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Skin-like self-assembled monolayers on InAs/GaSb superlattice photodetectors

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Received 16 May 2012, in final form 24 July 2012
Published 21 August 2012
Online at stacks.iop.org/JPhysD/45/365102

Abstract
We report on the effects of monolayer (ML) thick skin-like octadecanethiol (ODT, CH₃(CH₂)₁₇SH) on type-II InAs/GaSb MWIR photodetectors. Circumventing the ageing effects of conventional sulfur compounds, we use ODT, a self-assembling, long molecular chain headed with a sulfur atom. Photodiodes coated with and without the self-assembled monolayer (SAM) ODT were compared for their electrical and optical performances. For ODT-coated diodes, the dark current density was improved by two orders of magnitude at 77 K under −100 mV bias. The zero bias responsivity and detectivity were 1.04 A W⁻¹ and 2.15 × 10¹⁵ Jones, respectively, at 4 µm and 77 K. The quantum efficiency was determined to be 37% for a cutoff wavelength of 5.1 µm.

(Some figures may appear in colour only in the online journal)

1. Introduction

Infrared photodetectors based on type-II superlattice (T2SL) InAs/GaSb have recently attracted considerable interest due to the ability to engineer the band gap [1], suppression of Auger recombination [2] and low interband tunnelling [3]. Such advantages make type-II super lattice photodetector technology a promising alternative to mercury cadmium telluride (MCT) and quantum well infrared photodetectors (QWIPs) in focal plane array (FPA) applications. However, as the confinement of photocurrent typically requires the fabrication of a mesa structure, surface states due to dangling bonds, inversion layers and interfacial traps are generated due to the abrupt termination of the crystal structure on the mesa side walls, which mediate surface leakage currents. In order to overcome this, various passivation methods such as sulfide passivation [4], deposition of a dielectric layer [5,6], overgrowth with wide band gap materials [7], polymer film coatings, such as polyimide [8], have been proposed. Passivation is expected to suppress the oxidation of the side walls and saturate dangling bonds to prevent the formation of surface states. Silicon dioxide deposition has been shown to be an effective technique [9], but it requires high temperatures for deposition or high RF powers to excite a plasma with potential for damage. Sulfur passivation replaces oxygen with sulfur and saturates the dangling bonds [10]. It is an effective passivation method and is easy to apply but the effect of passivation is temporary making it impractical [11]. It is usually applied using ammonium sulfide, sodium sulfide or zinc sulfide solutions. To increase the long-term stability of sulfur passivation, post-dielectric capping was used [12]. Alternatively, an electrochemical sulfur passivation (ECP) treatment was tried [4], but ECP does not prevent degradation with time, completely. Also, sulfur solutions etch GaSb layers (120 nm min⁻¹) at 60 °C [13] resulting in porous layers damaging the superlattice structure. For InAs/GaSb SLs, with mesas etched down to the GaSb buffer layer, sulfur may etch the GaSb buffer introducing a series resistance, which may lead to a superficial increase in the dynamic resistance, $R₀$ [10]. Sulfur reduces the reverse bias dark current density by two orders of magnitude and increases the zero bias resistance ($R₀$) by two orders of magnitude [4,5] while reducing photoresponsivity [10], which is masked by an increase in detectivity ($D^*$) due to higher dynamic resistance ($R₀$).

