Use of Raman Spectroscopy in Characterizing Soft X-ray Multilayers: Tools in Understanding Structure and Interfaces

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Use of Raman spectroscopy in characterizing soft x-ray multilayers: 
Tools in understanding structure and interfaces

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ABSTRACT

Our group is studying the structure and interfaces of soft x-ray multilayers by various techniques including
x-ray diffraction and Raman spectroscopy. Raman spectroscopy is particularly useful since it is sensitive to the
identity of individual bonds and thus can potentially characterize the abruptness of interfaces in multilayers.
Blocking interfacial mixing is very important in achieving and maintaining high reflectivity. We report our studies
of the as-deposited and postannealed structure of Mo/Si and W/C multilayers. The Mo/Si system is probably the
most widely studied multilayer currently because of its potential applications for soft x-ray projection lithography
for the range of 13 to 15 nm. A high normal-incidence, peak reflectance is mandatory for imaging applications that
involve many reflections. The reported theoretical and achieved reflectances of the Mo/Si system are 80% and
65% respectively. This loss of 15% can bring about a six-fold loss in system throughput in the eight-reflection
system contemplated. The interfaces in the Mo/Si system are thought to play a significant role in the degrading
reflectance so characterization techniques which have interfacial sensitivity are particularly important. The Mo/Si
multilayer system is susceptible to Raman characterization since both the a-Si spacer layer and the MoSi 2 compound
which forms at the interface have Raman active modes. In this paper we report the first Raman studies, to our
knowledge, of the a-Si layers and their crystallization and the crystallization of the Mo/Si interface of the
multilayer brought about by a one-hour 1000 °C anneal. These changes are apparent in the Raman spectra before
they can be unambiguously detected by x-ray diffraction.

1. INTRODUCTION

Understanding the formation of refractory metal silicides at the interfaces of thin films is an extremely
interesting basic science and an important technological problem. Technologically, the formation of silicide is
important in very large-scale integration (VLSI) where silicides are being used as interconnect films and in
producing better soft x-ray multilayers, since their presence at interfaces in real Mo/Si multilayers keeps the peak
reflectance from achieving the theoretical maximum of 80% which is predicted for multilayers containing atomically
abrupt and smooth interfaces.

Besides their applications in x-ray imaging technology and microelectronics, multilayers have also proven
to be a convenient way of studying the nanoscale properties of materials - for example, the crystallization and
interdiffusion of matter on atomic scales.

In the current x-ray imaging revolution periodic bilayer multilayers have played an important role. These
multilayer mirrors, which are used to reflect light with the wavelengths in soft x rays, typically consist of periodic
multiple bilayers, each bilayer composed of a low-Z and high-Z layer. The reflectance is achieved by the
interference occurring at the sharp interfaces between these layers. Generally speaking, the sharper the interfaces,
the higher the reflectance which can be obtained. Any modification of the interfaces will directly impact the optical
performance of the multilayer. Techniques which can detect and quantify the intermixing at interfaces are extremely
important for preparing the superior multilayers required for soft x-ray projection lithography, since a 10% gain
in the reflectance in each of the mirrors of the eight-mirror systems contemplated for lithography can triple the light
through the camera.
Small angle x-ray diffraction (SAXRD) and direct imaging by scanning transmission electron microscopy (STEM) have been successful in characterizing interfaces of soft x-ray multilayers but can give artificially wide estimates of interfaces. Raman spectroscopy has also been shown to be a useful technique for characterizing multilayers of amorphous semiconductors, such as Si/Ge, and multilayers of metal/carbon, such as W/C multilayers because it can directly "count" interfacial bonds in some cases. Observing the interfaces of the multilayers is especially important, but under some circumstances this is very difficult and may require special care. This is because of the small volume of material within the interfaces and the interface bonds may have a low Raman cross section.

