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Conference Paper · May 2016
DOI: 10.1117/12.2223389

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Gibbs Free Energy Assisted Passivation Layers

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ABSTRACT

Reduction of surface leakage is a major challenge in most photodetectors that requires the elimination of surface oxides on etched mesas during passivation. Engineering the passivation requires close attention to chemical reactions that take place at the interface during the process. In particular, removal of surface oxides may be controlled via Gibbs reactivity. We have compared electrical performance of type-II superlattice photodetectors, designed for MWIR operation, passivated by different passivation techniques. We have used ALD deposited Al₂O₃, HfO₂, TiO₂, ZnO, PECVD deposited SiO₂, Si₃N₄ and sulphur containing octadecanethiol (ODT) self-assembled monolayers (SAM) passivation layers on InAs/GaSb p-i-n superlattice photodetectors with cutoff wavelength at 5.1 µm. In this work, we have compared the result of different passivation techniques which are done under same conditions, same epitaxial structure and same fabrication processes. We have found that ALD deposited passivation is directly related to the Gibbs free energy of the passivation material. Gibbs free energies of the passivation layer can directly be compared with native surface oxides to check the effectiveness of the passivation layer before the experimental study.

Keywords: Superlattice, Photodetector, InAs/GaSb, ALD, Gibbs Free Energy, Passivation

1. INTRODUCTION

Photodetectors with dry or wet etched mesa side walls often lead to increased surface leakage currents that degrade detector performance and Type II InGaAs/GaSb superlattice (T2SL) photodetectors operating in MWIR and LWIR regions are no exception. Type-II InAs/GaSb superlattices has recently gained a lot of interest for the infrared photodetector applications due to many advantages like suppression of Auger recombination [1], and interband tunneling [2], and the ability to tailor the bandgap within most of the infrared region by changing the thickness of the constituent InAs and GaSb layers [3]. InAs/GaSb superlattices are periodic structures consisting of alternating layers of nanometers thick InAs and GaSb layers. Surface passivation is extremely critical in T2SL InAs/GaSb photodetectors, due to large number of very thin alternating layers. Different etch rates of InAs and GaSb during mesa definition lead to roughness on the mesa side walls. A conformal coating can be critical to cover all the tiny undulations on mesa side walls. Furthermore, components of InAs/GaSb SL are chemically very reactive. They are easily oxidized and a native oxide layer of several nanometers thick is quickly formed upon exposure to air. Adsorbed oxygen diffuses through the surface, reacts with Ga, Sb, In and As atoms then forms native oxides such as Ga₂O₃, Sb₂O₃, In₂O₃, As₂O₃ and InAsO₃ some of which are conductive [4,5].
This mechanism is responsible for the formation of additional conductive channels and consequently, leads to a large surface leakage. So far, many passivation techniques have been proposed to overcome these surface leakages. Dielectric material deposition [6, 7], sulfurization [8, 9], overgrowth with wide band gap materials [10] and polymer film coating [11] techniques can be listed as successful methods for passivation. Passivation is expected to limit oxidation of the side walls, saturate dangling bonds to avoid surface states and self clean [12] already formed native oxides at the etched surfaces. Sulfur containing solutions satisfies the dangling bonds and cleans the conductive native oxides by replacing the oxygen atoms with sulfur atoms. Although sulfurization is a very effective method, sulfur passivation does not have long term stability. In addition, sulfur solutions may damage the crystal surfaces causing losses in the optical performance. Dielectric deposition by plasma enhanced chemical vapor deposition (PECVD) is a very common technique in industrial production of photodetectors but it may create plasma damage and thermal damage on the crystal structure. On the other hand, atomic layer deposited (ALD) metal oxides seem to satisfy all passivation needs by providing damage free, low stress, low interface state density and low pin hole density coatings. In this work, we have compared the electrical performances of type-II superlattice pin photodetectors to determine the effectiveness of ODT SAM passivation, PECVD grown SiO₂ and Si₃N₄ passivation, ALD grown Al₂O₃, HfO₂, TiO₂ and ZnO passivation. Electrical performance of the photodetectors has been compared against unpassivated reference photodetectors to determine the effectiveness of the ALD deposited oxides. The same crystal structure and same fabrication processes were used to test different passivation materials.

