First-principles predictions of ruthenium-phosphorus and ruthenium-boron glassy structures and chemical vapor deposition of thin amorphous ruthenium-boron alloy films

Daniel E. Bost, Hyun-Woo Kim, Chia-Yun Chou, Gyeong S. Hwang, John G. Ekerdt *

Department of Chemical Engineering, University of Texas at Austin, Austin, TX 78712, USA

A R T I C L E   I N F O

Article history:
Received 20 August 2015
Received in revised form 22 November 2016
Accepted 12 December 2016
Available online 18 December 2016

Keywords:
Amorphous thin film
Ruthenium alloy
Copper diffusion barrier
Boron
First-principles density-functional calculations

A B S T R A C T

First-principles density-functional calculations are presented revealing that Ru(P) and Ru(B) alloys with moderate P or B content can result in a glassy structure exhibiting strong chemical short-to-medium range order. Amorphous phases are predicted to be energetically more favorable than the crystalline counterparts for the Ru(P) and Ru(B) alloys above ~ 20 at.% P and ~ 10 at.% B. The relative stability of amorphous and crystalline Ru(B) alloys is examined along with local atomic ordering in the amorphous alloys. The growth of ultrathin (3 nm) amorphous Ru(B) alloy films of varying B concentration via chemical vapor deposition is explored using Ru3(CO)12 and B2H6 as the Ru and B sources, respectively. Experiments reveal the films grown at 250 °C are amorphous at B contents in excess of 15 at.% and polycrystalline below 10 at.% B, consistent with first-principles predictions. Amorphous Ru(B) films remain amorphous following annealing at 450 °C and become polycrystalline at 500 °C. Film resistivity ranged from 40 to 120 μΩ-cm and was independent of B loading. Electric field stress tests to failure for Cu/3-nm Ru(B)/SiO2/Si stacks are used to indicate suitability of Ru(B) as a copper diffusion barrier layer.

© 2016 Elsevier B.V. All rights reserved.

1. Introduction

As the characteristic dimensions of microelectronic devices shrinks and RC delay becomes a more significant factor in device design, interest in extremely thin, platable, conformal diffusion barriers for copper interconnects has grown [1], with industry predictions indicating the necessity of a conformal barrier layer 1.9 nm thick for the 16-nm technology node [2]. A two-layer barrier of Ta and TaN is usually the copper diffusion barrier of choice 

Several responses to this have been proposed and tested. Atomic layer deposition (ALD) and plasma-assisted deposition have been used to grow single-crystal films with few or no grain boundaries using known materials such as TaN [8]. Other materials such as nitrogen have been used to stuff the grain boundaries, either by saturating the material during growth [5,9] or depositing a secondary layer of material that diffuses and fills the grain boundaries after growth [10]. A variety of metal alloys such as RuTa [11], Ti(N) [12], W-doped indium oxides [13], Co-W alloys [5], Ru(P) [14], and VOx and self-forming V-Cu alloys [15] have also been examined for barrier properties at low thickness. Some three-element alloys and non-metallic components have also shown promising results, such as TaSiC [16] and WGeN [17], and graphene [2]. Herein, we examine the elimination of grain boundaries by the addition of an alloying element during chemical vapor deposition (CVD) to cause the potential barrier material to grow in an amorphous rather than crystalline phase.

Ruthenium grown by CVD or ALD has arisen as a potential single-layer alternative to Ta/TaN stacks [6,11,18,19,20,21]. Previous studies of Ru in the 10-nm thickness range have shown the addition of P causes the Ru(P) alloy to grow as an amorphous film [6,20]. Ruthenium films containing ~ 17 at.% P remain amorphous upon annealing to 360 °C [14,22]. Further, 5 and 3-nm-thick amorphous Ru(P) films function as Cu diffusion barriers [7,14]. Ab initio molecular dynamics calculations [6] showed that the Ru(P) alloy with 20 at.% P can result in an amorphous structure exhibiting the topological and strong short-range order.

Generally, the atomic size ratio and negative formation energy are considered to be the main parameters to promote the properties of amorphous alloys. To gain quantitative understanding of the effect of the atomic size ratio (λ) and alloying elements on amorphous alloy formation ability we experimentally and computationally examine boron (B), which is different in atomic size and chemistry with P, in creating
an amorphous Ru(B) (**a-Ru(B)**) alloy as an alternative to amorphous Ru(P) (**a-Ru(P)**) alloys.

We use first principles density-functional theory calculations to predict the three dimensional (3D) amorphous alloy structures and bonding properties of Ru(P) and Ru(B) alloys. Analyzing the details of the structure changes in the different alloy systems reveals icosahedra dominate the medium range ordering in both systems, while the short range ordering is primarily governed by the atomic size ratio. Twenty at.% P was found to favor a-Ru(B) [6,14] and results reported herein indicate an a-Ru(B) film is favored above 10 at.% P. This study also reports the use of CVD to produce a-Ru(B) thin films at B content above 10 at.% using Ru(CO)_{12} as the Ru source and BiH_{3} as the B source. Diborane is selected because it decomposes completely on Ru(0001) between 230 and 400 K, depending on coverage [23], making it amenable to use at typical film growth temperatures.