So far, Mo/Si is the most efficient multilayer mirror for the soft x-ray range of 13-30 nm. A large number of groups are studying these multilayers and their applications because of the high theoretical and achieved peak reflectances of 80% and 65%, respectively, over the 12.5 to 16 nm range. This is the preferred range for projection soft x-ray lithography. We are the first group to apply a Raman spectroscopic technique to characterize soft x-ray Mo/Si multilayers. We have observed the presence of a-Si in as-deposited multilayers, its disappearance and the formation of crystalline MoSi₂ in annealed multilayers. The Raman spectroscopy study of commercial MoSi₂ powder has also been carried out to help identify the MoSi₂ peaks. Both large angle x-ray diffraction (LAXRD) and SAXRD measurements have been performed to aid in understanding Raman results. Scanning electron microscopy (SEM) measurements provide direct picture of the surface topology evolution during annealing.

2. EXPERIMENTAL

The multilayer interfacial regions consist of a mixture of molybdenum and silicon, so that the study of molybdenum silicide is required for Raman studies of Mo/Si multilayers and we began with a study of molybdenum disilicide (MoSi₂) powder. For this reason, this study of multilayers has applications beyond the study of soft x-ray reflectors since silicides are expected to be materials for high-speed electronics and very large-scale integration (VLSI). This is because many refractory metal silicides have been found to have very high thermal and chemical stability and high electrical conductivity. Because of these applications, silicides have received substantial attention, and a large number of studies have been carried out on them. Many silicides such as WSi₂, PdSi, and TiSi₂ have been extensively studied by Raman spectroscopy, but there has only been one published report on MoSi₂.

The molybdenum disilicide powder was purchased from Aldrich Chemical Company, Inc. The powder's average particle size is about 1 µm and its manufacturer quotes the purity as 99%. The bulk material was obtained to provide a standard with which to compare the thin film material.

In addition to studying the as-received MoSi₂ powder and the as-deposited thin film samples, both powder and films were subjected to vacuum annealing, to air annealing, and the powder to KOH etching. This was done to aid understanding of the observed features in Raman and x-ray diffraction spectra. The vacuum annealing consisted of sealing the sample in an evacuated quartz tube or glass ampoule. For the sample annealed in air, the ampoule was not sealed. The furnace was controlled to 1000 °C and the sample was placed directly in the hot oven at the beginning of the annealing time and air quenched by withdrawing the ampoule directly from the furnace at the end of the annealing time. As an alternative to annealing and to etch out the reacted materials and impurities, some of the powder was mixed with KOH solution (10 M KOH) filtered and dried in air at a 110 °C oven. This etching process was observed to remove most of the residue reacted silicon.

The multilayer samples were made by dc magnetron sputtering for a fixed target system. Layering was achieved by rotating the substrate in the system at a fixed rate over the sources. The Mo/Si multilayers were prepared for soft x-ray mirror (plasma diagnostic) applications and were shown to have excellent soft x-ray reflectance. The sputtering targets were molybdenum metal and p-type silicon. The sample substrates were ultrasmooth crystalline silicon wafers of [100] orientation. The period λ or d-spacing was measured to be 8.4 nm, as deposited, with a nominal γ = dSil/dSi+2dMo of 0.40. There were 30 pairs. The total multilayer thickness is thus about 250 nm. The top layer is amorphous silicon and is twice as thick as other layers to retard oxidation. The multilayer samples were also sealed in vacuum and annealed at temperatures ranging from 300 to 1000 °C.
From now on we refer it as thick multilayer sample.

Another group of Mo/Si multilayer samples were prepared to be very thin on purpose to magnify the interface effects. We will call them thin samples from now on to distinguish from the normal ones described above. The samples substrates were ultrasmooth crystalline silicon wafers of [111] orientation. To use wafer with [111] orientation, we can separate the weak silicides' x-ray diffraction peaks from the strong substrate signals which are around 30°. The period \( \Lambda \) was measured to be 2.0 nm, as deposited, with a nominal \( \gamma \) of 0.46. There were 60 pairs. The top is still the silicon protection layer. These samples were also sealed in vacuum and annealed at temperatures of 300, 600, 900, and 1000 °C.