In this work, we propose to use sulfur-containing self-assembled monolayer (SAM) octadecanethiol (ODT),
CH$_3$(CH$_2$)$_{17}$SH, as a passivation layer for the InAs/GaSb superlattice photodetectors. ODT SAMs provide a reactive sulfur head (SH) at the end of the 18-carbon long chain. Although thiol SAMs are mostly studied on Au surfaces, some studies on III–V materials have also been reported [14–21]. Majority of the work on thiol SAMs on III–V compounds investigate GaAs [14–17] but there are a few examples on InP [18], GaP [19] and InAs [20, 21]. Typically, the active sulfur end of the thiol creates sulfur bonds on the surface, protecting the surface from oxidation while satisfying dangling bonds. ODTs replace oxygen atoms by sulfur atoms and thereby clean the surface from conductive native oxides [20–22]. Using x-ray photoelectron spectroscopy (XPS) McGuiness et al showed that long chain ODT and dodecanethiols form a monolayer (ML) on etched GaAs surfaces, which strongly reduces As and Ga oxides and effectively prevents subsequent interface oxide re-growth [17]. Knoben et al showed that well-ordered SAMs of ODT can form on InAs and reduce surface oxidation [20, 21]. Work on InP photodiodes has demonstrated suppression of dark current due to ODT SAMs [18]. Conductive oxides on the mesa side walls can reduce device performance, dramatically. SAM thiols bring all the good benefits of sulfur passivation without any side effect. Thiol treatment can be applied by just dipping the sample in a thiol solution. Unlike sulfur passivation, self-terminating thiol does not damage III–V surfaces [21] and the large chain length of the ODT SAM increases the thermodynamic stability [23] where intermolecular forces lead to the formation of a skin-like hydrocarbon layer.

2. Experimental

The sample studied in this work was grown commercially (IQE Inc. USA) with molecular beam epitaxy on a GaSb substrate. The photodetector was designed as a p-i-n photodetector with a design cutoff wavelength of 5 µm. It starts with a 100 nm thick GaSb buffer layer and a 20 nm Al$_{1-x}$Ga$_x$As$_{0.5}$Sb layer as an insulator and etch stop layer, respectively, followed by a 1000 nm GaSb:Be ($p = 1.0 \times 10^{17} \text{ cm}^{-3}$) p contact layer. The p-i-n part of the design consists of 140-period 8 ML of InAs/8 ML of GaSb:Be ($p = 1.5 \times 10^{17} \text{ cm}^{-3}$) p-type layers, 40-period 8 ML of InAs/8 ML of GaSb i-layers, 60-period 8 ML of InAs:Te ($n = 5 \times 10^{17} \text{ cm}^{-3}$)/8 ML of GaSb n-type layers, and the structure is terminated by a 20 nm InAs:Te ($n = 5 \times 10^{17} \text{ cm}^{-3}$) cap layer. Single-pixel photodetectors were fabricated with mesa sizes of 400–600 µm$^2$. To minimize surface damage, mesas were fabricated by standard lithography and wet etching. Mesa-isolated photodiodes were defined with H$_3$PO$_4$/C$_6$H$_8$O$_7$/H$_2$O$_2$/H$_2$O. Top and bottom ohmic contacts were made by evaporating 5 nm titanium and 200 nm gold. One was bonded to a chip carrier as reference and the other used for ODT passivation. The sample was immersed in a solution of HCl in isopropanol (3M) for 1 min and rinsed in isopropanol for 5 min prior to immersion in 1mM solution of ODT (Aldrich, 99%) in ethanol and was left there for 48 h at 60 °C. Then the sample was rinsed with ethanol and isopropanol and dried with dry N$_2$.

3. Results and discussion

Figure 1(a) shows a schematic of the InAs/GaSb SL p-i-n photodiode structure and figure 1(b) shows a schematic of ODT on the superlattice surface. ODT molecules form a smooth, continuous, crystalline packed hydrocarbon layer with a tilt angle of ∼30° from the surface normal [20]. Figure 1(c) shows the scanning electron micrograph of a 400 µm$^2$ photodetector. The SEM image clearly shows that the ODT SAM solution does not damage the superlattice layer or the underlying GaSb layer.

To investigate the electrical response of ODT SAM passivation, we compared dark currents of passivated samples with unpassivated devices at 77 K. The diodes were placed in a metal box and immersed into liquid nitrogen. The I–V characteristics were measured using a HP4142OA source-measure unit. Figure 2(a) shows the measured dark current density versus applied bias voltage characteristics of the
Figure 2. (a) Dark current density versus applied bias of the unpassivated and ODT SAM-passivated $400 \times 400 \mu m^2$ diodes at 77 K. (b) Zero bias differential resistance versus applied bias voltage characteristics for the unpassivated and ODT SAM-passivated samples at 77 K.