2. Equipment and procedures

The equipment and procedures used to produce and analyze Ru(B) films are similar to those discussed in previous work on Ru(P) films [6,14] in order to minimize potential processing differences when comparing the films. Briefly, Ru(B) films were grown in a cold-wall CVD chamber on 15k Å SiO_{2} deposited using tetraethyl orthosilicate on pure Cu, Al, and Ru targets as needed (Kurt J. Lesker & Co) and using sputtering guns (AJA International 320-2a) loaded with 99.999 at.% Ru precursor, Ru_{3}(CO)_{12} (Aldrich, 99%), was a volatile solid-state compound distributed from a saturator into the growth chamber using H_{2} as the carrier gas, with the saturator kept at 85 °C. The Ru precursor and carrier gas was regulated to a partial pressure in the chamber. B_{2}H_{6} (Voltaix, research grade) at 15 ppm in H_{2} was used at typical gas pressures between 0 and 24 Pa when not in use. The barrier testing method for copper diffusion under field stress is very similar to that outlined in Reference [7] and described in Reference [14]. A Cu/Ru(B)/SiO_{2}/Si stack was manufactured by growing an Ru(B) film on the SiO_{2} substrate, then depositing a thick (~10 μm) Cu film on top of the film by PVD without exposing the film to atmosphere. After removing the binary film from vacuum, test devices consisting of 0.3-mm diameter Cu dots on top of 1-mm diameter Ru(B) dots were fabricated. A single 20 mm × 20 mm square of wafer could produce as many as 36 device stacks for testing, but in practice about half of these were typically damaged at some point in the course of transportation and processing and failed immediately under testing, leaving 10 to 15 stacks for testing from a given film. In the failure test, a device stack was placed under a strong electric potential, and the leakage current monitored (Agilent 4156C Semiconductor Parameter Analyzer) until capacitive breakdown was observed, resulting in a sudden abrupt jump in current flow.

Our calculations were performed using density functional theory (DFT) [29] within the Perdew-Berke-Ernzerhof generalized gradient approximation (GGA-PBE) [30] as implemented in the Vienna **ab initio** Simulation Package (VASP) [31]. We employed Vanderbilt-type ultra-soft pseudopotentials to describe the interaction between core and valence electrons, and a plane-wave basis set with a kinetic energy cutoff of 300 eV. The model structures for amorphous Ru(P) and Ru(B) were obtained using **ab initio** molecular dynamic (AIMD) simulations. We first created the amorphous structure of Ru, and then replaced a given fraction of Ru with P or B. The Ru-P or Ru-B alloy systems, each containing 72 atoms in a periodic supercell, were melted at 3500 K for 3 ps with a time step of 1 fs, and then quenched to 500 K at a rate of 1.5 K/fs, followed by static structural optimization. Here the temperature was controlled using velocity rescaling. The structures of crystalline Ru(P) and Ru(B), as well as ruthenium, phosphorus and boron, are summarized in Table 1. For each amorphous system, 3–5 independent samples were considered to obtain good statistics in predicting its energetics and structure. The Brillouin zone integration for periodic 72-atom supercells was performed using one k-point (at Gamma) for AIMD simulations and 2 × 2 × 2 Monkhorst-Pack k-point mesh [32] for geometry optimization and energy calculations. For further structural and electronic property analysis, the Ru_{80}P_{20} and Ru_{87}B_{13} alloy structures were modeled using 144-atoms (consisting of 115 Ru and 29 P atoms in Ru_{80}P_{20}) and 19 Ru and 13 B in Ru_{87}B_{13}) in a periodic supercell volume of 2 nm³. While no simulation study has been reported for the Ru(P) and Ru(B) systems, the N_{2}Ru_{P_{20}} structure obtained using the same procedure (1.49:1) [27]. The manual nature of the parameter re-adjustment introduces an error range of ±10 at.% to the final calculated C content using this technique due to the low sensitivity of the C 1s peak compared to the Ru 3d peaks.

Film thicknesses were obtained by comparing two measurements: Si 1s peak attenuation calculated as noted above, and an **ex situ** X-ray reflectivity (XRR) measurement made using a Bruker-Nonius D8 diffractometer equipped with a sealed-tube, long fine focus Cu K_{α} (1.5418 Å) source, a Si–X solid state detector, and operated in the Bragg-Brentano configuration. Film roughness was measured via atomic force microscopy (AFM) (Agilent Instruments 5500). Films were judged to be continuous when the XPS– and XRR-derived thickness estimates matched, the XRR measurement gave a clean and definite pattern, and the AFM-based film root mean square (RMS) roughness did not exceed half of the calculated thickness. Only continuous films are included in the data reported herein.

Film crystallinity was evaluated **ex situ** using X-ray diffraction (XRD) (Bruker-Nonius D8) operated at glancing incidence angle (0.5°). Films were designated X-ray amorphous based on the lack of appearance peaks associated with crystalline Ru [28]. As-deposited and annealed films were studied for crystallinity. Annealing was done in the XPS chamber using a stage heating bulb under 1.3 mPa of H_{2}; the films were exposed to atmosphere between annealing steps due to the XRD apparatus being open to air.

