text{ eV} \), the values which are greater than that of bulk silicon (\( > 1.1 \text{ eV} \)) thus confirming the existence of nanodots in the samples. This is supported by the Raman analysis for 300/400 °C samples (Fig. 3) which reveals the coexistence of both nc-Si crystal at 517 cm\(^{-1}\) and amorphous Si at 480 cm\(^{-1}\) [9]. All the above results are different from those of Si quantum dots [8] fabricated using the sputtering method.

![Fig. 2. Typical AFM image of nc-Si nanodots formed at 573K/30s/10W/10 sccm](image)

![Fig. 3. Raman spectra of of nc-Si nanodots formed at different temperatures](image)

3. RESULTS AND DISCUSSION

The analysis obtained from AFM (Figure 2) shows that the nanodots fabricated at 300 °C(573 K) may start to form earlier, but they seem to be more defined and observable at the time of 30 s after the deposition. Under such a condition, less number of dots is formed on the substrate, approximately around 28 dots per \( \mu m^2 \). Further analysis on the cross sectional area reveals the dot’s patterns in dome-like shape, which measure around 45.11 nm in diameter. This appearance is contrary to a pole-like shape observed in Fig.2. For the nanodots fabricated at a temperature of 400 °C (673 K) and similar deposition time (Fig. 4a), marked changes occurs in both density and size of 134 dots per \( \mu m^2 \) and ~88.11 nm (Fig. 4b), respectively. The dependence of contact angle, \( \theta \) at 70° was found to be less significant as can be observed for the case of \( r^* \) in Fig. 7. Some selected samples were analyzed using photoluminescence (PL) and XRD in order to get better resolution of the spectrums. A theoretical investigation was done using the classical approach of heterogeneous nucleation in which the net energy change required to form a spherical nucleus is equivalent to the summation of energy terms associated with four different interfacial interactions.
substrate temperature, each for $T = 573$ K and $673$ K; undercooling temperature, each for $T_m - T = 1110$ K dan $1010$ K respectively, atomic volume, $v_o = 12.1 \text{ cm}^3/\text{mol}$ and contact angle, $\theta = 70^\circ$. The simulation results have been obtained and are shown in Table 1 (for $T = 573$ K).

Fig. 4. Typical AFM images of nc-Si nanodots a) formed at $673$ K/30 s/10 W/10 sccm - circled, and b) measured density $\sim 134$ dots per $\mu m^2$ and diameter $\sim 88.11$ nm.

Table 1. Experimental and computed growth parameters

<table>
<thead>
<tr>
<th>Experiment, $T = 573$ K</th>
<th>Simulation</th>
<th>Fixed, $\gamma_{NS} = 1.44 \times 10^{-4} \text{ J cm}^{-2}$</th>
<th>$\gamma_{LS} = 7.37 \times 10^{-3} \text{ J cm}^{-2}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$d$ (nm)</td>
<td>$r^*$ (nm)</td>
<td>$\gamma_{NS}$ ($\text{J cm}^{-2}$)</td>
<td>$r^*$ (cm)</td>
</tr>
<tr>
<td>32.17</td>
<td>16.09</td>
<td>$19.8 \times 10^{-4}$</td>
<td>$1.61 \times 10^{-6}$</td>
</tr>
<tr>
<td>45.11</td>
<td>22.56</td>
<td>$27.8 \times 10^{-4}$</td>
<td>$2.26 \times 10^{-6}$</td>
</tr>
<tr>
<td>54.56</td>
<td>27.28</td>
<td>$33.7 \times 10^{-4}$</td>
<td>$2.73 \times 10^{-6}$</td>
</tr>
<tr>
<td>78.38</td>
<td>39.19</td>
<td>$48.4 \times 10^{-4}$</td>
<td>$3.92 \times 10^{-6}$</td>
</tr>
<tr>
<td>88.11</td>
<td>44.06</td>
<td>$54.5 \times 10^{-4}$</td>
<td>$4.41 \times 10^{-6}$</td>
</tr>
<tr>
<td>95.07</td>
<td>47.54</td>
<td>$58.8 \times 10^{-4}$</td>
<td>$4.75 \times 10^{-6}$</td>
</tr>
</tbody>
</table>

