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## Bonding Geometries at the In<sub>2</sub>O and SiO/III-V Semiconductor Interface

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Oxide monolayers and submonolayers formed by vapor depositon of In<sub>2</sub>O and SiO oxides on InAs(001)-(4×2) were studied by scanning tunneling microscopy (STM). At low coverage, In<sub>2</sub>O molecules bond to the edges of the rows and most likely form new In-As bonds to the surface without any disruption of the clean surface structure. Annealing the  $In_2O/InAs(001)-(4\times2)$  surface to 380°C results in formation of flat ordered monolayer rectangular islands. The annealed In<sub>2</sub>O no longer bonds with just the As atoms at the edge of row but also forms new O-In bonds in the trough. SiO chemisorption on InAs(001)- $(4\times2)$  is completely different than In<sub>2</sub>O chemisorption. At room temperature, even at low coverage SiO adsorbates bond to themselves and form nanoclusters. For SiO/InA(001)-(4×2) post-deposition annealing (PDA) does not disperse the nanoclusters into flat islands. Both In<sub>2</sub>O and SiO depositions on InAs(001)-(4×2) surface do not displace surface atoms during both room temperature deposition and postdeposition annealing.

#### Introduction

The current silicon based integrated circuits (IC) technologies have dominated the high-performance logic industry for decades. The success with the semiconductor IC industries lies in device downsizing: scaling. However, continuing the scaling is becoming increasingly difficult, and silicon transistor technology is rapidly approaching its physical limits. According to the 2008 updated International Technology Roadmap for Semiconductors (ITRS) (1), to attain adequate drive current for highly scaled metal-oxide semiconductor field-effect transistors (MOSFETs), eventually other high transport channel materials (Ge and III-V) are needed for beyond 22 nm node generation so that semiconductor industries can continue keeping Moore's Law.

Recently, extensive studies have been carried out on III-V compound semiconductors for potential channel materials of MOSFETs (2-4). Several academic and industry groups have shown promising III-V MOSFETs results (5-12). However, the interface between oxides and III-V semiconductors is not well understood. Forming electrically passive interfaces between the gate oxide and the channel (13) is still a major issue for the development of MOSFETs devices. Unlike hydrogen passivated silicon and SiO<sub>2</sub> interfaces ( $D_{it} \sim 10^{10}~eV^{-1}cm^{-2}$ ) (14), oxide/III-V interfaces have large interfacial trap densities,  $D_{it}$  between  $10^{12}$  and  $10^{14}~eV^{-1}cm^{-2}$  (15,16), so enhancement mode III-V

MOSFETs devices require very high capacitance even to counteract the lowest reported the interfacial trap densities. In order to obtain a fundamental understanding of surface passivation of III-V compound semiconductors, it is critical to investigate and understand surface morphological evolution and the oxide/semiconductor interface bonding at the atomic level in order to facilitate gate oxide selection for III-V MOSFETs device and enable very aggressive oxide scaling.

InAs is an attractive channel material for high speed, low power MOSFET due to its very high effective electron mobility ( $\sim 15,000~\rm cm^2~V^{-1}s^{-1}$ ) compared to Si ( $\sim 300~\rm cm^2~V^{-1}s^{-1}$ ) at 300 K (17). InAs FET results have shown there is potential for high performance InAs based MOSFET (18,19). This report is focused upon the  $c(8\times2)/(4\times2)$  surface since other surfaces like As-rich (2×4) reconstruction readily undergo oxygen induced displacement reactions (20-23). Although InAs(001) (24-26) have been the focus of many scanning tunneling microscopy (STM) studies, only one STM study has been performed on the bonding geometry of oxide on InAs(001) surface (27).

To develop an atomic understanding of oxide morphologies on InAs(001)- $c(8\times2)/(4\times2)$ , the bonding geometries of two different oxides  $In_2O$  and SiO on InAs(001)- $c(8\times2)/(4\times2)$  were investigated. Scanning tunneling microscopy (STM) was used to determine the exact interfacial bonding geometries of  $In_2O$  and SiO/InAs(001)- $c(8\times2)/(4\times2)$  after room temperature oxide deposition. In addition, the effects of post-deposition annealing on oxide bonding sites were investigated. The results are focused on InAs, but the results are readily applied to InGaAs systems due to similar surface structure and oxide bonding geometry (28).