The barrier testing method for copper diffusion under field stress is very similar to that outlined in Reference [7] and described in Reference [14]. A Cu/Ru(B)/SiO_{2}/Si stack was manufactured by growing an Ru(B) film on the SiO_{2} substrate, then depositing a thick (~10 μm) Cu film on top of the film by PVD without exposing the film to atmosphere. After removing the binary film from vacuum, test devices consisting of 0.3-mm diameter Cu dots on top of 1-mm diameter Ru(B) dots were fabricated. A single 20 mm × 20 mm square of wafer could produce as many as 36 device stacks for testing, but in practice about half of these were typically damaged at some point in the course of transportation and processing and failed immediately under testing, leaving 10 to 15 stacks for testing from a given film. In the failure test, a device stack was placed under a strong electric potential, and the leakage current monitored (Agilent 4156C Semiconductor Parameter Analyzer) until capacitive breakdown was observed, resulting in a sudden abrupt jump in current flow.

In **in situ** XPS was performed using a Physical Electronics 3057 XPS system with an MgKα X-ray emitter and a fixed 30° angle in a third cold-walled chamber. This system included an **in situ** Ar⁺ ion sputtering filament that allowed the removal of material and the construction of composition depth profiles to judge the compositional uniformity. Film thickness was calculated by measuring the attenuation of the Si 1s signal in Ru as tabulated in the NIST database [24], with the pre-attenuation peak area regularly calibrated and sputter rates recalculated with each sample and checked for consistency with previous sputter rates. Film composition was derived by weighting the measured peak areas for the component substances by their known XPS sensitivity factors [25]: 4.273 for Ru 3d, 0.159 for B 1s, and 0.296 for C 1s. Because the C 1s peak overlaps the Ru 3d_{5/2} peak, those peaks were deconvoluted with freeware analytical software [26] using a Shirley background approximation with three end-points to either side of the peak, with parameters manually adjusted until the three component signal peaks reflect the known ratio of the Ru 3d_{5/2}:3d_{3/2} peaks in pure ruthenium.
shows excellent agreement with that from previous AIMD simulations [33].

3. Results and discussion

3.1. Relative stability between amorphous and crystalline Ru(P) and Ru(B) alloys

To examine the relative stability of the amorphous versus ordered structures of Ru(P) and Ru(B), we calculated and compared the total energies of Ru(P) and Ru(B) supercells at varying P and B contents. Here, we only considered the regions below 30 at.% of P and 20 at.% of B, where the transition from a more stable ordered to amorphous structure was found. As shown in Fig. 1, the calculation results demonstrate that the amorphous Ru(P) and Ru(B) structures become energetically more favorable than their ordered counterparts when the P and B contents are above 20 at.% and 10 at.% respectively. Here, the ordered structures were obtained by replacing Ru with P or B, starting with the hexagonal close packed structure of pure Ru (with a lattice constant of 2.70 Å). Similarly, the amorphous structures were constructed by replacing Ru with P or B atoms in α-Ru; the replacing sites were carefully chosen to ensure homogeneous distribution of P/B. Both ordered and amorphous alloys were modeled using a 72-atom supercell, and the atomic positions and the supercell volume were optimized to minimize the total energy.

Fig. 2 shows the predicted volume change exhibiting a distinctive nonlinear trend for the Ru(P) alloy but a linear decrease for the Ru(B) alloy. The Ru(P) alloy volume becomes a minimum at 20 at.%, yielding the highest packing density, while the Ru(B) alloy volume decrease, leads to increased packing density as B content increases. This is largely due to the large atomic size difference between Ru and B atoms. As expected, the crystalline phase is slightly denser than the amorphous alloy of corresponding composition. The smaller atomic size of B than P decreases the composition limit to form an amorphous structure.

For the sake of comparison, we also evaluated the crystalline-to-amorphous transition using the following universal correlation between glass formability and atomic volume ratio in a binary alloy [32]: 

$$C_{\text{min}} = \frac{0.1}{\gamma^3 - 1},$$

where $C_{\text{min}}$ is the minimum concentration of solute element needed to produce the amorphous phase and $\gamma$ ($= R_b/R_a$) is the radius ratio of the solute ($R_b$) to the solvent ($R_a$). The elastic model has been widely adopted in predicting the structural instability of a crystalline binary alloy (as a function of solute concentration) due to the size difference between solvent and solute atoms. Taking 1.338 Å [34], 1.06 Å [35], and 0.82 Å [36] for Ru, P, and B atomic radii, respectively, the required minimum concentrations for P (in an amorphous Ru(P) alloy) and B (in an amorphous Ru(B) alloy) are estimated to be 19.5 at.% and 11.5 at.% respectively, which is in good agreement with our DFT calculations.

