Low Temperature poly-Si TFTs for Mobile Display Applications

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Abstract: Mobile display products using poly-Si TFTs with integrated drive circuits are on the market. There is a demand to integrate more components monolithically to realize system on panel. To meet the demand, lower Vth and uniform characteristics for both n- and p-channel TFTs are required. To realize TFT with lower Vth, in this paper we present a technique to crystallize poly-Si in (100) surface orientation, with a rather uniform grain size on a glass substrate. A high-pressure water vapor annealing (HWA) technique can be used to terminate poly-Si grain boundaries more effectively than conventional hydrogenation. Of particular importance to the case of gate insulators, we describe deposition methods that can be used to form low damage thin films at glass compatible temperatures.

Keywords: laser crystallization; poly Si TFTs; (100) oriented poly-Si; interface energy; gate insulator; thin film deposition techniques; high pressure water vapor annealing;

1.0 Introduction

Products using low temperature poly-Si (LTPS) TFTs, like digital still cameras, mobile phones, personal digital assistants, car navigators are on the market with integrated drive circuits, digital analog converters (DAC), timing controller etc. on the same glass substrate as the active pixel matrix, offering higher resolution and better reliability. There is a demand to integrate more components like multi-bit memory on pixel, CPU interface on the same substrate to realize system on panel [1]. Figure 1 shows the system integration roadmap for LTPS TFT LCDs.

To achieve the roadmap, lower Vth and uniform characteristics for both n and p channel TFTs are required. The main reasons for the high Vth and variation in characteristics are random orientation of poly-Si grains, grain size variation, and incomplete termination of grain boundaries, damage to the interface during gate insulator deposition, and static charge in the gate insulator.

Almost all of LTPS TFTs used in the products are based on excimer laser crystallized poly-Si. Excimer laser crystallized poly-Si grains are randomly located and randomly oriented in plane and have various sizes. Therefore TFTs fabricated using this type of poly-Si have large variations in their parameters. To avoid excessive variation in characteristics, it is desirable that the active region of TFTs are grain boundary free, which means the grain boundaries are at predetermined positions so that they can be avoided when forming a device. The device can be fabricated inside the gains and characteristics similar to c-Si can be realized [2]. Therefore many researchers are trying to control grain boundary position using various techniques [3-7]. It is important to control the location of the grains, it is also equally important to control orientation of the grains. It is known that {100} normal exhibits 1/3 of dangling bonds than {111} normal at the surface, resulting in a better interface for (100) textured films hence lower Vth [8,9]. In section 2 a model is described and based on the model a technique to crystallize poly-Si in (100) surface orientation on a glass substrate is presented.

Hydrogenation of poly-Si is indispensable to reduce defect at grains boundaries and within the grains. An emerging technique, which is suggested to be more efficient in terminating defects in poly-Si TFTs than conventional hydrogenation is summarized in section 3.

![Figure 1. System integration roadmap for LTPS TFTs. [ref. 1]](image-url)
Controlling the grain position and orientation is not the only important technology to reduce Vth. The formation of high quality poly-Si/gate insulator (SiO₂) interface at glass compatible temperature is also a very crucial process. The characteristics of the gate insulator SiO₂ currently used in poly-Si TFT production, which is deposited by RF-PECVD, are much inferior to thermal SiO₂ used in LSI. The reason for the inferiority is damage caused by the plasma at the poly-Si/SiO₂ interface and SiO₂ film itself during deposition. In section 4 promising deposition techniques that can be used to form low damage, high quality SiO₂ thin films at glass compatible temperature are described.

