Asymmetric diffusion at the interfaces in multilayers

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Nanoscale diffusion at the interfaces in multilayers plays a vital role in controlling their physical properties for a variety of applications. In the present work depth-dependent interdiffusion in a Si/Fe/Si trilayer has been studied with sub-nanometer depth resolution, using x ray standing waves. High depth-selectivity of the present technique allows one to measure diffusion at the two interfaces of Fe namely, Fe-on-Si and Si-on-Fe, independently, yielding an intriguing result that Fe diffusivity at the two interfaces is not symmetric. It is faster at the Fe-on-Si interface. While the values of activation energy at the two interfaces are comparable, the main difference is found in the pre-exponent factor suggesting different mechanisms of diffusion at the two interfaces. This apparently counter-intuitive result has been understood in terms of an asymmetric structure of the interfaces as revealed by depth selective conversion electron Mössbauer spectroscopy. A difference in the surface free energies of Fe and Si can lead to such differences in the structure of the two interfaces.

Atomic diffusion is fundamental to many processes in material science such as microstructure development, non-martensitic phase transformation, stress relaxation etc. In multilayers, atomic diffusion at the interfaces plays a vital role in controlling their physical properties for a wide variety of applications. In x ray and neutron mirrors intentionally diffused interfaces have been used to reduce the higher order contamination [1]. In tunnel magnetoresistance multilayers, thermal annealing can increase magnetoresistance by orders of magnitude [2]. In giant magnetoresistance (GMR) multilayers, interdiffusion can significantly affect the GMR [3]. In spin valves with Mn based antiferromagnetic layer, Mn diffusion can seriously degrade the performance [4]. Co diffusion in Sm-Co/Fe exchange-spring magnet films is known to improve the exchange coupling [5]. While atomic diffusion in bulk solids is a widely studied and fairly well understood phenomenon, a reasonable understanding of the interfacial diffusion in multilayers has yet to emerge. Several factors like a steep concentration gradient at the interfaces, interfacial stresses and disorder may significantly modify the diffusion in multilayers. This has resulted in unexpected interfacial phenomenon like a non-parabolic shift of phase boundaries in the presence of strong composition dependence of diffusivity [6].

X ray standing waves generated by total external reflection of x rays from buffer layer [7], or in a multilayer [8, 9] have been used for concentration profiling of various elements. In the present work, we exploit the depth selectivity of x ray standing waves for studying the depth dependent interdiffusion in a Si/Fe/Si trilayer with sub-nanometer depth resolution. High depth sensitivity of the technique allows one to measure diffusion at the two interfaces of Fe namely Fe-on-Si and Si-on-Fe independently. An intriguing finding of the present work is that the Fe diffusivity at the two interfaces is not symmetric; diffusion is faster at the Fe-on-Si interface. In order to understand this apparently counter-intuitive result, conversion electron Mössbauer spectroscopy (CEMS) has been used to study the interfacial structure. It is found that there is a significant difference in the structure of the two interfaces, resulting in different diffusivities at the two interfaces. A difference in the surface free energies of Fe and Si can lead to such difference in the structure of the two interfaces.

The structure of the multilayer used for diffusion measurements is \([W (2.0 nm)/Si (3.1 nm)]_{10}/Si (3.8 nm)/Fe (2.7 nm)/Si (7.0 nm)\) (referred as SW-ML). The deposition was done using ion beam sputtering in a vacuum chamber with a base pressure of \(1 \times 10^{-7}\) mbar [10]. A broad beam Kaufman type ion source was used with Ar ions of energy 1 keV and a beam current of 20 mA. The bottom \([W (2.0 nm)/Si (3.1 nm)]_{10}\) multilayer is used to generate x ray standing waves [9]. On the top of this multilayer Si (3.8 nm)/Fe (2.7 nm)/Si (7.0 nm) structure was deposited without breaking vacuum. A thickness of 3.8 nm for the first Si layer is chosen in such a way that the Fe layer lies roughly midway between two antinodes of the x ray standing waves generated in W/Si multilayer at the Bragg peak. This point is clear from the inset of Fig. 1 which shows the contour plot of x ray intensity as a function of depth and the scattering vector \(q\). At \(q = 1.35\) nm\(^{-1}\) which corresponds to the center of the Bragg peak of W/Si multilayer, Fe layer is midway between the two antinodes.