<table>
<thead>
<tr>
<th>Lattice constants (Å)</th>
<th>Volume(Å³)</th>
<th>K-pts.</th>
<th>#Ru</th>
<th>#P(B)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ru</td>
<td>a = 2.732, c = 4.290, γ = 120</td>
<td>27.65</td>
<td>96</td>
<td>2</td>
</tr>
<tr>
<td>P</td>
<td>a = 9.274, b = 9.260, c = 24.159, β = 105.74</td>
<td>2009.85</td>
<td>4</td>
<td>84</td>
</tr>
<tr>
<td>B</td>
<td>a = 10.738, b = 11.385, c = 26.249</td>
<td>1002.56</td>
<td>4</td>
<td>131</td>
</tr>
<tr>
<td>Ru₄P</td>
<td>a = 5.595(5.902), b = 3.968(3.859), c = 7.181(6.896)</td>
<td>159.43</td>
<td>64</td>
<td>8</td>
</tr>
<tr>
<td>RuP</td>
<td>a = 5.535(5.52), b = 3.968(3.859), c = 7.181(6.896)</td>
<td>110.01</td>
<td>216</td>
<td>4</td>
</tr>
<tr>
<td>RuP</td>
<td>a = 5.140(5.117), b = 4.526(3.903), c = 3.923(2.871)</td>
<td>91.26</td>
<td>216</td>
<td>4</td>
</tr>
<tr>
<td>Ru₂B₆</td>
<td>a = 11.139</td>
<td>362.68</td>
<td>216</td>
<td>23</td>
</tr>
<tr>
<td>Ru₄B₅</td>
<td>a = 9.17, c = 4.59</td>
<td>195.19</td>
<td>105</td>
<td>12</td>
</tr>
<tr>
<td>Ru₂B₃</td>
<td>a = 7.519, c = 4.749, γ = 120</td>
<td>233.65</td>
<td>216</td>
<td>14</td>
</tr>
<tr>
<td>RuB₅</td>
<td>a = 11.673, b = 11.940, c = 2.886</td>
<td>399.45</td>
<td>27</td>
<td>22</td>
</tr>
<tr>
<td>RuB₃</td>
<td>a = 2.877, c = 2.867, γ = 120</td>
<td>81.10</td>
<td>216</td>
<td>4</td>
</tr>
<tr>
<td>RuB₅</td>
<td>a = 2.924, c = 12.83, γ = 120</td>
<td>95.70</td>
<td>216</td>
<td>4</td>
</tr>
<tr>
<td>RuB₅</td>
<td>a = 4.675, b = 2.879, c = 4.051</td>
<td>81.10</td>
<td>216</td>
<td>2</td>
</tr>
</tbody>
</table>

Table 1: Equilibrium lattice parameters for crystalline Ru(P) and Ru(B) alloys obtained from GGA calculations.

**Fig. 1.** Variations in total energy per atom for amorphous and ordered Ru(P) and Ru(B) alloys as a function of P(B) content (at.%). For each amorphous structure, the reported average value was obtained from three independent 74-atom supercell calculations.

**Fig. 2.** Variation in volume for α-Ru(P) and α-Ru(B) alloys as a function of P(B) content (at.%). For each amorphous alloy, the reported average value was obtained from three independent 64-atom supercell calculations.
calculation results. These results demonstrate that the amorphization of binary alloys is mainly driven by the elastic strain contribution arising from the atomic size difference between Ru and P (B) atoms.

Fig. 3 shows a variation in the mixing enthalpy for amorphous and crystalline Ru(P) and Ru(B) alloys as a function of the Ru:P(B) composition ratio, with respect to crystalline Ru (c-Ru) and crystalline P or B (c-P or c-B). The mixing enthalpy per atom (ΔE_{mix}) is given by:

\[ ΔE_{mix} = E(\text{alloy}) - (1-x)E_{\text{Ru}} - xE_{\text{P(B)}} \]

where \( E(\text{alloy}) \) is the total energy per atom of the Ru(P) or Ru(B) alloy examined, \( x \) is the number fraction of P(B), and \( E_{\text{Ru}} \) and \( E_{\text{P(B)}} \) are the total energies per atom of crystalline hcp Ru and P(B) (c-Ru, c-P or c-B), respectively. The results indicate that the Ru(P) alloy forms the most favorable structure when the P content is around 40 at.% P, crystalline phases than their amorphous counterparts; hence \( fi \) 20 at.% P. This was observed experimentally in previous studies that can easily be incorporated into the amorphous Ru matrix above Ru(P) alloy, the large negative mixing enthalpies suggest that P atom exhibits a minimum value of \( ~1 \) at.% P. For the Ru(P) alloy, the large negative mixing enthalpies suggest that P atom can easily be incorporated into the amorphous Ru matrix above 20 at.% P. This was observed experimentally in previous studies that showed amorphous Ru(P) thin films formed above approximately 17 at.% P [14]. We also note that the mixing enthalpy becomes positive for low P content, which might indicate the presence of a barrier for incorporation of P into pure Ru.

On average, for Ru(P) the total energies are \(-0.3\) eV/atom lower for crystalline phases than their amorphous counterparts; hence α-Ru(P) alloys may be expected to undergo crystallization at elevated temperatures. Experiments showed 3 to 5 nm-thick films remained amorphous upon annealing at 400 °C, but began to crystallize under a 450 °C 3-h anneal [14]. The Ru(B) alloy mixing enthalpy is lower for the crystalline phase than the amorphous phase between 20 and 50 at.% B (Fig. 3). The thermal stability of the amorphous Ru(B) phase is reported herein and films with 13 to 20 at.% B remained amorphous at 450 °C and formed polycrystalline films at 500 °C.