Figure 3. Ageing of the ODT SAM-passivated InAs/GaSb p-i-n photodetector at 77 K up to 65 days. (a) Dark current versus applied bias, (b) dynamic resistance–area product versus applied bias. The ODT SAM-passivated devices do not change with time for the duration of our experiment.

unpassivated and ODT SAM-passivated $400 \mu m^2$ diodes at 77 K. The passivated detectors show at least two orders of magnitude reduction in dark current density compared with the unpassivated detectors. The dark current density (at $-0.1$ V) reduced from $1.1 \times 10^{-5}$ to $3.1 \times 10^{-7} A cm^{-2}$. These measurements yield $R_0 A$ product values of $1.8 \times 10^4 \Omega cm^2$ and $3.2 \times 10^5 \Omega cm^2$ for the unpassivated and ODT SAM-passivated samples, respectively (figure 2(b)). A comparison of dark currents clearly shows that ODT SAM is a very effective way for passivation of type-II InAs/GaSb p-i-n photodetectors. The prominent reduction in dark current due to ODT SAM passivation is very encouraging when compared with the recently published type-II InAs/GaSb passivation works [11, 24].

For successful passivation, another important parameter, in addition to low dark current and high dynamic resistance values, is the long-term stability. Stability measurements were performed at 77 K. The samples were mounted on a helium-cooled cold finger and covered with a Au-coated radiation shield. $I$–$V$ measurements were repeated every 10 days to monitor the effects of ageing of the passivated devices. After each measurement, the cryostat was opened to air to provide atmospheric ageing conditions. Figure 3 shows the measured dark current density versus applied bias voltage and dynamic resistance–area product versus applied bias voltage characteristics of the ODT SAM-passivated $400 \mu m^2$ diodes at 77 K with at least a ten-day interval between the measurements. Dark current measurements indicate that the ODT SAM does not show any sign of ageing even after 65 days. Strong intermolecular interactions between the long chains of ODT molecules form a closely packed and thermodynamically stable skin-like layer. This reinforced sulfurization method is expected to form an inert surface. Small fluctuations are due to the noise floor of our instrument.

To understand the nature of the dark current, temperature-dependent $I$–$V$ measurements were made. Dark current densities versus inverse temperatures under $-0.1$ V bias are shown in figure 4. The $I$–$V$ curve is dominated by a
The diffusion current for temperatures higher than 100 K and by a generation–recombination (G–R) current for temperatures lower than 100 K. The diodes with ODT SAM passivation show a lower dark current than the unpassivated photodetectors for temperatures lower than 100 K. This suggests that the passivation process reduces native oxides eliminating surface states and preventing current flow through the surface channel. Figure 4 also shows the summary of dark current density at $-0.1$ V as a function of temperature measured at different stages of ageing. The temperature dependence of ageing over 60 days is consistent with our prior measurements. For temperatures higher than 100 K, Arrhenius-type behaviour yielded an activation energy of 0.23 eV, which is close to the device band gap. For lower temperatures, current begins to divert from the Arrhenius type of behaviour. The G–R current becomes dominant for temperatures below 100 K. The passivated photodiodes show reduced G–R currents, which may originate both from the bulk and the surface. Our results indicate that passivation has led to a reduction in the surface contribution of the G–R current. Alternatively, to analyse the surface-related dark current behaviour, the dynamic resistance is measured for different mesa sizes. The inverse of the dynamic resistance–area product at zero bias as a function of the perimeter-to-area ratio at 77 K for the passivated and unpassivated detectors is shown in the inset of figure 4. The surface resistance can be calculated from the slope of the graph [24]. The calculated surface resistivity ($\rho_{\text{surface}}$) values at zero bias are $1.2 \times 10^5 \Omega \text{cm}$ and $3.9 \times 10^6 \Omega \text{cm}$ for the unpassivated and passivated diodes, respectively. The results reveal an improvement of at least a factor of 30 in $\rho_{\text{surface}}$ for the ODT SAM-passivated devices at 77 K.