The Raman spectra were recorded on a SPEX 1877 Triplemate spectrometer. The spectrometer consists of a modified Czerny-Turner zero-dispersion double spectrometer as a filter stage and a modified Czerny-Turner spectrograph. The dispersion element used in the spectrograph stage is a 1800 grooves/mm holographic grating with 0.92 nm/mm dispersion. The spectral bandpass selected was \( -15 \) cm\(^{-1}\). The 514.5 nm line \( (-400 \) MW) of an argon ion laser (LEXEL 96) was used as the excitation source in most studies. The other laser lines were used on occasion to confirm peak identification. A 1200 grooves/mm ruled grating with a small aperture placed outside of the laser cavity was used as a presample monochromator to eliminate the laser plasma lines. The laser beam was brought to line focus \((1 \times 10 \) mm\(^2\)) on the sample's surface using a cylindrical lens to match the entrance slit of the triplemate monochromator; the scattered signals were analyzed by the spectrometer and detected by a photomultiplier (EMI 7981B). Then photon-counting electronics were used to convert the light signal into the electrical signal. A 386SX PC was used to control the spectral scanning and data acquisition through a digital-analog interface box (TransEra-MDAS-7000).

Both LAXRD and SAXRD measurements were performed before and after annealing on a SCINTAG DMS-2000 powder diffractometer using a CuK\( \alpha \) source \((\lambda = 0.154 \) nm) with power of 55.5V-25mA. SAXRD provides an accurate measurement of the average bilayer spacing of a multilayer mirror.\(^1\) A general rule that a large number of orders is indicative of sharp boundaries between the layers.

### 3. RESULTS AND DISCUSSION

The Raman spectra of the commercial molybdenum disilicide powder samples are shown in Figure 1. The raw powder (as-received) spectrum (see (a)) possesses four peaks: two small broad peaks located at 323 and 438 cm\(^{-1}\) (the higher frequency one is stronger), one fairly sharp peak located at 520 cm\(^{-1}\), and one small peak located at 822 cm\(^{-1}\). We conclude that the sharp peak at 520 cm\(^{-1}\) arises from the unreacted polycrystalline silicon in the powder. This is suggested by its position and was confirmed by two observations: first, the fact that the 520 cm\(^{-1}\) peak is lower after vacuum annealing at 1000 °C (see (b)) and, second, the fact that it is entirely absent after etching in KOH (see (c)). Strong alkali will etch crystalline silicon but will not etch MoSi\(_2\). Other investigators have also reported the presence of unreacted silicon in as-received silicide powders.\(^1\) We conclude that the high-frequency peak is due to MoO\(_3\) impurity. This was suspected because of the position of the line and was confirmed by annealing in air (see (d)), as will be discussed below. The two low-frequency peaks are due to MoSi\(_2\). Doland and Nemanich\(^2\) recently reported the position of MoSi\(_2\) prepared by annealing Mo on Si wafers as being 325 and 440 cm\(^{-1}\), which are close to our measurements of 323 and 438 cm\(^{-1}\). These peaks are also close to those of WSi\(_2\) at 331 and 451 cm\(^{-1}\), which has been studied more extensively.\(^3\) Both MoSi\(_2\) and WSi\(_2\) have tetragonal structure with space group 14/mmm \((D_{4d})\), with the Mo atoms at 2(a) Wyckoff positions and the Si atoms at 4(e). The vibrational mode analysis indicates two active Raman modes for MoSi\(_2\), \(\Gamma_{2g}^{1}\) (A\(_{1g}\)) and \(\Gamma_{2g}^{1}\) (E\(_{g}\)).

After the powder sample was annealed in vacuum at 1000 °C for one hour, we saw that the two MoSi\(_2\) peaks become sharper and stronger (see (b)). This can be interpreted as meaning that during the annealing process the original small grains of crystalline MoSi\(_2\) grew into larger grains which possess a larger Raman cross section. Analogous behavior, grain growth with concurrent sharpening and growth of the Raman signal, has been seen in the annealing of amorphous and microcrystalline Si. Note that even after annealing the high-frequency peak is still observable though smaller. We interpret this decrease in the silicon peak intensity as the partial reaction of some
of unreacted silicon during annealing.

