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Author(s)	Kohama, Kazuyuki; Ito, Kazuhiro
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Synthesis Using Cu(Ti) Alloy Films for Self-Forming Functionality in Electronic Devices[†]

ITO Kazuhiro*, KOHAMA Kazuyuki**

Abstract

Cu interconnects have been used extensively in ULSI devices. However, large resistance-capacitance delay and poor device reliability have been critical issues as the device feature size has reduced to nanometer scale. In order to achieve low resistance and high reliability of Cu interconnects, we have proposed a process forming a thin Ti-based self-formed barrier (SFB) after annealing a Cu(Ti) alloy film deposited on dielectrics at elevated temperatures. Identification of Ti-based SFB was conducted using the electron diffraction, X-ray photoelectron spectroscopy, and the Rutherford backscattering spectrometry techniques. Those indicate that the Ti-based SFB consists of mainly amorphous Ti oxides for any kinds of dielectrics. Small amount of crystalline Ti compounds such as TiC, TiN, TiSi was dependent on a kind of dielectrics. The Ti-based SFB growth was concluded to be controlled by a thermally activated process. The activation energy and the pre-exponential factor values decreased with decreasing C concentration in dielectrics, suggesting that the growth is controlled by a chemical reactions of Ti atoms with dielectrics. The process was applied to 45 nm-node dual damascene interconnects and evaluated its performance. The microstructure analysis by transmission electron microscope and energy dispersive X-ray fluorescence spectrometer showed the 2 nm-thick Ti-based SFB at the interface between a Cu wire and low-k dielectrics. The line resistance and via resistance significantly decreased compared with those when using the conventional Ta/TaN barrier. The stress migration performance was also drastically improved using the SFB process. A performance of time dependent dielectric breakdown revealed superior endurance. These results suggest that the Ti-based SFB process is one of the most promising candidates for advanced Cu interconnects. This process using Ti as an alloy element can be applied for Cu-interconnect formation in electronic devices such as TFT-LCDs.

KEY WORDS: (Cu(Ti) alloy films), (Cu interconnects), (Resistivity), (Diffusion barrier), (Adhesion)

1. Introduction

Ultra large-scale integration (ULSI) technology contribute significantly to our electronic information society. ULSI consists of transistors as the switching devices and interconnects that connect between switches. Scaling down of transistors significantly contributes to device performance. For example, electron goes from source to drain, and the gate voltage controlled the electron flow due to formation of channel under the gate. Scaling down the transistor size by, for example, one and a half, both the voltage and gate delay similarly scaling down, and the power dissipation becomes one forth. Then device feature size has reduced from micron-scale era into nano-scale era. On the contrary, interconnect delay increased with decreasing gate size, although the gate delay decreased. This results that total device delay reaches the minimum at a critical gate size, suggesting further scaling down makes it increase. Thus, we need

further improvement in Cu interconnects in a view point of metallurgy, with device future size has reduced to smaller than 50 nm.

Cu interconnects form layer structures and Cu easily diffuses into dielectrics by which Cu interconnect is surrounded. Thus, refractory metals and compounds such as Ta and TaN are deposited between Cu and dielectrics as a barrier. Cu dry etching is somehow difficult, and thus a dual damascene method is

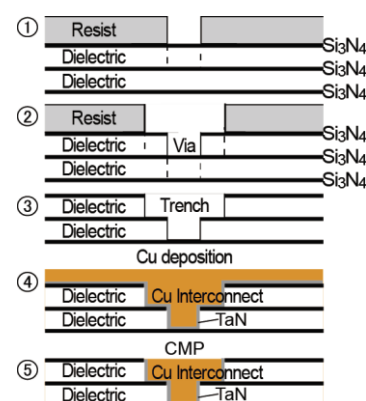


Fig. 1 Schematic illustration of dual damascene method.

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* Professor

** Assistant Professor

used for making Cu interconnects, as shown in **Fig. 1**. Digging a groove so-called via and trench is the first, and then Cu film is electrochemically deposited, and finally chemical-mechanical polishing was conducted.

In fabrication of Cu interconnects, large resistance-capacitance (RC) delay and poor device reliability have been critical issues as the device feature size has reduced to nanometer scale¹⁾. In the RC delay, resistivity increase due to barrier layers is becoming significant with the reduction in line width of the Cu interconnects. Moriyama et al.²⁾ and Shimada et al.³⁾ reported that significant resistivity increase is observed when the line width decrease to smaller than 100 nm and a very thin barrier layer (< 5 nm) is required for the interconnects with a line-width of ~ 45 nm and an average grain size of ~ 450 nm to achieve an effective interconnect resistivity of less than $4 \mu\Omega\text{-cm}$ (**Fig. 2**). Thus, our goal is how to make a thin barrier and large-grained Cu interconnects.