The spectral response of the photodetectors was measured with a Fourier transform infrared spectrometer (FTIR, Bruker Equinox 55). The photoresponsivity of the photodetectors was measured at 77 K using a calibrated blackbody source at 450 °C (Newport, Oriel 67000), a lock-in amplifier (SRS, SR830 DSP) and a mechanical chopper (SRS, SR540) system. The photodetectors were illuminated with a 300 K background with a 2π field of view. A 3–5 μm blackbody filter was used to eliminate unwanted illumination. Figure 5(a) shows the quantum efficiency (QE) versus wavelength graph under 0 V bias voltage at 77 K. The QE of the passivated photodetector is determined as high as 37% for these single pass detectors under front illumination around 3.5 μm wavelength. Since the QE is directly related to the thickness of the absorbing layer, it is difficult to make direct comparisons with other published data due to differences between structures. In any case, 37% QE in our photodetectors is high among type-II InAs/GaSb superlattice papers [25, 26]. Figure 5(b) shows the responsivity and Johnson-noise-limited detectivity ($D^*$) versus applied bias voltage graph for the ODT SAM-passivated photodetector. The zero bias responsivity of the ODT-passivated photodetector was equal to 1.04 A W$^{-1}$ at 4 μm and 77 K. Under zero bias, the peak $D^*$ was found to be $2.15 \times 10^{13}$ Jones for the passivated photodetector at 4 μm and 77 K. When we compare our results with recently published type-II InAs/GaSb passivation works
[11, 24], the ODT-passivated T2SL photodetectors are very promising.

The influence of the ODT SAM on the performance of the T2SL is closely related to the surface chemistry of the superlattice. Chemically reactive InAs/GaSb SL surfaces are easily oxidized and a native oxide layer of several nanometres thick is quickly formed upon exposure to air. These form conductive channels and lead to surface leakage. However, sulfur is chemically preferred over oxygen, and replaces thick is quickly formed upon exposure to air. These form easily oxidized and a native oxide layer of several nanometres superlattice. Chemically reactive InAs/GaSb SL surfaces are promising.

4. Conclusions

We have demonstrated the suppression of dark current and increase in optical response of InAs/GaSb superlattice photodetectors with cutoff wavelength at 5.1 μm (MWIR). We have used ODT SAM passivation of InAs/GaSb p-i-n superlattice photodetectors. The passivated superlattice photodetectors reduced the dark current from 1 nA to 3 × 10−10 A cm−2 compared with the unpassivated photodetector at 77 K and under −0.1 V applied bias condition. The corresponding zero bias–area product (R0A) is improved by two orders of magnitude (from 1.8 × 104 to 3.2 × 106 Ω cm2). The zero bias sensitivity and detectivity (D*) are determined to be 1.04 A W−1 and 2.15 × 1013 Jones, respectively at 4 μm and 77 K. The quantum efficiency (QE) of the passivated photodetector is determined to be 37% under the single pass front illumination condition. Temperature-dependent dark current and dynamic resistance–area product as a function of the perimeter-to-area ratio measurements revealed that passivation prevents photodetectors from being dominated by surface currents for temperatures below 100 K. The devices show Arrhenius type of behaviour at higher temperatures, an indication that the dominant current is the bulk diffusion current for high temperatures. The calculated activation energy is equal to 0.23 eV, which is close to the device band gap. The dark current performance of the photodiodes did not degrade for 65 days after the passivation, indicating long-term stability. This work shows that ODT SAM is a good passivation material for p-i-n InAs/GaSb superlattice photodetectors.

References