2. EXPERIMENTAL

The SL structure was designed for MWIR operation with design cutoff wavelength of 5 μm. Figure 1a. illustrates structure of the p-i-n design type-II superlattice crystal design with corresponding thicknesses and doping concentrations. The pin part contains 140 periods 8 monolayers (MLs) of InAs/8 MLs of GaSb:Be (p=1.5x10¹⁷ cm⁻³) p-type layers, 40 periods 8 MLs of InAs / 8 MLs of GaSb i-layers, 60 periods 8 MLs of InAs:Te (n: 5x10¹⁷ cm⁻³)/8 MLs of GaSb n-type layers. 1000 nm GaSb:Be (p=1.0x10¹⁷ cm⁻³) and 20 nm InAs:Te (n: 5x10¹⁷ cm⁻³) layers introduced as p contact and n contact layers, respectively [13]. Figure 1b shows measured cut-off wavelength of the designed SL structure after optical response measurements which are done at 77 K by using a FTIR. Fotodetectors with mesa sizes from 100 x 100 μm² to 700 x 700 μm² are fabricated using phosphoric acid based solution (Figure 1c). Ohmic contacts were deposited by evaporating 5 nm titanium and 200 nm gold [14]. For ODT passivation, the sample was immersed in 1mM solution of ODT (Aldrich, 99%) in ethanol and was left in the solution for 48 hours at 60 °C. SiO₂ and Si₃N₄ films were coated in PlasmaLab 8510C reactor at 160 °C and the process was carried out under the pressure of 0.5 Torr and RF power of 9 W. Flow rates were 180 sccm SiH₄ (%2 in N₂), 225 sccm N₂O for SiO₂ and 45 sccm NH₃ for Si₃N₄. Final thickness of the passivation layer was about 250 nm. Al₂O₃, HfO₂, TiO₂ and ZnO depositions carried out in atomic layer deposition system (Savannah 100) with 150 °C as the substrate holder temperature under constant 20 sccm N₂ gas flow. Growth of thin films have been done by delivering 0.015 s water vapor (H₂O) and related precursor gas pulses into the chamber in a sequential manner. A wait time of 20 s was added after each pulse to ensure surface reactions to take place. 0.015 s trimethylaluminum (TMA), 0.15 s tetrakis (dimethylamido) hafnium (TDMAHf), 0.15 s tetrakis (dimethylamido) titanium (TDMATi) and 0.2 s dimethylzinc (DMZn) precursor gas pulses used for Al₂O₃, HfO₂, TiO₂ and ZnO film depositions, respectively. Passivated samples along with reference samples were bonded to a 40 pin dual in-line package (DIP) chip carrier for electrical measurements. Figures 1d shows image of the sample bonded to a 40 pin DIP chip carrier.
Figure 1: (a) structure of the p-i-n design superlattice crystal with corresponding thicknesses and doping concentrations. (b) Measured cut-off wavelength of the SL structure. (c) Schematic cross section of the InAs/GaSb SL p-i-n photodiode structure. (d) Image of the sample bonded to a chip carrier, inset shows close view of the bonded sample.