To test the hypothesis that the 822 cm\(^{-1}\) peak is due to MoO\(_3\), we annealed the same sample again in air at 500 °C for one hour. The peak became more intense (see (d)). Note that in concert with the intensification many other weak peaks also appear. Their positions are 996, 665, 438, 374, 329, 286, 245, 218, 199, 159, and 129 cm\(^{-1}\). Because the purpose of this experiment was only to verify that the peak at 822 cm\(^{-1}\) is from MoO\(_3\), the resolution was not optimized. This led to the undistinguishable peaks at 374 and 329 cm\(^{-1}\). Jeziorowski reported peaks at 380, 367, 337, and 292 cm\(^{-1}\).\(^{21}\) He also reported another peak at 117 cm\(^{-1}\) which we identify as an Ar\(^+\) laser plasma line. We have seen this line regardless of the sample we used. The aperture of the plasma-line eliminator is not sufficiently narrow to completely eliminate plasma lines near the elastically scattered laser light. Excluding this 117 cm\(^{-1}\) line and the MoSi\(_2\) signal, curve (d) represents a perfect Raman spectrum of MoO\(_3\).\(^{21,22,23}\) The 338 cm\(^{-1}\) peak has been covered by the stronger MoO\(_3\) signal. In traces (a) and (b) the oxide impurity was too small to produce a large Raman signal, and only the strongest peak of the oxide was observed. In (b), because the powder was annealed in vacuum, there is no way for oxide to be removed, so the oxide's signal remains.

Raman spectra from Mo/Si thick multilayers are shown in Figure 2. The spectrum of the as-deposited multilayer sample only gives a broad bump located approximately at 480 cm\(^{-1}\) (see Figure 2 (a)). We attribute this feature to amorphous silicon.\(^{24}\) The silicon layers of Mo/Si multilayers have previously been reported to be amorphous on the basis of transmission electron microscopy.\(^{25}\) Curve (a) provides direct evidence that these layers are amorphous and, furthermore, indicates that the amorphous silicon is highly defective since the peak is broad and weak. The metal Mo belongs to space group Im\(_3\)m (O\(_h\)) and the Mo atoms are at Wyckoff 2(a) positions. This structure does not allow any active modes and we see no evidence of them. We also do not see evidence of Mo-Si bonds in the as-deposited multilayers.

We subjected our sample to successively higher temperature anneals starting at 200 °C and eventually reaching 1000 °C. The amorphous silicon peak disappears at about 300 °C and no other changes are seen until 1000 °C. Curve (c) of Figure 2 shows the Raman spectra after the sample's final vacuum annealing at 1000 °C. The disappearance of the Raman peak due to amorphous silicon is noteworthy since it is opposite to what is often observed with the annealing of thicker layers of room-temperature deposited amorphous silicon. This can produce a better bonded or structurally more perfect anneal-stable amorphous silicon, possessing a Raman spectrum which is more intense and shows two humps, one centered at about 480 cm\(^{-1}\) and the other one lower at 170 cm\(^{-1}\). The intensity of the peaks typically increases and the peaks narrow with annealing.\(^{24}\) The fact that this same phenomenon does not occur for the metal/silicon samples indicates that annealing may produce significantly different changes in them on an atomic level than it does in pure silicon films and silicon/dielectric multilayers. The loss of signal indicates that the Raman cross section of the silicon layers has decreased. It may be that the material is becoming more defective -- perhaps because metal atoms are diffusing into the silicon.

After this 1000 °C annealing three peaks were seen: two peaks at 323 and 438 cm\(^{-1}\), respectively, which can be attributed to crystalline MoSi\(_2\), and one peak at 520 cm\(^{-1}\), which can be attributed to crystalline silicon.