## **Experimental and Computational Method**

MBE was employed to grow 0.2  $\mu$ m of 1×1018 cm-3 doped InAs layer on 500  $\mu$ m thick InAs(001) substrates with 1×10<sup>18</sup> cm<sup>-3</sup> doping. Experiments were performed on both n-type and p-type wafers. The re-grown wafers were capped in situ with a 50 nm protective arsenic cap. The wafers were transferred to a vacuum container for transporting to the STM chamber. Experiments were performed in two ultrahigh vacuum (UHV) chamber systems with base pressures of 2×10<sup>-10</sup> Torr. Each chamber was equipped with low energy electron diffraction (LEED) and either a Park Scientific room temperature or an Omicron variable temperature (VT) STM. The InAs(001)- $c(8\times2)/(4\times2)$  surface was prepared by thermal decapping method. Details concerning the decapping method used in this study have been discussed elsewhere (28,29).

After the  $c(8\times2)/(4\times2)$  surface reconstruction was verified by LEED, the sample was transferred to the STM chamber, where STM measurements were performed at room temperature. All of the STM measurements were carried out at room temperature (RT) using electrochemically etched tungsten tips. The InAs(001)- $c(8\times2)/(4\times2)$  clean surfaces could be observed with both filled and empty state STM. Subsequently, In<sub>2</sub>O or SiO was deposited in the main chamber by evaporating sintered In<sub>2</sub>O<sub>3</sub>(s) (Aldrich Chem. Co.) or SiO(s) (Alfa Aesar) from a high temperature effusion cell (Applied EPI) contained in a differentially pumped chamber. In<sub>2</sub>O<sub>3</sub>(s) sublimates as In<sub>2</sub>O (g) and O<sub>2</sub> (g) at 1000 - 1025 °C (30). However due to the low sticking probability of O<sub>2</sub> on group III-rich  $c(8\times2)/(4\times2)$  surfaces (31), the majority of the adsorbed species are In<sub>2</sub>O. SiO(s)

evaporates congruently as SiO(g) molecules at 950 - 975 °C (29). During evaporation, the main chamber pressure is  $2 \times 10^{-7}$  Torr or  $5 \times 10^{-9}$  Torr for In<sub>2</sub>O and SiO, respectively.

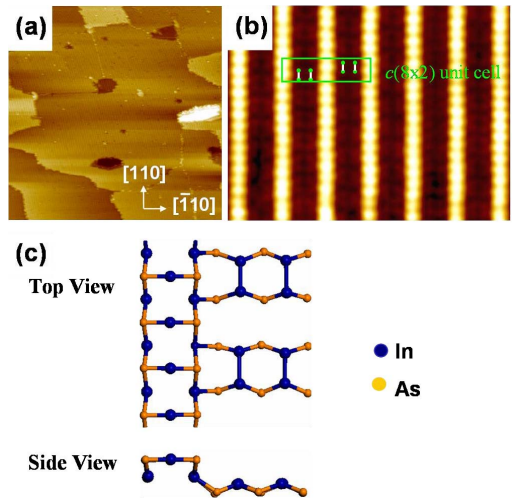


Figure 1. Structure of clean InAs(001)-(4×2) at 300 K. (a) A filled state (Vs = -2 V, It = 0.2 nA, 2000 Å × 2000 Å) STM image obtained from the InAs(001)-(4×2) clean surface. (b) A high resolution filled state (Vs =-2 V, It = 0.2 nA, 100 Å × 85 Å) STM image obtained from the InAs(001)-(4×2) clean surface. (c) Ball-and-stick diagram of the InAs(001)-(4×2) reconstruction. The atomic InAs(001)-(4×2) reconstruction structure model has been discussed in Ref. 22. The green rectangle presents the unit cell of  $c(8\times2)$ . Note that In atoms on the row are undimerized, and the In row atoms have the same height as the As atoms at the edge of row.

### **Results and Discussions**

In-rich InAs(001)-(4×2) can readily be prepared by decapping of As<sub>2</sub> capped InAs(001) wafers. A typical STM image of a clean InAs(001)- $c(8\times2)/(4\times2)$  surface prepared by decapping method can been seen in Fig. 1(a). All STM images in this study were taken with a negative sample bias using the constant-current tunneling mode. The InAs(001)-(4×2) surface has a unit cell consisting of two undimerized group III atoms on the row and two group III homodimers in the trough. Details of the InAs(001)-(4×2) clean surface structure are discussed elsewhere (26).