Using this thermodynamic perspective, the amorphous structure forming ability is predicted by the comparison of the mixing enthalpy and the mixing enthalpy difference between crystalline and amorphous phases. The lower the mixing enthalpy, the higher the glass forming ability, and the lower the mixing enthalpy difference between the crystalline and amorphous phases, the higher the glass forming ability. Based on this correlation, the lower mixing enthalpy and lower mixing enthalpy difference for the Ru(P) alloy than the Ru(B) alloy may enhance its glass forming ability.

To gain understanding of the structural stability between constituent atoms, we explored the atomic distribution of selected Ru80P20 and Ru87B13 alloys as these were near the transition where amorphous alloys should form. Fig. 4 shows a set of the pair distribution functions from our AIMD simulations, together with corresponding crystalline structures for comparison. The amorphous structures were characterized using the pair distribution function (PDF), \( g(r) \), which is defined as [37]

\[ g(r) = \frac{V}{4\pi r^2} \frac{n(r)}{N} \]

where \( n(r) \) represents particles in a shell within the region \( r \pm \Delta r/2 \), where \( \Delta r \) is the shell thickness, and \( N \) denotes the number of particles in the model volume \( V \).

The pair distribution function \( g(r) \) was computed using 3 different 144-atom supercells for good statistics. No sharp second-neighbor peak is present, which confirms the amorphous nature (i.e., a lack of long-range order) of the Ru(P) and Ru(B) alloys. The first peak in \( g_{\text{Ru-Ru}} \) of a-Ru80P20 is around 2.7 Å, which nearly corresponds to the bond distance of amorphous Ru (2.7 Å), while the first peak of \( g_{\text{Ru-Ru}} \) in a-Ru87B13 is 2.6 Å, which indicates that the a-Ru87B13 is more closely packed than a-Ru80P20 even at lower B content than P content.

For the pair distribution functions for P(B), the statistics for the \( g_{\text{P-P}} \) and \( g_{\text{B-B}} \) are not good comparing with the crystalline counterparts with coordination number (CN) = 0, but the small intensity of a partial peak and a calculated coordination number of 0.85 (at cut-off of 3.03 Å) for P and B, respectively, describe the existence of dimers (P₂ and B₂). The pair distribution functions between Ru and P (B), \( g_{\text{Ru-P}} \) and \( g_{\text{Ru-B}} \), are also shown in Fig. 4. The first peaks in a-Ru80P20 and a-Ru87B13 are at 2.4 Å and 2.3 Å, respectively. This suggests the strong intermixing between the Ru and P (B) atoms, namely, a strong chemical short-range order. This result is similar to the model proposed by Lee and Hwang [38] for the Au-Si alloy, which shows strong short-range order with the negative mixing enthalpy. For the total pair distribution function, the a-Ru80P20 and a-Ru87B13 alloys show distinct amorphous character as an evidenced by the split second peak, where the average coordination number of Ru around P and B are 9.10 and 8.74, respectively.

From these \( g(r) \) features we can estimate that the structures of the amorphous alloys are governed by a chemical ordering effect due to the chemical interaction between the Ru and the P (B) atoms. More insight into the structural changes is gained by analyzing the structures and characterizing the local environment surrounding a P (B) atom by Ru atoms. As displayed in Table 2, we calculated the first-neighbor CN of the Ru atoms around P and B at selected Ru80P20 and Ru87B13 alloys as a function of normalized cutoff radius, \( r^* \). With increasing \( r^* \), the average CN increases. These results also show that pure Ru is less closely packed than the alloys. This is attributed to the relatively smaller size of P or B compared to Ru. The CN of Ru80P20 is somewhat lower than that of Ru87B13 within the nearest neighbor distance \( r^* < 1.2 \). However, when \( r^* \) is large (>1.3), the CN of the Ru80P20 is somewhat greater than that of the Ru87B13. The radius of B is smaller than that of P; therefore the average CN of Ru around the B atoms within the nearest neighbor distance is lower than that of Ru around the P atoms.
3.2. Voronoi analysis

The local atomic ordering of Ru(P) and Ru(B) were examined using the Voronoi tessellation method [39,40]. The Ru(P) or Ru(B) alloy with a moderate P or B content results in a glassy structure exhibiting a distinct topological and chemical short-range order (SRO). The type of coordination polyhedron around a P(B) atom can be specified using the Voronoi index \(\langle i_3, i_4, i_5, i_6, \ldots \rangle\), where \(i_n\) indicates the number of \(n\)-edged faces of the Voronoi polyhedron and \(\Sigma i_n\) is the total CN, to designate and differentiate the type of coordination polyhedron surrounding the center of solute atoms. For the \(\alpha\)-Ru80P20 structure (Fig. 5a), the solute coordination polyhedra form the TTP (tri-capped trigonal prism packing), which corresponds to a Voronoi index of \(\langle 0,3,6,0 \rangle\); mono-capped square Archimedean antiprism (slightly distorted from the TTP), which corresponds to a Voronoi index of \(\langle 0,5,4,0 \rangle\); and CN10 polyhedra, which has a Voronoi index of \(\langle 0,4,6,0 \rangle\). For a \(\alpha\)-Ru87B13 structure (Fig. 5b), the solute coordination polyhedra form the CN8 Kasper polyhedron, with a Voronoi index of \(\langle 0,4,4,0 \rangle\); and CN9 TTP, which corresponds to a Voronoi index of \(\langle 0,3,6,0 \rangle\). The dominant polyhedra of the solute in alloys are shown in Table 3. Voronoi analysis indicates that the average CN is 9.3 for the Ru(P) alloy, and 8.4 for the Ru(B) alloy, which corresponds to

---

**Table 2**

Average coordination number of P and B atoms as a function of cutoff radius. Here, the cut-off radius \(r^*\) is normalized with respect to 2.4 Å, which was the average nearest neighbor Ru-P distance in the Ru(P) alloy considered.