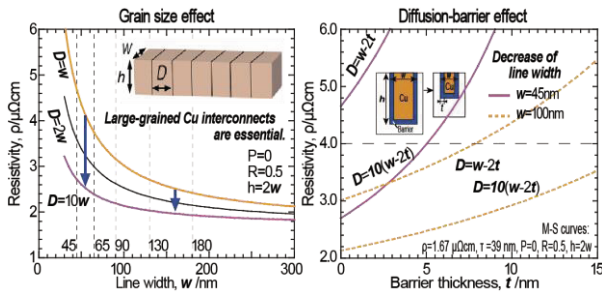


Fig. 2 Theoretical resistivities as a function of grain sizes and effective resistivities as a function of barrier thickness calculated using the MS model with $P=0$ and $R=0.5$ for the Cu interconnects³⁾.

Various methods have been reported to reduce the volume of barrier layers in Cu interconnects by fabricating an ultra thin barrier⁴⁻¹²⁾. Formation of a thin barrier layer by annealing Cu alloy seeds at elevated temperatures is conventionally called a self-formed barrier (SFB) technique. The Cu alloy films with various solutes (Mg, Ti, Al, Cr, etc.) were extensively investigated for the possibility of the self-formation of the passivation layers¹³⁻²¹⁾, and those with Mn^{7,8)} and Ti for SFBs⁹⁻¹²⁾. The SFB technique is essential for reaction of solute atoms with dielectrics as shown in **Fig. 3**. The Ti

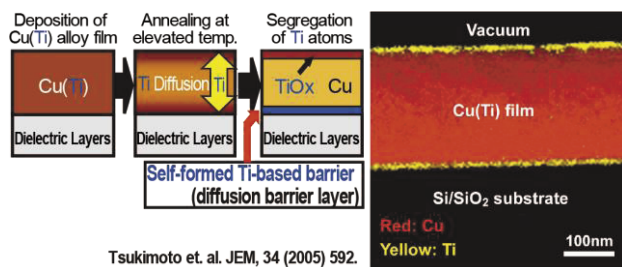


Fig. 3 Schematic illustrations of Ti-based self-formed barrier (SFB) synthesis using Cu(Ti) alloy films annealed at elevated temperature and its TEM/EDS mapping result⁹⁾.

atoms in the Cu(Ti) alloy seeds can react with many kinds of dielectrics such as SiO_2 , SiOC , SiCO , and SiCN , resulting in formation of SFB with different composition¹¹⁾. However, the technique using a Cu(Ti) alloy seed has not been applied to dual-damascene interconnects. In this paper, we firstly show characterization of Ti-based SFB using transmission electron microscope (TEM)/energy dispersive X-ray spectroscopy (EDS), X-ray photoelectron spectroscopy (XPS), and Rutherford backscattering spectrometry (RBS), and examples applied Ti-based SFB to the Cu dual-damascene interconnect for 45-nm node of ULSI.

2. Ti-based SFB characterization

2.1 Identification of Ti-based SFB using XPS

TEM/selected area diffraction (SAD) observation showed that the Ti-based SFB consisted of crystalline Ti compounds (TiC , TiSi or TiN)⁹⁻¹²⁾. **Figure 4** shows cross-sectional TEM images and SAD images taken from an area marked with a broken circle in the TEM images for an annealed Cu(Ti)/ SiO_2 (**Fig. 4(a)**) and Cu(Ti)/Low-k1 (**Fig. 4(c)**) samples. Thin Ti-based SFB layers formed at the interfaces between a Cu(Ti) alloy film and dielectrics. They consisted of polycrystalline TiSi on SiO_2 (**Fig. 4(b)**) and TiC on Low-k1 (**Fig. 4(d)**). The kinds of the Ti compounds formed in the Ti-based SFB depended on the kinds of dielectrics. The thickness, dielectric constants, and compositions of the dielectrics are shown in Table 1. The TEM/SAD observation and SIMS analyses suggested that the Ti-based SFB consisted of amorphous Ti oxides (TiO_x) as well as the crystalline Ti compounds.