Simultaneous x ray reflectivity (XRR) and x ray fluorescence (XRF) measurements were done using Bruker D8 diffractometer fitted with a Göbbel mirror on the incident beam side in order to obtain a parallel monochromatic beam of Cu Kα radiation. Fluorescence spectrum was measured using a Ketur detector with an energy resolution of 200 eV. Thermal annealing of multilayer struc-
ture was done in a vacuum of $10^{-6}$ mbar in order to induce interdiffusion at the interfaces of Fe and Si layers. A separate study on the annealing behavior of W/Si multilayer showed that at least up to 623 K, the W/Si multilayer structure is stable with almost no change in its x-ray reflectivity. Thus the thermal annealing in the present multilayer essentially induces interdiffusion at the interfaces of Fe and Si layers only.

In order to study the Fe-on-Si and Si-on-Fe interfaces using CEMS, the following two multilayer structures were also prepared using ion beam sputtering: (i) substrate/ [Si(6 nm)/$^{57}$Fe(2 nm)/Fe(2 nm)]$_{10}$, and (ii) substrate/ [Si(6 nm)/Fe(2 nm)/ $^{57}$Fe(2 nm)]$_{10}$, designed as ML1 and ML2 respectively. In the first multilayer the 2 nm thick Fe layer lies on Fe-on-Si interface while in the second multilayer it lies on Si-on-Fe interface. Since Mössbauer measurements are sensitive only to $^{57}$Fe, the first multilayer gives information preferentially about the Fe-on-Si interface, while the second one gives information about Si-on-Fe interface. The CEMS measurements were done using a Wissel Mössbauer spectrometer and a gas flow proportional counter with 95% He + 5% CH$_4$.

SW-ML was isochronally annealed at 473 K, 498 K, 523 K, 598 K and 623 K for 1 h each. Figure 1 gives some representative XRR of SW-ML after various stages of annealing. The XRR of this multilayer is dominated by that of W/Si multilayer mirror, with only small modulation in the $q$ region below the Bragg peak attributable to the top Si/Fe/Si trilayer. Figure 2 shows Fe-fluorescence data of SW-ML measured simultaneously with XRR. The region around the Bragg peak in which x-ray standing waves are generated, is highlighted in the figure by shaded area. One may note that in the as-deposited sample Fe-fluorescence exhibits two well-defined peaks in the region where x-ray standing waves are generated. The origin of these two peaks can be understood from the contour plot of x-ray intensity as a function of $q$ and depth from the surface of the multilayer. The rectangle represents the position of Fe layer.
ence data it is difficult to estimate the roughnesses of the two interfaces separately. However, at 498 K and above roughnesses of the two interfaces can be determined individually with good reliability. In order to demonstrate this point, the fitting of 498 K data obtained by taking the roughnesses of the two interfaces to be equal is also shown in Fig. 2 (dashed curve). The best fitted curve clearly deviates from the experimental data.

The variation in the roughnesses of the two interfaces with the thermal annealing has been used to estimate the diffusivity of Fe at the two interfaces using the relation: $$D(T) = \frac{\sigma^2(T_2) - \sigma^2(T_1)}{2t},$$ where $D(T_2)$ is the diffusion coefficient at temperature $T_2$, $\sigma(T_1)$ and $\sigma(T_2)$ are the roughnesses of an interface before and after annealing at temperature $T_2$ for time $t$. Figure 3 gives the Arrhenius plot of $\ln D$ versus $1/T$ at the two interfaces of Fe. One finds that there is a significant difference in Fe diffusivities at the two interfaces. A linear fit to the experimental data as expected from the Arrhenius temperature dependence of the diffusivity $D(T) = D_0 \exp(-E/K_B T)$, yields the activation energy $E$ as well as the pre-exponent $D_0$ for the diffusion. The experimentally obtained values are shown in Table I. One finds that the activation energies for diffusion at the two interfaces are comparable within experimental errors. However, the pre-exponent $D_0$ is significantly high for Fe-on-Si interface as compared to that for Si-on-Fe interface, resulting in a significantly high diffusivity at Fe-on-Si interface in the temperature range studied in the present experiment.

At first glance the above results appear to be counter intuitive as both interfaces ought to be identical, having Fe layer on one side and Si layers on the other. In order to understand the possible reason for this difference in the diffusivity at the two interfaces, a detailed study of the structure of the two interfaces has been done using CEMS on the samples ML1 and ML2. As discussed in the experimental section CEMS of ML1 is sensitive to Fe-on-Si interface while that of ML2 is sensitive to Si-on-Fe interface.