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Basis atom</th>
<th>(r^* = 1.1)</th>
<th>1.2</th>
<th>1.3</th>
<th>1.4</th>
<th>1.5</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\alpha)-Ru</td>
<td>Ru</td>
<td>5.0</td>
<td>8.6</td>
<td>10.9</td>
<td>12.2</td>
<td>12.5</td>
</tr>
<tr>
<td>(\alpha)-Ru80P20</td>
<td>P</td>
<td>3.6</td>
<td>9.1</td>
<td>11.8</td>
<td>12.7</td>
<td>13.5</td>
</tr>
<tr>
<td>(\alpha)-Ru87B13</td>
<td>B</td>
<td>4.3</td>
<td>9.4</td>
<td>10.9</td>
<td>12.1</td>
<td>13.2</td>
</tr>
</tbody>
</table>

---

**Fig. 4.** Pair distribution functions \(g(r)\) for (a) \(\alpha\)-Ru80P20 and (b) \(\alpha\)-Ru87B13 alloys. For each plot, the average CN with corresponding cut-off radius in parenthesis is shown.
the average CN calculated from integration of the first peak of the pair distribution function (PDF).

It is well known that the preference polyhedra and CN are governed by the effective atomic size ratio, \( \lambda \), between solvent and solute atoms. For instance, an earlier study shows that with decreasing \( \lambda \), the preferred polyhedra changes from the Frank-Kasper type (for \( \lambda \approx 1 \)) to the icosahedral type (for \( \lambda \approx 0.732 \)). Considering the dominant CN of 9, we can expect that the polyhedron with a Voronoi index of \( \langle 0,3,6,0 \rangle \), TTP phase found in the Ru(B) alloy, which has the CN of 8, is also a part of polyhedra found in NiB alloys. While no simulation study has been reported for Ru(P) and Ru(B), we also calculated the Ni$_{80}$P$_{20}$ and Ni$_{80}$B$_{20}$ as part of the verification of Ru(P) and Ru(B). The dominant polyhedron found in the Ni$_{80}$P$_{20}$ and Ni$_{80}$B$_{20}$ alloys are \( \langle 0,2,8,0 \rangle \), \( \langle 0,4,6,0 \rangle \), \( \langle 0,2,8,1 \rangle \), and \( \langle 0,3,6,0 \rangle \), respectively. In addition, the average CN of the P and B atoms is 10.5 and 9.1 for Ni(P) and Ni(B), respectively. These are very similar to previous findings, that show the dominant polyhedra in Ni$_{80}$P$_{20}$ and Ni$_{80}$B$_{20}$ are \( \langle 0,2,8,0 \rangle \) and \( \langle 0,3,6,0 \rangle \), respectively. Therefore, it is evident that our approach to elucidate the SRO and CN for Ru(P) and Ru(B) is reasonable in predicting the glasses properties as well.

As is shown in Fig. 5, the formation of ‘quasi-equivalent’ P-centered Ru clusters arising from topological and chemical SRO is also likely to lead to the medium-range order (MRO) in the binary alloy. In fact, the short-to-medium range order is seen in other metallic glasses, particularly in transition metal–metalloid and transition metal–transition metal systems where the chemical short-range-order is significant. In Ni$_{80}$P$_{20}$, the P atom-centered clusters (Fig. 5c) are packed with the icosahedral order, which is very similar to the topological configuration with the AIMD simulation. In the same manner, Ru(P) has shown the icosahedral type regardless of the type of SRO. These results indicate that the MRO found in the metal-metalloid binary alloy has the icosahedral ordering, which has the most stable packing in metallic glasses.

### 3.3. Film growth using \( \text{B}_2\text{H}_6 \) as the B source in Ru(B) CVD

As with the \( \text{PH}_3 \) hydride used in Ru(P) film growth, \( \text{B}_2\text{H}_6 \) appears to decompose rapidly and completely on Ru and Ru(B) under the film growth conditions at 250 °C. This was expected since \( \text{B}_2\text{H}_6 \) decomposes on Ru(0001) between 230 and 400 K depending on the \( \text{B}_2\text{H}_6 \) exposure level. Unexpectedly, much higher B incorporation levels were found in the films grown with \( \text{B}_2\text{H}_6 \) when compared to the \( \text{P} \) incorporation levels found with the same \( \text{PH}_3 \) partial pressure. In practical terms the high reactivity of \( \text{B}_2\text{H}_6 \) on Ru presents challenges to limit the B incorporation levels during film growth. The 15 ppm \( \text{B}_2\text{H}_6 \) in \( \text{H}_2 \) concentration used herein was the lowest concentration commonly sold by the manufacturer. Control of film composition also relied on \( \text{B}_2\text{H}_6 \) source gas flow rate adjustments during growth. Other CVD chambers will need to establish protocols for limiting and regulating the \( \text{B}_2\text{H}_6 \) partial pressure to very low levels.

