SUMMARY  Microwave photoconductivity decay (μ-PCD) method was applied to evaluate the effects of chemical composition and Ar⁺ plasma induced damage on the bulk and the surface states in amorphous In-Ga-Zn-O (a-IGZO) films. It was found that the peak reflectivity signal in the photoconductivity response increased with decreasing the Ga content, and had a strong correlation with the a-IGZO transistor performances. In addition, the peak reflectivity signals obtained after various Ar⁺ plasma treatment duration were well correlated with the transistor characteristics. With Ar⁺ plasma treatment, the peak reflectivity signal decreased in accordance with degradation of transistor characteristics. The μ-PCD method was found to be a very useful tool not only to evaluate the bulk and the surface states, but also to predict the performance of a-IGZO transistors subjected to various plasma processes in the production.

key words: microwave photoconductivity decay, a-IGZO, oxide semiconductor, photoconductivity response

1. Introduction

Amorphous In-Ga-Zn-O (a-IGZO) is attracting considerable attention as it could be a high-performance channel material in thin film transistors (TFTs) for next generation displays [1]–[5]. The control and monitor of the channel composition, the surface states and the interface states between the protect and the channel layer is of importance because the channel composition directly influences the TFT performance [6]–[8], and the surface and the interface states are often created by various manufacturing processes such as plasma process [9], [10]. Microwave photoconductivity decay (μ-PCD) method is known as a non-destructive and contactless technique and widely used for process control and characterization of defects in various semiconductors [11]–[15]. In the previous study, for a-IGZO films, we found that the peak reflectivity signals of the photoconductivity responses measured by μ-PCD were governed by the film quality (bulk states) that affects the a-IGZO transistor performance [16]. In the present study, we carried out μ-PCD measurements to evaluate the bulk states of a-IGZO films with various Ga contents. In addition, we applied the μ-PCD measurements to evaluate the effect of Ar⁺ plasma induced damage to the surface states of a-IGZO films and the influence of the SiO₃ passivation process. We discuss the correlation between the peak reflectivity signals of photoconductivity responses and transistor performances in terms of chemical composition and the surface states.

2. Experimental

The μ-PCD measurements were carried out using a commercially available lifetime measurement system (KOBELO Research Institute, LTA-1820SP). To obtain a sufficient signal-to-noise ratio, the differential μ-PCD technique for microwave detection was used [15]. In this system, excess carriers are generated by a laser irradiation to increase the conductivity, resulting in a change in the microwave reflectivity of the specimen. The change in the reflected microwave power is directly related to the change in the excess carrier density [11], [12], [15]. A semiconductor laser-pumped third harmonic yttrium lithium fluoride (YLF) pulse laser was used for the carrier generation. The wavelength was 349 nm (3.55 eV), and the pulse width was 15 ns. The energy of the exciting light was 3.55 eV and higher than the band gap energy of the a-IGZO film (approximately 3.2 eV) [17], [18]. The penetration depth of the laser light was estimated to be approximately 300 nm. The photon flux of the laser pulse was 1 × 10¹⁵ cm⁻². The microwaves (26 GHz) were guided to both the signal and the reference waveguides, and then irradiated on the specimen. The reflected microwaves passed through the waveguides to the detector, and the reflectivity differences between the irradiated and non-irradiated (reference) areas were measured.

The X-ray photoelectron spectroscopy (XPS) measurements were applied to investigate the electronic structure in a-IGZO films. A Physical Electronics PHI Quantera SXM with a monochromatic Al Kα source (energy 1486.7 eV) was used. The energy calibration was performed by setting the binding energy (BE) of the Au 4f₇/₂ = 84.0 eV, Cu 2p₃/₂ = 932.7 eV and Ag 3d₃/₂ = 368.3 eV. The measurement was carried out at room temperature. The analysis area was 100 µm in diameter. The emitted photoelectrons were detected at a take-off angle of 30°.

Figure 1 shows a schematic illustration of the cross-sectional structure of an a-IGZO TFT with a bottom gate structure. An n-type crystalline Si wafer with a 200-nm thick thermally-grown SiO₂ layer was used as the gate electrode and the gate insulator, respectively. The a-IGZO
films with a thickness of 50 nm were then deposited by co-sputtering of Ga2O3, In2O3 and ZnO targets at room temperature. Three samples were prepared. The compositions of the films confirmed by XPS measurements were In:Ga:Zn = 1:0.7:1, 1:1:1, and 1:2:1, respectively. The input power was controlled in order to obtain the intended chemical composition, while the process pressure of a mixed Ar and O2 gases was fixed at 5 mTorr. For the evaluation of the effect of Ar+ plasma induced damage and the influence of the SiOx passivation process, a-IGZO film with a thickness of 50 nm was deposited by DC sputtering using an InGaZnO4 target. The input power was 200 W and the process pressure of a mixed Ar and O2 gases was 1 mTorr. The a-IGZO films were photolithographically patterned, and the specimen was annealed at 350°C for 1 h in air. The Mo source and drain electrodes were then deposited at room temperature by shadow-mask sputtering. The channel length (L) and width (W) were 10 μm and 200 μm, respectively. For the evaluation of the effect of Ar+ plasma induced damage, Ar+ plasma treatments were carried out under the following conditions: RF power of 50 W, process pressure of 0.8 Torr at room temperature. The transfer curves were measured using a semiconductor parameter analyzer (Keithley 4200) at room temperature under dark condition.