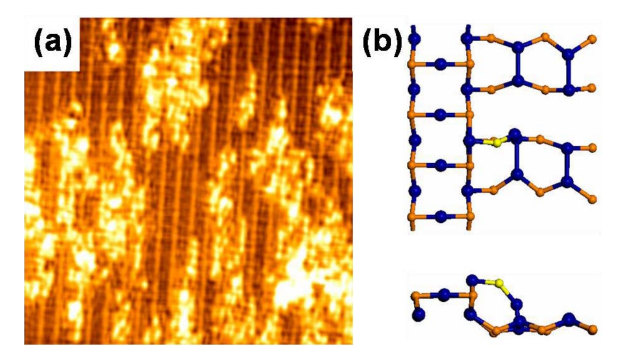


Figure 2. Bonding of RT deposited  $In_2O/InAs(001)$ -(4×2). (a) A filled state (Vs = 1.5 V, It = 0.2 nA, 35 Å × 35 Å) STM image obtained from the InAs(001)-(4×2) surface with  $\sim 25\%$  coverage of  $In_2O$ . (b) Possible atomic bonding structure diagrams of  $In_2O/InAs(001)$ -(4×2) surface along single  $In_2O$  chemisorption bonding sites with top view (upper) and side view (lower) . Note that this image is taken in the Park Scientific RT-STM system.

# <u>Indium Monoxide (In<sub>2</sub>O)</u>

Once the surface structure of InAs(001)-(4×2) was characterized, the gate oxides were deposited to investigate the exact atomic bonding structures of gate-oxide/III-V interfaces. STM results showed that the initial deposited In<sub>2</sub>O molecules bond to the edges of the rows and most likely form new In-As bonds to the surface without any disruption of the clean surface structure, shown in Fig. 2(a). In filled state STM images, In<sub>2</sub>O is imaged as a bright feature on the clean InAs(001)-(4×2) surface. Figure 2(b) shows a structural model illustrating the possible oxide bonding structure for room temperature deposition. Density function theory (DFT) results shows the binding energy is 0.84 eV/In<sub>2</sub>O for InAs surface (28). The binding energy (32) was calculated by subtracting the total energy of the most stable clean surface structure and gas phase oxide molecule from the total energy of the oxide/InAs complex:  $E_{bind}^{oxide/InAs} = E^{oxide/InAs} - (E^{InAs} + E^{oxide})$ , where  $E_{bind}^{oxide/InAs}$  is the binding energy between oxide molecules and InAs surface,  $E^{oxide/InAs}$  is the total energy of oxide deposited InAs double unit cell slab,  $E^{InAs}$  is the total energy of the clean InAs double unit cell slab, and  $E^{oxide}$  is the total energy of free oxide molecules (28).

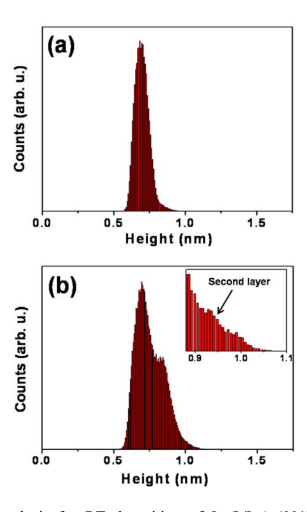


Figure 3. Height analysis for RT deposition of  $In_2O/InAs(001)-(4\times2)$ . (a) Height distribution of only the oxide-free regions of the  $In_2O/InAs(001)-(4\times2)$  surface in Fig. 2(a). (b) Height distributions of the entire  $In_2O/InAs(001)-(4\times2)$  surface in Fig. 2(a). There are two position peaks corresponding clean surface region and oxide covered surface region. The distance between two peaks is 1.5 Å. Inset in Fig. 3a is an expanded height distribution of  $In_2O$  deposited surface at the higher height region that reveals distinct second layer growth.

A height distribution analysis was performed to determine the room temperature growth mechanism. At medium coverage (  $\sim 25\%$  monolayer), the In<sub>2</sub>O formed islands that are in the [110] direction as shown in Fig. 2(a). Prior to all the first layer bonding sites being occupied with oxides, distinct second layer growth is observed on the oxide islands. Figure 3 shows In<sub>2</sub>O height distribution for room temperature oxide deposition. For only clean surface region, there is height distribution peak near 7.0 Å, shown in Fig. 2(a). For the whole surface, there is a bimodal height distribution shown in Fig. 3(b) with a lower position peak which is identical to the clean region height distribution of the InAs(001)-c(8×2)/(4×2) surface. The higher position peak corresponds to the oxide covered surface. The 1.5 Å height difference between the oxide and clean surface is consistent with single monolayer growth consistent with the line scan analysis performed on the Fig. 2 STM image. Inset in Fig. 3(b) is expanded height distribution of In<sub>2</sub>O deposited surface at the higher height region that consistent with second layer oxide

growth on the oxide islands. The first and second layers of oxide have the distinct heights corresponding to distinct layers of oxide. However, the second and subsequent oxide layers appear to be amorphous since these oxides do not form ordered structures.