Although the barrier structure and amorphous phases in the barrier were not directly identified by the TEM/SAD observation and SIMS analyses, an XPS

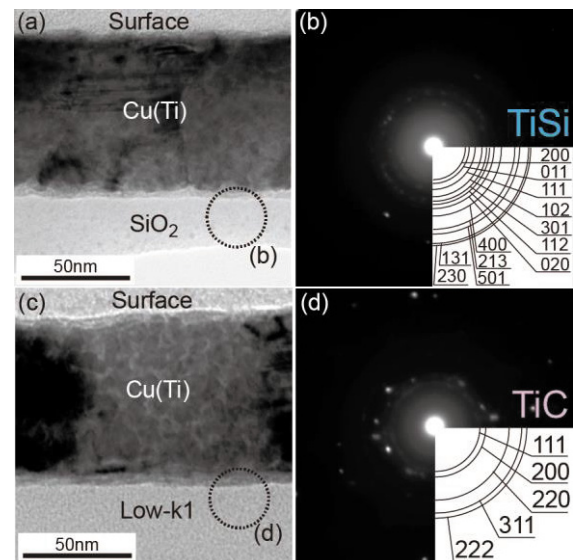


Fig. 4 Cross-sectional TEM images of the (a) Cu(Ti)/ SiO_2 and (c) Cu(Ti)/Low-k1 after annealing at 600°C for 2h in Ar, respectively. (b) and (d) are SAD images taken from the areas marked with broken circles in (a) and (c), respectively.

Table 1 Dielectric constants (k) and thickness of dielectric layers and their C, O, Si and N concentration (at.%)¹²⁾.

Dielectrics	k	t (nm)	C (at.%)	O (at.%)	Si (at.%)	N (at.%)
th-SiO ₂	3.9	100	-	66.7	33.3	-
Low-k1	3.0	440	17.0	24.9	18.8	-
Low-k4	2.6	450	~14	~29	~18	-
SiCN	4.8	200	21.4	0.5	25.0	12.8

technique with simultaneous Ar etching was employed to investigate the structure of the Ti-based SFB and its compositions systematically with dielectrics. **Figure 5(a)** shows typical elemental depth profiles of the Cu(Ti)/SiO₂ sample after annealing in ultrahigh vacuum (UHV) at 600°C for 2h obtained by simultaneous Ar etching and XPS measurements. Ti segregation was observed around etch times between 18000s and 25000s (at the interface), where oxygen and Si atoms were also detected. To identify composition of the Ti-based SFB, XPS measurements around the binding energies of Ti 2p, C 1s, Si 2p, and N 1s were performed. Portions of the XPS profiles around the binding energies of Ti 2p (460 eV^{22,23)}) obtained from the middle of Ti-based SFB (etch time of about 21600 s) are shown in **Fig. 5(b)**. They varied with dielectrics and were fitted by the sum of the XPS profiles of TiO/TiC, TiO₂, Ti₂O₃, and a background. Similarly, XPS profiles around the binding energies of C 1s, Si 2p, and N 1s were fitted. Based on the fitting, volume fractions of TiC, TiSi, TiN, TiO₂, Ti₂O₃, and TiO phases were determined by a ratio of each peak area to sum of peak areas.

Figure 6 shows depth dependence of composition of the Ti-based SFB in the annealed Cu(Ti)/dielectrics samples. In the Cu(Ti)/SiO₂ sample, Ti-based SFB mainly consisted of Ti oxides. The Ti oxides were in amorphous state, since crystalline Ti oxides were not detected by TEM/SAD analyses (**Fig. 4(a)**). Similarly, Ti-based SFBs in the other samples (Low-k1, Low-k4, SiCN) consisted of a large amount of amorphous Ti oxides (50~75%), as shown in **Fig. 6**. The crystalline TiC was observed in the annealed Cu(Ti)/Low-k1 and Cu(Ti)/SiCN samples, and amorphous TiC was observed

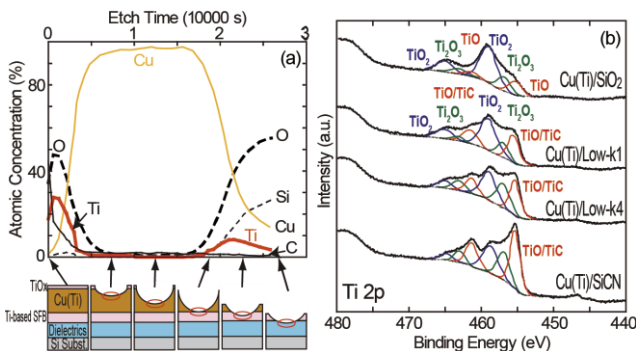


Fig. 5 (a) A XPS depth profile of the Cu(Ti)/SiO₂ sample after annealing in UHV at 600°C for 2h and (b) portions of XPS spectra around the binding energies of Ti 2 obtained from the middle of the Ti-based SFB formed in Cu(Ti)/dielectrics after annealing in UHV at 600°C for 2h¹²⁾.