The a-IGZO films for μ-PCD and XPS measurements were deposited on glass substrates and n-type crystalline Si wafers, respectively. The deposition was done either by co-sputtering of 3 targets or DC sputtering using an InGaZnO4 as mentioned above. The thickness of the film was 50 nm or 100 nm. Then, the films were annealed and treated with Ar+ plasma under the same condition described above.

3. Result and Discussion

3.1 Evaluation of Bulk States as a Function of Ga Content

Figure 2 shows the transfer curves of IGZO TFTs with various Ga contents. Table 1 summarizes the transistor performances. The saturation mobilities (μSAT) were extracted from $I_{DS} = \mu_{SAT} \frac{W}{2L} (V_{GS} - V_{TH})^2$, where $I_{DS}$ is the drain-source current, $V_{GS}$ is the gate-source voltage, and $C_{OX}$ is the gate insulator capacitance per unit area. The a-IGZO TFT with the lowest Ga content showed the highest driving current and μSAT and the lowest (best) sub-threshold swing (SS) values. The similar compositional dependence has been reported in the literatures [6]–[8], where the same trend was found that $\mu_{SAT}$ increased with decreasing Ga content.

Figure 3 shows the photoconductivity responses of a-IGZO films with various Ga contents. Each film exhibited a non-exponential decay profile containing a slow decay component. It can be seen that the peak reflectivity signals increase with decreasing the Ga content. On the other hand, the slope of the slow decay hardly varies with the Ga contents. We infer that the slow decay is related with the trapping processes by defects and/or by other factors. In the previous study, we evaluated the μ-PCD mapping over a wide compositional space of a-IGZO film deposited on a 6-inch glass substrate by co-sputtering of 3 targets [16]. In the map of the peak reflectivity signal, the same tendency depending on the Ga content was observed.

Figure 4 plots the values of $\mu_{SAT}$ against the peak reflectivity signals. The peak reflectivity signals become higher with $\mu_{SAT}$, and the linear relationship between $\mu_{SAT}$ and the
peak signals is clearly seen. This is consistent with previous report showing that the peak reflectivity signal has a strong correlation with $\mu_{\text{SAT}}$, and implies that the peak reflectivity signal directly corresponds to the density of the sub-gap states of a-IGZO bulk region [16]. Also, it has been reported that Zn contributes to the modulation of the shallow tail states below the conduction band [6]. It is reasonable to suppose that the sub-threshold characteristics were improved with decreasing the Ga content and/or increasing the Zn content. In addition, electron mobility strongly depends on the carrier concentration [2]. The carrier concentration of the present films are around $10^{18}$ cm$^{-3}$ and increases with decreasing Ga content. Within the present carrier concentration range, the number of effective recombination and/or trapping centers decreases with increasing carrier concentration as a result of the Fermi level shift toward the conduction band minimum. This leads to an increase in the saturation mobility. Therefore, we infer that the density of the sub-gap states, including the tail states, that act as trapping and/or recombination centers, is lower with decreasing the Ga content. Hence, in the present study, the a-IGZO film with low Ga content exhibited more photo-generated carriers survived during the laser pulse illumination, that leads to higher peak reflectivity signal, compared to that with high Ga content.

3.2 Evaluation of Surface and Interface States

3.2.1 Ar$^+$ Plasma Induced Damage

Figure 5 shows the transfer curves of a-IGZO TFTs as a function of Ar$^+$ plasma treatment duration. As shown in the figure, the untreated (0 s) TFT shows good transfer curve. $\mu_{\text{SAT}}$ was found to be 8.3 cm$^2$/Vs and $SS$ was 0.33 V/dec. On the other hand, most TFTs with the Ar$^+$ plasma treatment show a drastic increase in the off-current. Conducting behaviors with no turn-off is also observed already for the 3 s plasma treatment. It has been reported that the plasma treatment creates oxygen deficiency for the metal oxide materials, resulting in an increase in electron carriers on the surface region [19], [20].