In another similarity to Ru(P) films grown with \( \text{PH}_3 \), \( \text{B} \) accumulates on the growth surface more rapidly than it can be incorporated by the growing Ru(B) film, resulting in a steeply increasing B concentration within the film. The Ru(B)/SiO$_2$ starting interface to the growth surface if the \( \text{B}_2\text{H}_6 \) partial pressure is kept static during growth. To keep the film composition uniform with thickness the \( \text{B}_2\text{H}_6 \) partial pressure in the CVD chamber was varied over the course of the growth cycle for each film, beginning high and then gradually being stepped down to zero over the course of film growth (shown in Fig. 6 for a film grown at 250 °C). Fig. 7 presents the concentration profile corresponding to the pressure settings in Fig. 6 with the position based on sputtering times as described in Section 2. There is <5 at.% variation in the calculated B content of the film over 3 nm of total film thickness. The specifics of the concentration step sizes and times were based on previous Ru(P) CVD studies with \( \text{PH}_3 \) and were found to produce a low variation in the mean B concentration with position relative to the Ru(B)/SiO$_2$ interface. Overall film composition is reported using a mean B concentration.

Films were grown at substrate temperatures of 250 °C, 300 °C, and 350 °C, and no dependence in B the concentration profile was observed within this temperature range. This was expected since \( \text{B}_2\text{H}_6 \) decomposes on Ru at a much lower temperature than the lower limits of the

---

**Table 3** The dominant polyhedra of the solute atoms in the metallic alloy. The values are the fraction of solute atoms in the metallic alloys and only the polyhedra relevant to our work are shown.

<table>
<thead>
<tr>
<th>Voronoi index</th>
<th>( \text{Ru}<em>{80}\text{P}</em>{20} )</th>
<th>( \text{Ru}<em>{80}\text{B}</em>{13} )</th>
<th>( \text{Ni}<em>{80}\text{P}</em>{20} )</th>
<th>( \text{Ni}<em>{80}\text{B}</em>{20} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \langle 0,4,4,0 \rangle )</td>
<td>0.05</td>
<td>0.6</td>
<td>–</td>
<td>0.21</td>
</tr>
<tr>
<td>( \langle 0,3,6,0 \rangle )</td>
<td>0.24</td>
<td>0.4</td>
<td>–</td>
<td>0.43</td>
</tr>
<tr>
<td>( \langle 0,5,4,0 \rangle )</td>
<td>0.24</td>
<td>–</td>
<td>–</td>
<td>0.07</td>
</tr>
<tr>
<td>( \langle 0,2,8,0 \rangle )</td>
<td>–</td>
<td>0.7</td>
<td>–</td>
<td>0.21</td>
</tr>
<tr>
<td>( \langle 0,4,6,0 \rangle )</td>
<td>–</td>
<td>0.1</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>( \langle 0,2,8,1 \rangle )</td>
<td>–</td>
<td>0.1</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

---

**Fig. 6.** Partial pressure of diborane versus growth time for a film grown at 250 °C.
atomic force microscopy was used to measure the roughness of various 3 nm-thick amorphous Ru(B) films (not shown). RMS roughness of continuous films was approximately 0.1 nm, ±0.05 nm on average, which is similar to amorphous Ru(P) films of comparable thickness [14]. The AFM technique was primarily used to check film continuity. Attempts to produce films <3 nm in thickness resulted in discontinuous films with large RMS roughnesses indicating that the films were not continuous.

Temperature stability of the Ru(B) amorphous films was evaluated by alternating 3-h anneals of the Ru(B) films with X-ray diffraction analysis. Four films were tested, all 3-nm Ru(B) films were grown at 250 °C, one with 13 at.% B, two with 15 at.% B, and one with 20 at.% B. Each film was amorphous as-grown, and checked for crystallization using XRD following a 300 °C anneal, then a 400, 450, and 500 °C anneal in sequence. All films remained amorphous through the first three anneal cycles, and all films exhibited diffraction features associated with the Ru crystal structure following the 500 °C anneal as illustrated in Fig. 9 for the Ru(P) film. This is similar to the crystallization of 3 nm Ru(P) films, which were found to occur at 450 °C following a similar anneal sequence [14].

3.4. Barrier performance of Ru(B) and comparison to amorphous Ru(P)

The electrical failure test described in Experimental Methods was performed on several 3 nm Ru(B) films, to provide a direct comparison to the performance of previously-studied Ru(P) ultrathin films and a 5-nm Ta/TaN stack produced using in-house PVD in the study in reference [7]. Further discussion of the method and previous related work can be found in several of the references [7,14,47]. Three films were tested: a 15-at.% B Ru(B) film a similar 15-at.% B Ru(B) film plated with an additional 2 nm of PVD Ru to bring its overall thickness up to the 5 nm to allow comparison with Ru(P) results [7,14], and a 3-nm 25 at.% B Ru(B) film to provide an idea of whether increasing B content beyond the point where Ru(B) grows as an amorphous film provides any benefit to the possible Cu diffusion barrier application.