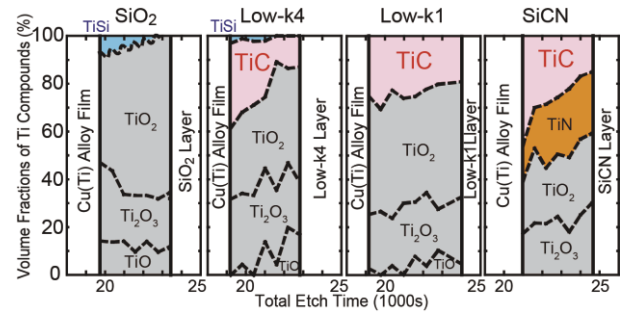


Fig. 6 Depth dependence of volume fractions of Ti compounds in the Ti-based SFB formed on the SiO₂, Low-k4, Low-k1, and SiCN layers after annealing in UHV at 600°C for 2h¹²⁾.

in the annealed Cu(Ti)/Low-k4 sample. TiN observed in the annealed Cu(Ti)/SiCN sample was in the amorphous state. The volume fractions of TiC and TiN were estimated to be about 20%, and both phases occupied a small part of the Ti-based SFB. While, a small amounts of TiSi was observed in the Cu(Ti)/SiO₂ and Cu(Ti)/Low-k4 samples, but not observed in the other samples¹²⁾. This is in good agreement with the previous study¹¹⁾. The crystalline Ti compounds such as TiC and TiSi tended to form beneath the Cu(Ti) alloy films. Some SAD patterns obtained in Cu(Ti)/Low-k1 and Cu(Ti)/SiCN samples showed that orientation relationship between crystalline TiC and Cu grains (not shown). This is one of the evidences that TiC formed beneath the Cu(Ti) alloy films. On the other hand, the amorphous phases such as TiO_x tended to form above the dielectric layers, and play an important role to prevent Cu diffusion into the dielectrics.

2.1 Ti-based SFB growth characterized by RBS

Growth of the Ti-based barrier was characterized using Rutherford backscattering spectroscopy (RBS). **Figure 7** shows RBS spectra of the annealed Cu(Ti)/low-k1 samples with various Ti contents. Ti segregation was observed at the interface in all samples

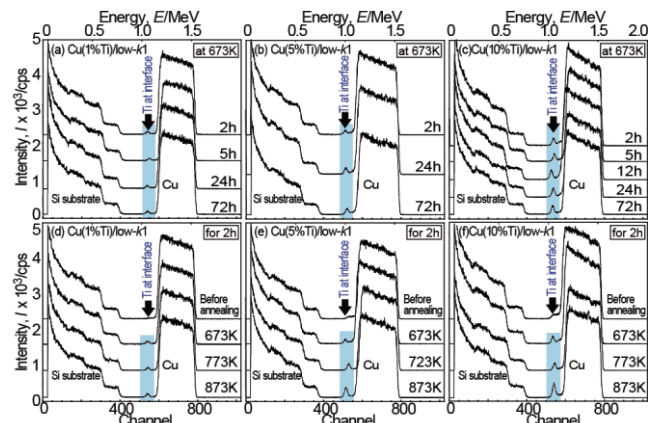


Fig. 7 RBS profiles of the Cu(Ti)/Low-k1 after annealing in UHV (a)-(c) at 400°C for various time (t) and (d)-(f) at various temperatures (T) for 2h²⁴⁾.