The inset of Fig. 5 shows the peak reflectivity signals of photoconductivity responses as a function of Ar$^+$ plasma treatment duration. The peak reflectivity signals become lower with increasing the treatment duration. It is clear that the peak reflectivity signal have a strong correlation with the Ar$^+$ plasma treatment duration. This result indicates that the peak reflectivity signal strongly reflects surface state, since the bulk state is hardly influenced by the plasma treatment. Considering this result and the previous report [16], it is suggested that the peak reflectivity signal consists with not only the bulk states but also surface states. It is reasonably assumed that the peak reflectivity signal is attributed to the recombination and/or trapping processes that originated in bulk and surface states. Therefore, the peak reflectivity signal decreased due to the increasing the surface states caused by Ar$^+$ plasma treatment.

XPS analysis was carried out in order to investigate the surface electronic structure of a-IGZO films with Ar$^+$ plasma treatment. Figure 6 shows the valence band (VB) spectra of a-IGZO films with various Ar$^+$ plasma treatment duration. The VB of a-IGZO consists mainly of O 2p and Zn 3d states. The VB edge was defined as the point where the linear extrapolation along the lower BE side of O 2p (around
4–5 eV) intersects the baseline. The values of the VB edge are 3.8 eV and 3.4 eV for a-IGZO films with and without 300 sec Ar plasma treatment, respectively. Similar energy shifts were also observed in Zn 3d, O 1s, In 4d and Ga 2p (the spectra of O 1s and Ga 2p are not shown) depending on the Ar+ plasma treatment duration. It is likely that this observation correlates with Fermi level shift [21]–[23] rather than the charging up effect. Min et al. have reported that Fermi level shifts by oxygen deficiency in the ZnO films [21]. In our study, the films exposed to Ar+ plasma showed conducting behavior as shown in Fig. 5. Thus, it is inferred that the Fermi level approaches to the conduction band (CB) edge. As the Fermi level shifts closer to the CB edge, the conduction electrons increase due to thermal excitations. Hence, we suggest that the BE shift with Ar+ plasma treatment is mainly due to the Fermi level shift. In particular, it is likely to consider that the Fermi level of the a-IGZO for the 300 s Ar+ plasma treatment is located at the CB level, so that transport mechanism is nearly metallic. The inset of Fig. 6 shows enlarged VB edge of the a-IGZO films with various Ar+ plasma treatment duration. As shown in the figure, the intensity near BE ~ 6 eV decreases with the Ar+ plasma treatment, indicating an increasing of the oxygen deficiency by exposing to Ar+ plasma [24].

3.2.2 Effects of SiO2 Passivation

To confirm that the peak reflectivity signal reflects the surface states as well as the bulk states, we investigated the influence of SiO2 passivation layer on a-IGZO film for the photoconductivity response. It has been reported that the characteristics of TFTs are influenced by plasma-enhanced chemical vapor deposition (PECVD) process depending on the process conditions. Therefore, for passivation of the a-IGZO film, SiO2 layer was deposited by PECVD at 150°C on the a-IGZO film since the low temperature (~ 150°C) PECVD of SiO2 passivation layers suppress the surface states at the back channel and improved TFT characteristics [25]. The thickness of the SiO2 layer was 50 nm. Figure 7 shows the photoconductivity responses of a-IGZO films with and without SiO2 passivation layer. It can be seen that the a-IGZO film with the SiO2 passivation layer exhibits higher peak reflectivity signal compared to that without the SiO2 passivation layer. It is inferred that the optimized SiO2 passivation layer decreases the surface states of a-IGZO film, that leads to more photo-generated carriers survived during the laser pulse illumination. This result supports the mechanism proposed above that the peak reflectivity signal consists with the bulk states and the surface states. Therefore, contrary to the Ar+ plasma irradiation, it was observed that the peak reflectivity signal of the a-IGZO with the SiO2 passivation layer was increased due to the reduction of the surface states.

4. Conclusion

We investigated the μ-PCD measurements to evaluate the a-IGZO films as functions of the chemical composition and the Ar+ plasma treatment duration. The peak reflectivity signal increased with decreasing the Ga content in the film, and the correlation was confirmed between the peak reflectivity signals and the transistor performances of the a-IGZO TFTs. It is suggested that the peak reflectivity signal directly corresponds to the density of sub-gap states of a-IGZO bulk film. In addition, it was found that the peak reflectivity signal decreased with the Ar+ plasma treatment due to the increase of the surface states caused by the defects such as oxygen deficiency. On the other hand, the peak reflectivity signal increased due to the reduction of the surface states originating from the SiO2 passivation layer. Hence, it is concluded that the peak reflectivity signal is strongly affected by the surface states as well as the bulk states. μ-PCD measurement is really effective to characterize the compositional dependence and the effect of the plasma process of a-IGZO films in research and development stage and expected to apply to the process control and management in both in-line and off-line inspection tools.

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References


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