Figure 4(a) shows Mössbauer spectra of ML1 and ML2 in as-deposited state. The spectra of both the specimens are fitted with three overlapping components: (1) a sharp sextet with hyperfine field about 33 T, (2) a broad magnetic component, having a distribution of hyperfine fields and (3) a non-magnetic doublet. The sharp sextet represents the bulk of α-Fe, while the broad magnetic component and the doublet represent the Fe atoms in the interfacial region. Further it is known that in Fe-Si alloy, if iron concentration is less than 50%, it becomes non-magnetic [12]. Therefore, the area under the doublet represents the fraction of Fe atoms in the interfacial region having iron concentration less than or equal to 50%. Results of fitting are given in Table I. One may note that the area under the sharp sextet in both the specimens is about 30%, within experimental errors. However, the relative areas of broad magnetic component and the doublet as well as the shape of the hyperfine field distribution of broad magnetic component [Fig. 4(b)] are very different in the two specimens. This suggests that the structure of the two interfaces namely Si-on-Fe and Fe-on-Si interface may be different. A rough estimate of the interface roughness as obtained from the width of the intermixed region comes out to be 0.78 nm, which is in agreement with that obtained from x ray measurements. From Table (1), one finds that at Si-on-Fe interface, the fraction of Fe atoms in the doublet is 18%, which is about 23% of the total Fe atoms in the intermixed region. This area fraction agrees very well with that expected for an error function concentration profile [Fig. 5]. Thus, Mössbauer measurements suggest that the concentration profile at
Thus, at Fe-on-Si interface diffusion occurs via the FeSi$_2$ self-diffusion in stoichiometric D$_{2}$bcc Fe(Si) phase. In literature the diffusivity data for interlayer, while at Si-on-Fe interface diffusion occurs via present work at the two interfaces with bulk diffusivities, meaningful to compare the diffusivities obtained in the metallic compounds of FeSi are given. However, it is not suggesting that the composition of interlayer is FeSi$_2$ doublet of ML1 match very well with those of FeSi$_2$. This suggests that at Fe-on-Si interface, there is an interlayer of Fe$_{1−x}$Si$_{2x}$, resulting in a plateau in a concentration profile. The isomer shift and quadrupole values of the doublet of ML1 match very well with those of FeSi$_2$, suggesting that the composition of interlayer is FeSi$_2$. Thus, at Fe-on-Si interface diffusion occurs via the FeSi$_2$ interlayer, while at Si-on-Fe interface diffusion occurs via bcc Fe (Si) phase. In literature the diffusivity data for self-diffusion in stochiometric DO$_3$ phase as well as intermetallic compounds of FeSi are given. However, it is not meaningful to compare the diffusivities obtained in the present work at the two interfaces with bulk diffusivities, since it is known that the multilayer diffusivities at the interfaces are very different because of high concentration of defects and possible concentration gradient.

The difference in the structure of the two interfaces can be understood in terms of the difference in the surface free energies of Fe(2.9 Jm$^{-2}$) and Si(1.2 Jm$^{-2}$). During the deposition of Fe on Si, the surface free energy of Si being lower, Si atoms try to move to the surface guided by the chemical driving force. This would lead to a stronger mixing at the interface and a possible formation of FeSi$_2$ compound. On the other hand, during the deposition of Si on Fe, no such chemical driving force exists, therefore the intermixing at Si-on-Fe interface would take place as a result of random thermal motions only and hence concentration profile is expected to be an error function.

In conclusion, x-ray standing wave technique has been used to get concentration profile of Fe layer in Si/Fe/Si trilayer. The precision of this technique is sufficient to differentiate between the two interfaces of the Fe layer. This allows one to study interdiffusion at the two interfaces namely Fe-on-Si and Si-on-Fe independently. Interestingly the diffusivities at the two interfaces are significantly different. This seemingly counter-intuitive result can be understood in terms of a difference in the structure of the two interfaces in the as-deposited film itself. CEMS measurements show that while at Si-on-Fe interface, the Fe concentration profile is an error function, at Fe-on-Si interface an interlayer exists with approximate composition of FeSi$_2$. This difference in the structure of the two interfaces is the cause of different diffusivities at the two interfaces. Besides being of fundamental importance in understanding the interfacial diffusion at nanometer scale, present results have important implications on the use of controlled thermal annealing for tailoring the properties of multilayers for a wide variety of applications.

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