We believe that the appearance of molybdenum silicide peaks means that extensive interdiffusion has occurred and that in the interdiffusion region crystallization has begun, and that the crystallites formed are thick enough to produce detectable Raman signals. The annealing process may have produced a portion of the silicon component of multilayer crystalline. There is some possibility of the 520 cm\(^{-1}\) peak coming from the substrate through the film. The 250 nm thickness of the multilayer is sufficient to block any Raman emissions from the substrate in the as-deposited film. Interdiffusion and reaction at 1000 °C may have made the film more transparent at the laser's 514 nm line, however. We have also inspected the sample's surface for cracks or pinholes that would give the laser probe access to the substrate. In addition, the ratio of the amplitude of the 520 cm\(^{-1}\) peak to the MoSi\(_2\) peaks does not vary as the sample's position is changed. If it were due to pinholes it would vary. The SEM pictures of as-prepared and after 1000 °C of multilayer surfaces were shown in Figure 3. Comparing to the result of as-prepared multilayer sample, the 1000 °C annealing made bigger grains appeared on the surface and no pinholes or cracks has been seen. This indicates that some crystallization does happen. At the valley parts of the surface, the effective thickness decreased which made the substrate signal detectable.
The SAXRD measurement results of thick multilayer sample are shown in Figure 4. Six peaks can be seen before annealing. 1000 °C annealing makes the multilayer structure deteriorate but there is still one peak remains which means the multilayer structure has not been totally destroyed yet. The LAXRD measurement results are shown in Figure 5. Before annealing, only two broad peaks have been observed (excluding substrate crystalline silicon signals). Peak at 14.2 ° is Si(111) from Si layers and peak at 20.3 is metal Mo(110) from Mo layers. After 1000 °C annealing, the spectrum displayed multiple new lines instead of the broad Mo peak. The identifications are shown in Table I according to standard powder diffraction file. From the LAXRD data we can see the evidence of existing molybdenum silicides which are caused by interdiffusion and interaction at multilayer interfaces.

Table I. Identification of LAXRD data of 1000 °C annealed Mo/Si multilayer. The first column lists the peaks detected. The following columns list the signal identifications [hkl] from the possible silicides.

<table>
<thead>
<tr>
<th>Peak#</th>
<th>d(nm)</th>
<th>Mo2Si</th>
<th>t'-MoSi2</th>
<th>Mo3Si2</th>
<th>h-MoSi2</th>
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<tr>
<td>1</td>
<td>0.3207</td>
<td></td>
<td></td>
<td>211</td>
<td></td>
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<tr>
<td>2</td>
<td>0.301</td>
<td></td>
<td>101</td>
<td>310</td>
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</tr>
<tr>
<td>3</td>
<td>0.2715</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>0.2447</td>
<td>200</td>
<td></td>
<td>002</td>
<td></td>
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<td>5</td>
<td>0.2275</td>
<td></td>
<td>110</td>
<td>112</td>
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<td>6</td>
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<td>202</td>
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<td>0.2107</td>
<td>211</td>
<td>103</td>
<td>112</td>
<td>222</td>
</tr>
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</table>

The same Raman and SAXRD measurements have been performed on the thin multilayer sample. The comparisons of the results between thick and thin samples provide some useful information. The Raman spectra of the thin multilayer sample (see Figure 6) shows a similar tendency of evolution as that of thick multilayer sample. The silicide signal from interface did not appear until after 1000 °C annealing. But there are also some differences in thin sample spectra (see Figure 7). These differences can be explained by the fact that the film is very thin. First of all, the peak at 520 cm⁻¹ from crystalline silicon signal was observed even before annealing. We attribute this signal to the substrate. Based on its location we presume that the film is too thin to block the substrate signal entirely. Secondly, after 1000 °C annealing, the silicide's signal is stronger than that of the thick film because there is less attenuation in it. In addition to the conclusion made above, it should be noted that the silicon signal is not as strong in the thin sample case as for the thick sample. This supports the assertion that this peak originates both from the substrate and crystallized Si layers in the thick, annealed multilayer sample. In the thin sample annealing has made most of the Si layer react to form silicide which then crystallized.

Figure 8 shows the SAXRD results of the thin multilayer sample. As the annealing temperature increases the diffraction peak produced by the multilayer structure decreases until it disappears. This indicates that the multilayer structure has been totally destroyed. Annealing at the same temperature did not eliminate the multilayer structure in the thick sample. The conclusions obtained from these data agree with those obtained from the above discussions.