2.0 Crystallization of (100) Oriented Poly-Si
For the XeCl pulse laser crystallization three regimes are seen, a low laser fluence regime where the small grains up to twice the film thickness are obtained. This energy range is characterized by a combination of explosive crystallization and vertical solidification. A high energy density regime in which the laser intensity is sufficiently high so that film completely melts and epitaxial growth is suppressed and microcrystalline film is obtained. In between is a regime where large grains that exceed the film thickness by up to a factor of 30 are obtained is so called super lateral growth regime. In this regime grains as large as 200 times the film thickness are observed for multiple pulses. The SEM micrograph of a 30 nm thick film in structure a-Si(30nm)/SiO₂(40nm)/SiNx(50nm)/glass, crystallized in this regime for a hundred pulses is shown in Fig. 2. The film was Secco etched to reveal the grain boundaries. As is seen from the micrograph grains larger than 6μm are obtained. The film was crystallized at atmospheric pressure in nitrogen and no thermal heating was carried out. The x-ray diffraction spectrum of the film is shown in Fig. 3 for energies 340, 350 and 360 mJ/cm². The energy density measurement for excimer laser not quite accurate, so energy density reported here should be taken as typical values rather than absolute values. We see very strong (111) peak, other peaks like (400) or (220) are not seen, showing film has preferential {111} surface normal orientation. The intensity of the X-ray peak increased with increasing energy density up to 360 mJ/cm². If the irradiation energy was further increased no signal was observed. The film became microcrystalline or amorphous like. We found a fourth regime just below the {111} preferential orientation regime. The grains in this region have square shape as shown in Fig. 4[10].

It has been reported that, the films with (100) texture have a rectangular shape (often approximately square) with edges parallel to the <110> or <100> direction. [11] The grains in this region some time have preferential {100} orientation. This is in contrast to the observation made by other researchers [12, 13], who reported that poly-Si crystallized by excimer laser shows only {111} preferential orientation.

It has been reported that during a-Si crystallization on SiO₂ surface a (100) texture minimizes Si-SiO₂ interfacial energy, on the other hand a (111) texture minimizes the energy of the Si surface. [14] The above results can be explained considering this fact.

Figure 2. SEM micrograph of a film crystallized using 28 ns pulse width laser, showing grains as large as 6μm for 30 nm thin film.

Figure 3. X-ray diffraction (θ-2θ) of a sample crystallized at different laser fluences. 360 mJ/cm² Corresponding to SEM micrograph in Fig. 2

Figure 4. Plan view TEM micrograph of the film crystallized in the square grain region for laser pulse width 28 ns.

When irradiation energy is set such that it is just near complete melting, and the peak power is very high like in short pulse laser non homogenous melting occurs. More un-melted nuclei can be obtained in side the melt rather than at interface, during growth {111} preferential orientation is obtained to minimize the surface energy.
If the energy is set a little lower than this value then interface plays a major role and \{100\} orientation is possible as explained by our model explained hereinafter. [17] The difference in interface energy results in a difference in the melting temperature [\(\Delta T_m = T_{m(100)} - T_{m(111)}\)] and is reported to be 0.016 °C for a 0.5μm thick film. The difference in melting temperature increases with decreasing thickness, as is seen in eq. (1) [14].

\[
\Delta T_m = \frac{2\Delta\gamma V}{t\Delta S_f},
\]

where \(\Delta\gamma\) is the difference in surface energy between the (100) texture and (111) texture, \(V\) is the volume fraction, \(t\) is the thickness of the film and \(S_f\) is the entropy of fusion per atom [14]. As the temperature difference is inversely proportional to thickness, it is expected to be about an order higher for 40nm-thick Si film.

The difference in the melting temperature between (100) texture and other textures seems small but when taking into account the large increase in the reflectivity of Si when it melts (a semiconductor–to-metal transition) the difference is significant and a partial molten configuration is possible in which most of the (100)-textured grains remain unmelted/solid while grains with a texture other than (100) melt. The energy density required to maintain a pool of fully molten Si is higher than the energy density required to heat solid Si up to the threshold of melting [14]. In fact solid (100) lamella floating in the melted Si has been observed in the CW Ar laser crystallization of Si films much thicker than 40 nm [15, 16].