Figure 6 shows the variations of $r^*$ and $\Delta G^*$ with $\gamma_{NS}$ for $T = 573$ K. The graph clearly indicates an increasing mode with a linear and curvy variations in $r^*$ and $\Delta G^*$, respectively as predicted from the theory. There are also points on the graphs indicating the measured values of $r^*$ ranging between 16-48 nm corresponding to nanodot’s sizes of 32-95 nm. These results strongly suggest that both critical parameters are the functions of surface energy at the nucleus-surface interface and contact angle between the nucleus and substrate. The V-W growth mode was found to fulfill equation (1), since the sum of $\gamma_{LS} \cos \theta + \gamma_{NS}$ is always greater than $\gamma_{LS}$. An example of simulated parameters for $T = 573$ K is tabulated in Table 1, together with the measured diameters and $r^*$. Analysis results for $T = 673$ K (not shown) are generally similar in the trends of graphs but slightly different in the appearance of nanodots. It is important to note that, there is a very good compromise between the theoretical and experimental data, especially the value $\gamma_{LN}$ $= 1.44 \times 10^{-4} \text{ J cm}^{-2}$ [10,11].
It is recommended that from the above discussion the self-assembled growth occurring in nc-Si nanodots is dominated by the Volmer-Weber growth mechanism. Accordingly, the oval-shaped islands are formed due to atomic binding contributed by neighbouring atoms according to a relationship given by equation (1). However, the expected growth process satisfies the classical nucleation theory on the assumption that the nucleus formed must be in a cap-like shape (part of spherical shape). The contradiction between the oval and the cap-like shape can be resolved by adapting reduced amount of atomic binding amongst the neighbouring atoms, with the original relationship remain unchanged. In other words slight modification is required in order that the sum of $\gamma_{LS}\cos \theta + \gamma_{NS}$ is just reasonably greater than $\gamma_{LS}$. Experimental verification has been successfully done and found to be in a very good agreement.

3.1 CRESE point

There is an interesting feature that can be extracted from the plot in Fig. 6. Solving the mathematical regression the following functions are obtained for the linear and curve plots, respectively, i.e., $y_1 (= r^*) = 4 \times 10^{-4} x + 10^{-8}$ and $y_2 (=\Delta G^*) = 4 \times 10^{-8} x^{2.984}$. Assuming that 2.984 ~ 3.0 ($y_2$: experimental linear equation) and both functions are equal, i.e. $Y_i = y_2$, then a cubic equation, $4 \times 10^{-8} x^3 - 4 \times 10^{-4} x - 10^{-8} = 0$ is formed. There are two possible solutions for surface energy $x$, namely $x_1 \approx 10^{-2}$ jcm$^{-2}$ and $x_2 \approx -61.6$ jcm$^{-2}$. Except $x_2$, $x_1$ is an extreme point representing positive $r^*$ and $\Delta G^*$ which lies within the practical limit. It can be considered as a point of equilibrium between the critical radius-energy and surface energy or so-called CRESE point. Further analysis on different temperatures could result in different CRESE points and a new relationship could be developed. A general relationship can be established as follows:

Rewriting eqs. (3) and (4) as

$y_1 (= r^*) = Ax^3$ and $y_2 (=\Delta G^*) = Bx + C$

Then, equating both equations to form a general CRESE equation,

$$A x^3 - Bx - C = 0$$

where

$$A = \frac{16 \pi T_m^2}{3 L_f^2 \Delta T^2} \left( \frac{2 - 3 \cos \theta + \cos^3 \theta}{4} \right),$$

$$B = \frac{-2 T^2}{L_f^2 \Delta T^2}$$

and $C$ is a constant.

Fig. 5. PL spectrum for a selected sample ($T=673$ K)

Fig. 6. Variations of $r^*$ and $\Delta G^*$ as a as function of surface energy, $\gamma_{NS}$ for $T=573$ K

Fig. 7. Variations of $r^*$ and $\Delta G^*$ as a function of contact angle, $\theta$
Equation (5) can be solved mathematically in order to determine the two CRESE points for each set of temperature.

4. CONCLUSION

The early stage of nanocrystalline silicon nanodot formation can be well described using the Volmer-Weber growth mode mechanism, based on the non-wetting oval shape cluster. Slight modifications of this oval to dome-like shape are needed in order to satisfy the liquid-solid nucleation theory. Accordingly, the sums of $\gamma_{LN} \cos \theta + \gamma_{NS}$ are just reasonably greater than $\gamma_{LS}$, and these have been verified experimentally. The self-assembly method can be applied for the fabrication of quantum devices, such as quantum dot single-electron transistor and nanobiosensor.

ACKNOWLEDGEMENT

The authors would like to thank the Ministry of Higher Education for providing financial support under the FRGS grant, Vot No. 78461.

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