To form more ordered oxide structures, the effect of annealing In<sub>2</sub>O/InAs(001) $c(8\times2)/(4\times2)$  was studied. Post-deposition annealing induced the oxide to spread out. As shown in Fig. 4, the 380°C annealed samples were more ordered with respect to both island shape and the structure within the islands consistent with the lowest energy structure being an ordered first monolayer of oxides. The ordered structures contained oxide in the troughs with rows in the [110] direction. Comparisons of oxide height distributions before and after annealing show the oxide height is reduced from 1.5 Å to 1.1 Å, shown in Fig. 5(a). For the oxide covered regions of the surface, there is a Gaussian height distribution shown in Fig. 5(b) with a full width at half max of only 1.8 Å which is consistent with formation of flat monolayer island structures with the 1.8 Å corrugation being primarily due to the apparent STM height difference between the indium and oxygen atoms in In<sub>2</sub>O adsorbates. Most importantly, for both room temperature deposition and post-deposition annealing, the In<sub>2</sub>O adsorbates never cause the displacements of any surface atoms on the  $InAs(001)-c(8\times2)/(4\times2)$  surface; if the annealing temperature is raised above 450°C using cyclic annealing (29) then the In<sub>2</sub>O molecules completely desorb leaving behind the clean (4×2) surface (28). The 380°C annealing induced formation of dense flat square mono-atomically tall islands indicating that a new bonding geometry forms after post-deposition annealing. DFT simulations show that the annealed In<sub>2</sub>O no longer bonds with just the As atoms at the edge of row but also forms new O-In bonds in the trough, shown in Fig. 4(b). For In<sub>2</sub>O, In<sub>2</sub>O-In trough sites are more stable than In-O-In-As edge sites by about ~0.77 eV/In<sub>2</sub>O (28).

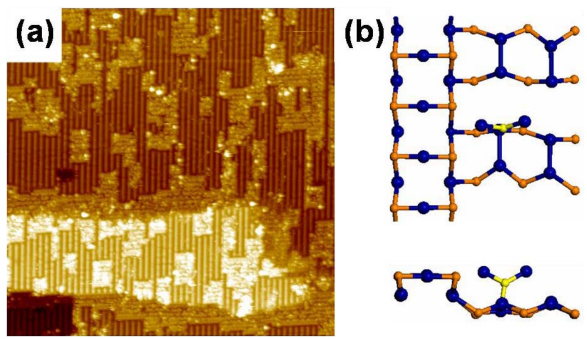


Figure 4. Bonding of RT deposited  $In_2O/InAs(001)$ -(4×2) after a 380°C anneal. (a) A filled state (Vs = -2.5 V, It = 0.2 nA, 1000 Å × 1000 Å) STM image obtained from the InAs(001)-(4×2) surface. (b) Possible atomic bonding structure diagrams of annealed  $In_2O/InAs(001)$ -(4×2) surface along single  $In_2O$  chemisorption bonding sites with top view (upper) and side view (lower).

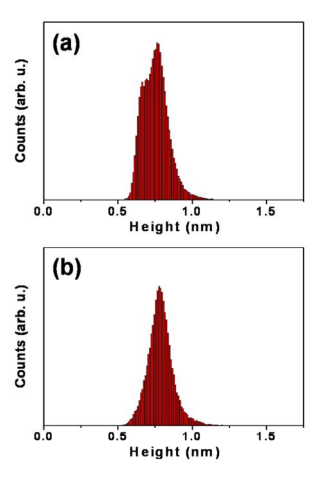


Figure 5. Height analysis for RT deposition and 380°C annealing of In<sub>2</sub>O/InAs(001)-(4×2). (a) Height distributions of In<sub>2</sub>O deposited and annealed InAs(001)-(4×2) in Fig. 4(a). (b) Height distribution of only the oxide covered regions of In<sub>2</sub>O deposited and annealed InAs(001)-(4×2) in Fig. 4(a) There are two position peaks corresponding clean surface region and oxide covered surface region, respectively. The distance between two peaks is 1.1 Å.