Our model reflects the above observation. In our model, a-Si film is irradiated at energy close to the melting point of poly-Si film. As the melting point of poly-Si is about 200 °C higher than of a-Si film, in the first pulse the whole film is melted. It is very difficult to control energy close to melting point using 28 ns pulse width laser, therefore to demonstrate the model a laser with 150ns pulse width was used. The energy density at which textures other than (100) melt in a 40nm a-Si/SiO2/SiN/glass structure was found to be 450 mJ/cm² for a 150 ns pulse width laser. Again this energy should be considered typical values rather than absolute values. This energy varies with the irradiation conditions, such as pulse duration, the irradiation environment and the underlying layers. In the next pulse, most of the textures other than (100) melt, while some of (100)-textured grains remain, because the (100) texture has a higher melting point than other textures. When the third pulse is irradiated, the number of (100)-textured grains increases much more than after the second pulse, as the solid (100) texture/lamella from the previous pulse may act as a seed and also new (100) grains will be added as a matter of course. This process is repeated after subsequent pulses, as is seen schematically in Fig.5. With a sufficient number of pulses, the whole film eventually becomes (100) textured [17].

Figure 6 shows SEM photographs of film crystallized at an energy of 450 mJ/cm² using 200 pulses. As said above, the films with (100) texture have a rectangular shape (often approximately square) with edges parallel to
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moved in 200 nm steps so that about 2000 measurements were performed. Figure 8 shows the standard triangular representation of the orientation of the film irradiated with 200 pulses as obtained from EBSP measurement. For this measurement about 2000 points were taken for 10 µm×10 µm area. Almost all the 2000 points cluster within 10 degrees of the (100) surface normal. This was the first time that such a strong (100) texture was obtained for excimer-laser crystallized Si film on a glass substrate [17]. As stated in the introduction, the (100) surface has 1/3 of dangling bonds than that of the (111) surface, therefore, lower interface states are expected. With lower interface states we can expect sharp subthreshold swing, hence lower Vth.

3.0 High-Pressure Water Vapor Annealing

High-Pressure Water Vapor Annealing (HWA) post annealing of top gate TFTs has been shown to improve characteristics [18]. In our group we have focused on bottom gate TFTs. As bottom-gate poly-Si TFT structure is considered beneficial to form a good-quality Si/SiO2 interface. This is because a SiO2 layer and an amorphous silicon layer can be consecutively deposited by PECVD without breaking the vacuum, hence clean Si/SiO2 Interface [19]. Recently we have shown that HWA is also effective in the case of bottom gate TFTs. Figure 9 shows schematic cross section of the bottom-gate TFT used in this study. A Mo layer was formed by magnetron sputtering and then patterned to form a gate electrode. For gate insulator 50-nm-thick silicon nitride (SiNx) and 40-nm-thick SiO2 were deposited using RF PECVD. A 40-nm-thick a-Si film was subsequently deposited by PECVD. A 150-mm-long, 0.5-mm-wide pulsed excimer laser beam at 99% overlap was used to crystallize a-Si. Phosphorus ions (dosage $7 \times 10^{12}$ cm$^{-2}$) were doped to form lightly doped drain (LDD). Then, phosphorus ($3 \times 10^{14}$ cm$^{-2}$) and boron ($1 \times 10^{15}$ cm$^{-2}$) ions were implanted to form the source and drain for N and P-channel TFTs, respectively. The source and drain regions were activated by thermal annealing, followed by delineation of poly-Si islands. Two different post annealing processes were carried out to evaluate the effect of HWA on hydrogenation. One was conventional hydrogenation, in which hydrogen extracted by thermal annealing (at 400°C for 2 h in N2 gas) from the capped hydrogen rich SiNx interlayer was used. The other process was HWA processing after the deposition of the interlayer, in which TFTs were annealed by high pressure steam at 350 °C and 1 MPa for 2h. Finally, aluminum metallization was performed [19].