3. RESULTS AND DISCUSSIONS

Dark current and dynamic resistance measurements have been done at 77K to investigate electrical performance of the passivation layers. We have used unpassivated control samples along with passivated samples. The photodetectors were put into an aluminum box and immersed into liquid nitrogen filled container. HP4142OA source-measure unit has been used to measure dark current voltage-current characteristics of the samples. Figure 2 shows the measured dark current density vs. applied bias voltage characteristics of the unpassivated and passivated 400x400 µm single pixel test diodes at 77K. Dots show dynamic resistance area product (RA) vs. applied bias voltage for selected photodiodes. At -0.1 V reverse bias, dark current density is measured as 3.3 x 10^-8 A/cm², this measurement yielded RA product value of 2.5 x 10^6 Ωcm², for the Al₂O₃ passivated photodetectors. Photodetectors passivated with a thin layer of Al₂O₃ shows the lowest dark current and highest dynamic resistance area product values for all bias voltages. Si₃N₄ is clearly the worst passivation material in this group. Photodetectors with ODT passivation show almost the same electrical characteristics with Al₂O₃ passivated photodetectors up to 0.1 V reverse bias beyond which the slope of the dark current increases when compared with detectors that are passivated with Al₂O₃. At low temperatures, currents in the photodetector are diffusion limited near zero bias. At larger biases, weak fluctuations in the already low current in ODT and Al₂O₃ are suggestive of trap assisted tunneling. Our results suggest that trap assisted currents are dominant for higher voltages for ODT passivation. Since the ODT chain is 2 nm long and the process is self-limited, some trap states on surface of ODT layer may still affect the electrical performance of the detector at high voltages through tunneling. Our electrical measurements show that Al₂O₃ passivation results in lower dark current than industry standard SiO₂ passivation at the same temperature.
Relationship between dark current densities versus inverse temperatures under 0.5 V reverse bias is shown in Figure 3. The temperature dependent I-V curve is dominated by diffusion current at high temperatures, generation-recombination current at mid temperatures and trap related currents at low temperatures. For temperatures lower than 70 K, all samples are dominated by surface related trap currents which give information about quality of the passivation. $\text{Al}_2\text{O}_3$ passivation results in at least an order of magnitude better performance than the closest competitor. This suggests that $\text{Al}_2\text{O}_3$ passivation reduces native oxides [12] eliminating surface states and preventing current flow through the surface channel. ODT and $\text{SiO}_2$ passivated samples show almost the same performance in this voltage range. Arrhenius type behavior has been observed for all samples which is an indication of diffusion dominated behavior. Arrhenius type behavior yielded activation energies of approximately 0.23 eV which is close to the device band gap.

Figure 2: Dark current density vs. applied bias voltage curves for passivated and unpassivated 400 x 400 µm diodes at 77K. Dots show differential dynamic resistance area product vs. applied bias voltage characteristics for selected diodes (same color coding).

Figure 3: Temperature dependent dark current density for unpassivated and passivated type-II InAs/GaSb superlattice 400 x 400 µm photodiodes at -0.5 V bias voltage.
Figure 4 shows surface resistivity of the photodetectors studied in this work versus Gibbs free energy. Vertical lines show Gibbs free energy of the conductive native oxides. Surface and bulk contributions to the dark current can be approximated as $1/R_0A = [(1/R_0A)_{\text{bulk}} + (1/r_{\text{surface}})(P/A)]$. Using this equation, the surface part of the resistance-area product can be calculated from the slope of the graph. Calculated surface resistivity values at zero bias are $2.31 \times 10^7$, $6.49 \times 10^5$, $2.99 \times 10^5$, $1.04 \times 10^4$, and $6.07 \times 10^4 \ \Omega \ cm$ for Al$_2$O$_3$, HfO$_2$, TiO$_2$, ZnO passivated, and reference samples, respectively. It is clear that Al$_2$O$_3$ is much better than the other passivation materials. From the graph it is also clear that surface resistance is increasing with decreasing Gibbs free energies. Lower Gibbs free energy allows Al, Hf or Ti atoms to replace Ga, In, As, or Sb atoms in the native conductive oxides. This process is called self clean process. Al$_2$O$_3$ and HfO$_2$ have lower Gibbs free energies than all possible conductive native oxides which makes them better passivation material for this system. On the other hand, ZnO has higher Gibbs energy than Ga, In and As oxides, so that, it gives worse results than reference samples. TiO$_2$ has higher Gibbs free energy than Ga$_2$O$_3$, but it still gives better electrical results than reference samples. Figure 4 suggest that ALD deposited metal oxides should have lower Gibbs free energies than native surface oxides to perform better as passivation materials.