Several authors have reported that annealing Mo/Si multilayers under 600 °C will transform completely into polycrystalline mixtures of Mo3Si2 and both the hexagonal and tetragonal phases of MoSi2. We have not seen the appearance of any molybdenum silicide until after annealing at 900 °C or above, and only tetragonal silicide has been obtained at these temperatures. This transition temperature is probably affected by the sample deposition
environment (sputtering gas pressure etc.) and vacuum annealing. It has been reported that Mo/Si systems fabricated by different methods shows different phase transition temperatures. Further annealing studies may be required to see the hexagonal phase by Raman.

Raman spectroscopy may prove to be a useful and new method of characterizing Mo/Si multilayers. As the data acquisition speed increases dramatically (by using CCD detector instead of PMT) it can be further developed as an in-situ characterizing technique during multilayer fabrication.

4. CONCLUSIONS

The Raman spectroscopy technique has been first applied to study polycrystalline MoSi$_2$ powder and characterize the as-prepared and annealed Mo/Si soft x-ray multilayer mirrors. The two active Raman modes of MoSi$_2$ located at 323 and 438 cm$^{-1}$ have been observed in powder MoSi$_2$ and in annealed Mo/Si multilayers. The observation of the signal due to amorphous silicon in as-deposited multilayers and its loss later with relatively modest heating indicates that significant atomic reordering is occurring at relatively modest temperatures. This observation and the observation of MoSi$_2$ in the annealed multilayer indicate that Raman spectroscopy can be a useful technique in characterizing Mo/Si multilayers and studying their interfaces.

5. REFERENCES


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Figure 1. Raman spectra of MoSi₂ powder: (a) as-received, (b) annealed in vacuum at 1000 °C for 1 h, (c) etched in 10 M KOH and dried in air, (d) annealed in air at 500 °C for 1 h.

![Raman Spectra of MoSi₂ Powder](image)
Figure 2. The Raman spectra of thick Mo/Si multilayer: (a) as-deposited, (b) annealed in vacuum at 400 °C for 1 h, (c) annealed in vacuum at 1000 °C for 1 h.

![Raman spectra diagram](image-url)
Figure 3. SEM images of thick Mo/Si multilayer sample: (a) as-deposited, (b) annealed in vacuum at 1000°C for 1 h.
Figure 4. Small angle x-ray diffractions ($\lambda = 0.154$ nm) measured from thick Mo/Si multilayer: 
(a) as-deposited, (b) annealed in vacuum at 1000 °C for 1 h.
Figure 5. Large angle x-ray diffractions measured from thick Mo/Si multilayer and MoSi$_2$ powder: (a) as-deposited multilayer, (b) as-received MoSi$_2$ powder, (c) multilayer annealed in vacuum at 1000°C for 1 h.
Figure 6. The Raman spectra of thin Mo/Si multilayer: (a) as-deposited, (b) annealed at 300 °C, (c) annealed at 1000 °C.
Figure 7. The Raman spectra of both thick and thin multilayer samples after 1000 °C vacuum annealing: (a) thin sample, (b) thick sample.
Figure 8. SAXRD measured from the thin Mo/Si multilayer: (a) as-deposited, (b) annealed at 300 °C for 1 h, (c) annealed at 600 °C for 1 h, (d) annealed at 900 °C for 1 h, (e) annealed at 1000 °C for 1 h.
DISCUSSION

Q This assumption that these very thin 0.2 nm layers are going to be representative of the interface may not be correct. You have diffusion and sputtering effects that are different. Would you comment.

A We have made even thinner layers. In multilayers with low angle measurements you can actually see both layers; there are no diffraction peaks. For this one there are diffraction peaks, which indicates that even though it is very thin the multilayer structure is still maintained. The sputtering involvement will give some defect on the multilayer. The purpose of doing this sample is to try to magnify the interface effect, not to care about the reflectivity and those kind of things.

Q The 1000° C annealing temperature is probably not a temperature that these mirrors are ever going to see in the application, lithography or whatever. Are you going to see any effect on real mirrors or is there an integrated effect of like 100° C times 10 hours rather than 1000° C times one hour?

A People report that even at lower temperature the longer times of annealing (some people anneal for 1000 hours) does produce some silicide from the interface. We see it for 5 hours annealing. Thus the lower temperature will produce some interface signals.