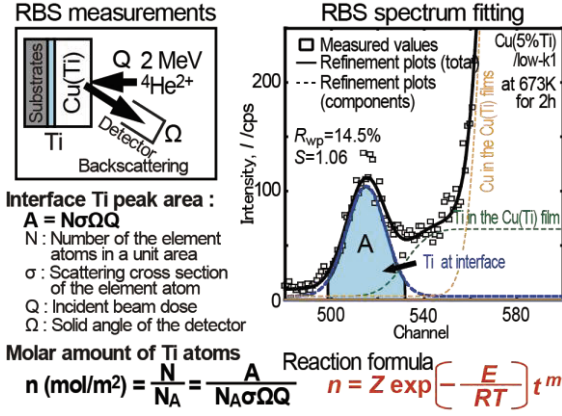


Fig. 8 RBS measurement principle and a fitting example of RBS spectrum for Cu(Ti)/Low-k1 after annealing at 400°C for 2h.

²⁴). Ti content segregated at the interface was estimated based on RBS theory. **Figure 8** shows the RBS measurement principle and an example of refinement plots placed upon the RBS spectrum of Cu(Ti)/Low-k1 after annealing at 400°C for 2h²⁴). The estimated molar amount of Ti atoms (n) is consistent with the molar amount of Ti atoms reacted with dielectrics, which follows the reaction formula

$$n = Z \exp(-E/RT) \cdot t^m, \quad (1)$$

where Z is a pre-exponential factor and E is the activation energy for the reaction.

Figure 9(a)-(c) show plots of $\log n$ versus $\log t$ for Cu(Ti)/dielectrics with the initial Ti concentrations of 1 at.%, 5 at.%, and 10 at.%, respectively, after annealing in UHV at 400°C. Those exhibit the growth behaviors of the Ti-rich interface layers. The $\log n$ values were found to be proportional to the $\log t$ values in all the samples, regardless of the initial Ti concentration in the alloy films or the kinds of dielectric layers. The m values were estimated from the slopes of the $\log n$ versus $\log t$ lines.

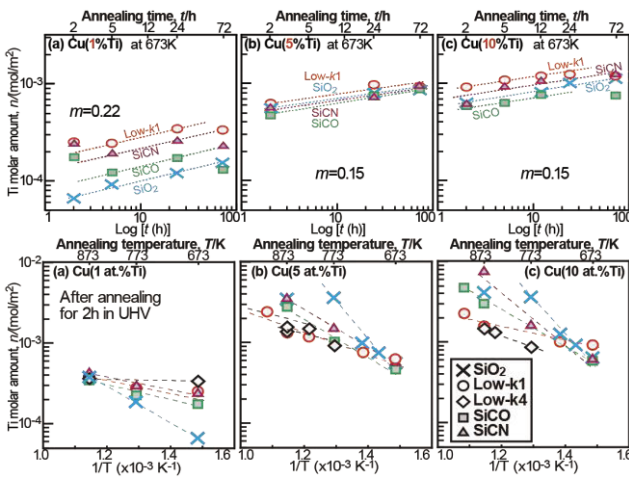


Fig. 9 (a)-(c) Plots of $\log n$ versus $\log t$ for Cu(Ti)/dielectrics after annealing in UHV at 400°C and (d)-(f) Arrhenius plots of $\log n$ versus $\log 1/T$ for Cu(Ti)/dielectrics after annealing in UHV for 2h²⁴).

They were almost similar for all the samples regardless of the kinds of dielectric layers at each initial Ti concentration in the alloy films. The m value estimated at the initial Ti concentration of 1 at.% was a little higher ($m=0.22$) than those ($m=0.15$) at 5 at.% and 10 at.%. The similar m values suggest that growth of the Ti-rich interface layers was controlled by a similar mechanism. The m values were around 0.2, lower than 0.5, suggesting that growth of the Ti-rich interface layers is not controlled by lattice diffusion but by grain-boundary and/or interface diffusion.

Figure 9(d)-(f) show Arrhenius plots of $\log n$ versus $1/T$ in Cu(Ti)/dielectrics for the initial Ti concentrations of 1 at.%, 5 at.%, and 10 at.%, respectively. These show a linear relationship, which suggests that the formation of the Ti-based SFB was controlled by a thermally-activated process. Slopes of the $\log n$ versus $1/T$ lines at the initial Ti concentration of 1 at.% were a little smaller than those at 5 at.% and 10 at.%, which were similar. This corresponds to the behaviors indicated in **Figs. 9(a)-(c)**. The initial Ti concentration at 1 at.% is insufficient to induce reaction of the Ti atoms with the dielectric layers. Activation energies, E , were estimated from the slopes of the $\log n$ versus $1/T$ lines. The E values were plotted as a function of C concentration of the dielectric layers (**Fig. 10(a)**). The E values at the initial Ti concentration of 5 at.%Ti and 10 at.%Ti were similar. The E values for the samples consisting of dielectric layers containing carbon (except SiO_2) tended to decrease with decreasing C concentration (decreasing k), and those without carbon (SiO_2) were much higher than others. This indicates that the reaction of the Ti atoms in the alloy films with the low- k layers is the most easily thermally-activated process. Also, composition of the dielectric layers is suggested to play an important role in the reaction of the Ti atoms with dielectric layers, and the carbon may be a key element to control the reaction. This is similar to the formation rule of Ti compounds (TiC or TiSi) in the Ti-rich interface layers.