## Silicon Monoxide (SiO)

Figure 6 shows a STM image of the InAs(001)-(4×2) surface after low coverage SiO deposition (~ 4% ML) at 300K with post-deposition annealing (PDA). The bright features on the surface are SiO molecules bonding with the InAs(001)-(4×2) substrate; some of the SiO bond to the rows and some of the SiO bond in the troughs. Magnified STM images show that there are three distinct types of bright features presented on the SiO/InAs(001)-(4×2) surface, which have been denoted by a hexagon, a circle and a square. These bright sites can be clearly differentiated by line scan analysis. The results show that these bright sites represent heights of three different oxide layers. The hexagon [Fig. 6(b)] is found to image as 1.8 Å above the surface row. This height represent monolayer height since it is close to the height of a single SiO molecule on GaAs(001)-(2×4) surface (29). The circle [Fig. 6(c)] is found to image as 3.8 Å above the surface row, indicating two layer SiO molecules on surface. The square [Fig. 6(d)] is found to image as 5.7 Å above the surface row, indicating three layer SiO molecules on surface.

These bright features are consistent with the higher self-binding energy of SiO molecules to form multilayer structures on InAs(001)-(4×2) (33).

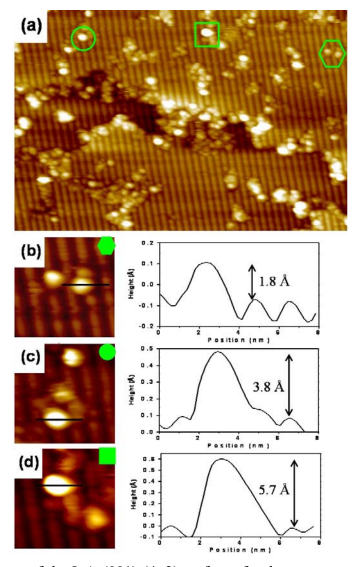


Figure 6. STM image of the InAs(001)-(4×2) surface after low coverage SiO deposition ( $\sim 4\%$  ML) at 300 K with PDA. (a) A filled state (Vs = -2.5 V, It = 0.2 nA, 80 Å × 55 Å) STM image obtained from SiO deposited InAs(001)-(4×2) surface. Three magnified STM images illustrate three different heights with corresponding geometric symbols, and line scan corresponding to the black line on the STM image revealing that SiO adsorbates bond themselves and form nanoclusters: (b) hexagon: 1.8 Å height with 12 Å × 12 Å size, (c) circle: 3.8 Å height with 16 Å × 12 Å size, and (d) square: 5.7 Å height with 20 Å × 16 Å size. Note that a single SiO molecule's size is about 4 Å × 4 Å according to Ref. 29.

At 35% coverage of SiO deposited on the InAs(001)-(4×2) surface at room temperature, there is no ordered oxide structure formation. SiO molecules bond to themselves and form nanoclusters shown in Fig. 7(a). The bonding configuration is nearly identical when the SiO/InAs sample was annealed to 380°C [Fig. 7(b)]. Height distribution analyses were performed on SiO deposited InAs(001)-(4×2) surface without and with PDA at 380°C. As shown in Fig. 7(c), for the whole surface without PDA, SiO deposited InAs(001)-(4×2) has a broad height distribution. For SiO/InAs(001)-(4×2)

shown in Fig 7(c), there is a lower position peak ( $\sim 7$  Å height) which is the clean region of the InAs(001)-(4×2) surface. If only the oxide covered surface is included in the height distribution analysis, the results [Fig. 7(e)] show the SiO/InAs(001)-(4×2) surface has much broader oxide height distribution (full width at half max (FWHM) ~ 7.4 Å) which is 3.7 × as large as the height distribution for the oxide covered regions of  $In_2O/InAs(001)-(4\times2)$  (FWHM ~ 2 Å) shown in Fig. 3. After PDA at 380°C, the height distribution becomes smoother consistent with the surface becoming slightly flatter. However, after PDA the oxide covered regions of SiO/InAs(001)-(4×2) have a FWHM of 5.9 Å which is still 3 × FWHM of the oxide covered regions for post-deposition annealed  $In_2O/InAs(001)$ -(4×2) (FWHM ~ 1.8 Å). This height distribution analysis is consistent with formation of nanoclusters for SiO deposited InAs surface. Further analysis of the SiO/InAs(001)-(4×2) surface after PDA shows no bright displaced As atoms nor dark oxygen sites. The common feature for both  $In_2O$  and SiO depositions on InAs(001)-(4×2) surface is that oxides do not displace surface atoms during both room temperature deposition and post-deposition annealing. After oxide deposition, InAs(001) surface retain the  $(4\times2)$  reconstruction even within a few angstroms of the adsorbates.