Fig. 10 shows the measured times to failure for the 3-nm 15-at.% B and 25-at.% B Ru(B) films. Fig. 11 contrasts the average failure times for the 15-at.% B Ru(B) film to those of a 3-nm amorphous Ru(P) film and a 5-nm CVD Ru film grown without alloying elements [14], and a 5-nm TaN film grown in situ for benchmarking in previous work [7]. Both amorphous Ru alloy films perform significantly better than pure CVD Ru despite lower thickness, indicating that amorphous phases mitigate the issue of grain boundary diffusion as predicted and demonstrating that both B and P alloys with Ru can serve in the role of Cu diffusion barrier/liner layer in interconnect devices.

Atomically resolved Cu diffraction and STEM imaging in addition to XPS and GIXD analysis can be performed on several 3 nm Ru(B) films, which were found to occur at 450 °C following a similar anneal sequence [14].

The Ru(B) films were grown at 250 °C, one with 13 at.% B, two with 15 at.% B, and one with 20 at.% B. Each film was amorphous as-grown, and checked for crystallization using XRD following a 300 °C anneal, then a 400, 450, and 500 °C anneal in sequence. All films remained amorphous through the first three anneal cycles, and all films exhibited diffraction features associated with the Ru crystal structure following the 500 °C anneal as illustrated in Fig. 9 for the Ru(P) film. This is similar to the crystallization of 3 nm Ru(P) films, which were found to occur at 450 °C following a similar anneal sequence [14].

3.4. Barrier performance of Ru(B) and comparison to amorphous Ru(P)

The electrical failure test described in Experimental Methods was performed on several 3 nm Ru(B) films, to provide a direct comparison to the performance of previously-studied Ru(P) ultrathin films and a 5-nm Ta/TaN stack produced using in-house PVD in the study in reference [7]. Further discussion of the method and previous related work can be found in several of the references [7,14,47]. Three films were tested: a 15-at.% B Ru(B) film a similar 15-at.% B Ru(B) film plated with an additional 2 nm of PVD Ru to bring its overall thickness up to the 5 nm to allow comparison with Ru(P) results [7,14], and a 3-nm 25 at.% B Ru(B) film to provide an idea of whether increasing B content beyond the point where Ru(B) grows as an amorphous film provides any benefit to the possible Cu diffusion barrier application.

Fig. 10 shows the measured times to failure for the 3-nm 15-at.% B and 25-at.% B Ru(B) films. Fig. 11 contrasts the average failure times for the 15-at.% B Ru(B) film to those of a 3-nm amorphous Ru(P) film and a 5-nm CVD Ru film grown without alloying elements [14], and a 5-nm TaN film grown in situ for benchmarking in previous work [7]. Both amorphous Ru alloy films perform significantly better than pure CVD Ru despite lower thickness, indicating that amorphous phases mitigate the issue of grain boundary diffusion as predicted and demonstrating that both B and P alloys with Ru can serve in the role of Cu diffusion barrier/liner layer in interconnect devices.
The ab initio molecular dynamics (AIMD) study shows that Ru(P) and Ru(B) alloys with moderate P (B) content can result in glassy structure exhibiting the topological and strong chemical short-to-medium range order. Amorphous phases above ~20 at.% of P and ~10 at.% of B are found to be energetically more favorable than the crystalline counterparts for Ru(P) and Ru(B) alloys, respectively. In the Ru$_8$P$_2$O$_2$ structure, the P-centered polyhedra prefer the TTP phase with Voronoi index (0,3,6,0), while in the Ru$_8$B$_{13}$, the B-centered polyhedra prefer the (0,4,4,0). In addition, the Ru-P and Ru-B systems show the icosahedral medium range ordering arising from packing the “quasi-equivalent” P-centered (B-centered) clusters in three dimensional spaces.

4. Summary and conclusions

Growth studies employed B$_2$H$_6$ to grow α-Ru(B) films with CVD and showed that it should perform well as an ultrathin Cu diffusion barrier. The films were amorphous above 10 at.% B, consistent with the first principles predictions. Film resistivity is <120 μΩ·cm and approaches that of single-crystalline pure Ru at similar thicknesses, and C contamination is consistently low. Consistent continuity and smoothness at 3 nm thickness suggest that the material is amenable to conformal deposition. Performance under failure testing shows that α-Ru(B) serves as a Cu diffusion barrier, while stability under annealing studies to 450 °C indicates that α-Ru(B) is stable under most processing conditions. Time-to-failure performance under field stress equals or exceeds previously studied α-Ru(P) amorphous films and exceed the measured time to failure of 5 nm films of pure Ru by 4–5 orders of magnitude. Overall, α-Ru(B) amorphous films produced using the hydride precursor appear promising as a liner material in technology node sizes requiring liners at <5 nm thickness.

Acknowledgements

This work was supported by the National Science Foundation (CBET-0854345 and CBET-1160195).

References

[41] P. Lamparter, Reverse Monte Carlo simulations of amorphous Ni$_{50}$P$_{50}$ and Ni$_{60}$P$_{40}$, Phys. Scr. T57 (1995) 72–78.