![Surface resistivity vs Gibbs free energy graph for the different passivation materials. Vertical lines show Gibbs free energies of the conductive native surface oxides.](image)

Removal of native oxides during passivation (self-cleaning) is critical for a successful passivation, but it requires passivation material to have lower Gibbs free energy than surface native oxides to work properly. Gibbs free energies of the ALD oxides are -377.9 kcal/mol, -260.1 kcal/mol, -211.0 kcal/mol and -149.4 kcal/mol for Al$_2$O$_3$, HfO$_2$, TiO$_2$ and ZnO molecules, respectively [15]. The Gibbs free energies of Ga$_2$O, Ga$_2$O$_3$, In$_2$O$_3$, As$_2$O$_3$, As$_2$O$_5$ and Sb$_2$O$_3$ are -75.3 kcal/mol, -238.6 kcal/mol, -198.6 kcal/mol, -137.7 kcal/mol, -187.0 kcal/mol and -151.5 kcal/mol, respectively [4,5]. Gibbs free energy of Al$_2$O$_3$ is lower than all ALD oxides and all surface native oxides investigated, in this study. Gibbs free energy analysis suggests that Al$_2$O$_3$ is the best candidate as a passivation material for InAs/GaSb system [6]. Due to favorable Gibbs free energies, Al$_2$O$_3$ formation is energetically preferred to native oxide of In, As, Ga and Sb atoms. Alternatively, As atoms that form As$_2$O$_3$ or In atoms in an In$_2$O$_3$ molecule are replaced by Al$^{3+}$ atoms of the TMA molecule. Similar reaction pathways with oxides of other metal atoms are also possible. These are the, so called, interfacial self cleaning reactions of surface oxides [16]. Timm et al. using X-ray photoelectron spectroscopy (XPS) has shown that deposition of Al$_2$O$_3$ on InAs strongly reduces In and As oxides [16]. Similar work on GaAs has also shown strong reduction of Ga and As oxides. Work on electrical characterization of MOS capacitors fabricated on GaSb has demonstrated...
strong suppression of Sb$_2$O$_3$ due to Al$_2$O$_3$ deposition [17]. We predict that these self-cleaning reactions remove surface oxides and reduce current channels on the mesa side walls. Furthermore, the conformal coating achieved by atomic layer deposition creates a perfect protective layer against environmental effects especially against long term oxidation of the uneven and rough etched mesa side walls. Conformal coating of the rough surfaces also satisfies dangling bonds more efficiently while eliminating metal oxides via self cleaning process. ALD passivation is a perfect candidate for future industrial photodetector technology due to its “self healing” ability.

IV. CONCLUSIONS

Different passivation methods tested for the type-II InAs/GaSb superlattice infrared photodetector system. Dark current densities and surface resistances of the photodetectors were measured to compare the performance of the passivation techniques. We found that ALD deposited HfO$_2$ and TiO$_2$ are better than reference detectors but they are not as good as Al$_2$O$_3$ passivated detectors. ZnO passivated detectors are worse than the reference detectors. These results showed that effectiveness of the ALD deposited passivation is directly related to the Gibbs free energy of the passivation material. Gibbs free energies of the passivation layer can directly be compared with native surface oxides to check the effectiveness of the passivation layer before the experimental study. The quest for better passivation should take into account the Gibbs rule while searching for better materials and techniques.

ACKNOWLEDGMENTS

This work was supported by Anadolu University, BAP research project, (1508F600) and Akdeniz University, BAP research project, (FKA-1015-918).

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