The transfer characteristics of 28 poly-Si TFTs were measured over the entire surface of the 300×350mm$^2$ glass substrate. The measurement was carried out with Vds ±10 V. The channel width W and channel length L of the TFTs were 20µm and 7µm, respectively. Performance variations over these TFTs are given in Fig. 10. Figure 10(a) shows characteristics for conventional hydrogenation, and Fig. 10(b) shows for HWA. Figure 10 superimposes twelve out of 28 TFT transfer characteristics for simplicity. Improvement was seen for Vth uniformity and mobility for both N- and P-channel TFTs. The Vth variation of TFTs decreased from 0.19 to 0.11V for N-channel and from 0.13 to 0.08V for P-channel in terms of standard deviations, and carrier mobility was enhanced by 18% for N-channels and 10% for P-channel TFTs compared with conventional hydrogenation. Subthreshold swing was also improved, which shows that HWA improves Si/SiO2 interface for the bottom-gate TFTs too which lies under the poly-Si.
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The improvement in the characteristics may be due to termination of dangling bonds by forming SiH, SiO and Si-OH bonds more effectively compared to conventional hydrogenation.

4.0 Techniques to deposit high quality SiO2

The reason for the poor quality of PECVD SiO2 is damage by plasma irradiation during deposition and a large number of Si-H and Si-OH bonds. It is essential to use a deposition technique which is damage free and has a minimum number of Si-H and Si-OH bonds. One way to realize this is to separate plasma and deposition regions and use a high density plasma to dissociate SiH4 more effectively. Such a plasma source is illustrated in Fig. 11. Recently a surface wave plasma CVD process using slotted waveguide antenna for deposition of gate insulators has been reported [20]. In the surface wave plasma, the plasma is concentrated near the anode therefore the substrate is not exposed to the plasma. It has a structure like that shown in Fig. 11. Another interesting characteristic of a surface wave plasma reactor is that the electron temperature decreases with increasing power. Lower electron temperature means, less damage to the depositing film. They used slotted waveguide antenna to introduced 2.45GHz microwave in to the 40cm×40cm chamber. They fabricated TFTs using microwave plasma CVD on an SOI wafer [20]. No difference in characteristics was found between TFTs fabricated using conventional thermal oxide and film deposited using surface wave excited PECVD. Although the TFTs were fabricated on an SOI wafer and high temperature was used and further work is required to utilize this technology for large substrate, the technique clearly showed potential for deposition of high quality SiO2.

Another interesting technique which has potential in deposition of high quality SiO2 film is Electron Cyclotron Resonance (ECR) CVD. Using parallel synchronized-phase micro-wave introduction, plasma stream with width 9cm, length 60cm was reported [21]. As plasma density in ECR CVD can be as high as 10^{12}/cm^3, even if the substrate is situated some distance from the plasma a high incident density of radicals is possible. In fact direct oxidation of Si at a rate as high as 6nm/min was reported in the initial stage of oxidation with good thickness uniformity on a large substrate [21].

High quality PECVD SiO2 films were also reported using the same ECR plasma source. Since the ECR plasma highly decomposes the gases, SiH4 and O2 can be used instead of SiH4 and N2O or tetraethylorthosilicate (TEOS) gas. They measured RF(1MHz) and quasi-static capacitance-voltage (C-V) characteristics for a 59nm thick PECVD deposited film. From the difference of the capacitance between high and quasi-static frequency curve, Dit as small as 7×10^0/cm^2V^-1 was reported at the mid gap of Si [21], which is surprisingly small for a low temperature deposited film making it suitable for LTPS TFTs.

Figure 10. Twelve superimposed TFTs (a) conventional hydrogenation (b) HWA [ref. 19]

Figure 11. A schematic illustration of a surface wave plasma source
techniques for damage free deposition of SiO₂. However a lot of work is required to make them suitable for LTPS TFTs. HWA seems to improve the characteristics for both bottom gate and top gate TFTs. Combining the techniques described above, we believe that it will be possible to achieve the characteristics dictated by the roadmap.

References
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