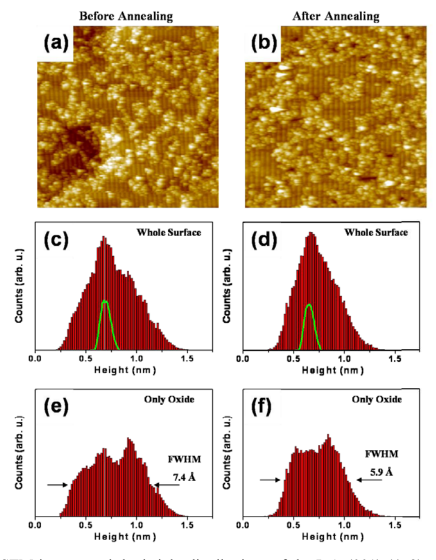


Figure 7. STM images and the height distributions of the InAs(001)-(4×2) surface after 35% coverage of SiO deposition. Filled state (Vs = -2.5 V, It = 0.2 nA, 700 Å × 700 Å)

STM images obtained from SiO deposited InAs(001)-(4×2) surface (a) without and (b) with post-deposition annealing (PDA) at  $380^{\circ}$ C. (c) Height distribution of the entire Si<sub>2</sub>O/InAs(001)-(4×2) surface without PDA shown in Fig. 7(a). (d) Height distribution of the entire Si<sub>2</sub>O/InAs(001)-(4×2) surface with PDA shown in Fig. 7(b). (e) Height distribution of the oxide covered regions of SiO/InAs(001)-(4×2) without PDA. (e) Height distribution of the oxide covered regions of SiO/InAs(001)-(4×2) with PDA. The green curves shown in Fig. 7 whole surface are the clean surface regions of SiO/InAs(001)-(4×2) surface.

The experimental results show that  $In_2O$  molecules completely desorb at 450 °C leaving behind the clean (4×2) surface [28], and SiO molecules remain on the surface even at 530 °C (not shown). The temperatures above 530 °C were not tried due to the heating limitations of the manipulator. Literature results show for pure substances (not InAs chemisorbates) SiO has higher sublimation energy than  $In_2O$  ( $\Delta H$ : SiO 3.3 eV vs.  $In_2O$  2.6 eV) (33). It is hypothesized that the lower self-binding energy allows  $In_2O$  to form monolayer flat ordered islands, while the higher self-binding energy induces SiO to form multilayer disorder amorphous structures on InAs(001)-(4×2). Figure 8 shows the vapor pressure for  $In_2O$  and SiO versus the temperature according to experimental data of Ref. (33-35) To desorb 1 ML of  $In_2O$  and SiO oxide from themselves (for example:  $10^{-8}$  Torr vapor pressure for 100 seconds), the oxide temperature should be 440 °C for  $In_2O$  and 595 °C, which is consistent with experimental results.

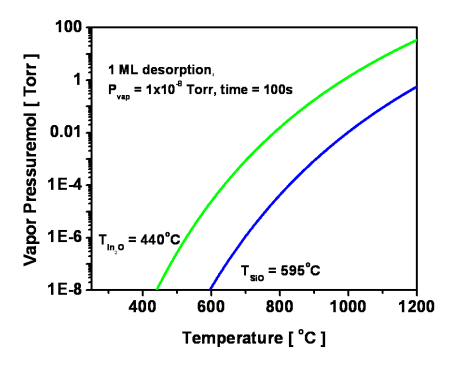


Figure 8. The vapor pressure vs temperature for In<sub>2</sub>O and SiO.

# **Summary**

The bonding geometries of two different oxides  $In_2O$  and SiO on InAs(001)- $c(8\times2)/(4\times2)$  were investigated using STM. The results show the completely different bonding geometries.  $In_2O$  molecules initially bond with As atoms at the edge of the row. After post-deposition annealing,  $In_2O$  molecules bond with the trough In atoms to form new O-In bonding sites.  $In_2O$  oxide form dense flat monolayer tall island structures on InAs(001)- $c(8\times2)/(4\times2)$  surface. Conversely, SiO molecules have higher self-binding energy so that they bond themselves and form nanoclusters on InAs(001)- $c(8\times2)/(4\times2)$  surface.  $In_2O$  and SiO depositions on InAs(001)- $c(8\times2)/(4\times2)$  surface do not displace surface atoms during both room temperature deposition and post-deposition annealing.

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