The pre-exponential factors, Z , were estimated from intercepts of the slopes in the $\log n$ versus $\log t$ lines (**Figs. 9(a)-(c)**) and $\log n$ versus $1/T$ lines (**Figs. 9(d)-(f)**). The Z values estimated from two different sets of data were the same at each initial Ti concentration and dielectric layer. The Z values the Cu(Ti)/dielectrics after annealing in UHV were plotted as a function of C concentration of the dielectric layers (**Fig. 10(b)**). This is similar to the E behavior as shown in **Fig. 10(a)**. The pre-exponential factor shows the frequency with which the Ti atoms meet elemental reactants in the dielectric layers. The frequency was low in the samples with low- k , and was high in the samples with SiO_2 . In consequence, coefficients of the reaction rate ($Z \exp(-E/RT)$) were insensitive to C concentration in the dielectric layers at the initial Ti concentrations of 5 at.% and 10 at.%. These factors lead to the conclusion that growth of the Ti-based SFB is controlled by chemical reactions, represented by the Z and E values, of the Ti atoms with the dielectric layers, although there are few diffusion processes

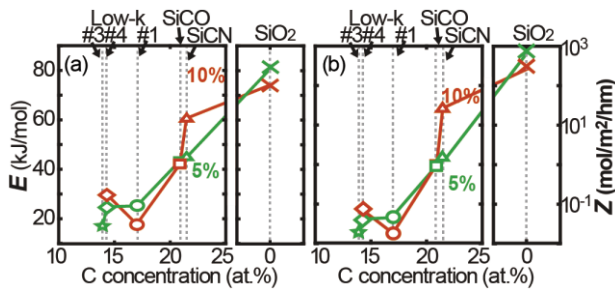


Fig. 10 (a) Activation energies of E^{24} and (b) values of Z of the Cu(Ti)/dielectrics after annealing in UHV as a function of C concentration of the dielectrics.

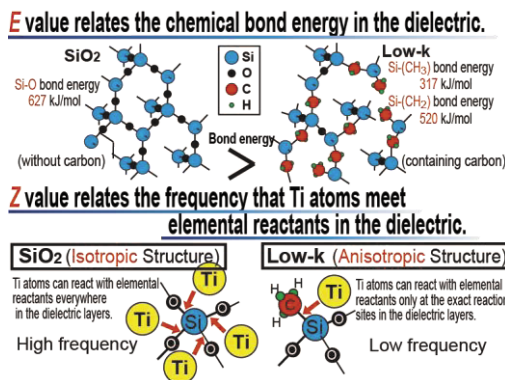


Fig. 11 Schematic illustration of the role of E and Z in reaction of Ti atom in Cu(Ti) alloy with dielectrics.

possible in all the samples since the m values suggest that it is controlled by grain-boundary and/or interface diffusion. We believe the activation energy relates the chemical bonding energy in the insulator, and Z value relates the frequency that Ti atoms meet elemental reactants such as O, C, N in the insulators, as shown in **Fig. 11**. We conclude the growth was controlled by chemical reaction of Ti atoms with the reactants in the insulators.

3. Cu dual-damascene interconnects using a Cu(Ti) alloy seed

We applied Ti-based SFB to a dual-damascene structure fabricated using 45nm-node interconnect technology with Cu and SiOCH (low-k) dielectrics. The Ti-based SFB was applied to a M2 layer between M1 and M3 layers, which were fabricated using a pure-Cu seed with a conventional Ta/TaN barrier (**Fig. 12**)²⁵. To fabricate the M2 layer, a 15nm-thick Cu(Ti) alloy film was deposited on trench and via patterning substrates using an ionized physical vapor deposition (PVD) method, and similarly a 30nm-thick pure Cu film was deposited on the Cu(Ti) alloy seed. A 600nm-thick Cu film was deposited by an electro-chemically plated (ECP) method, and followed by annealing at 450°C for 1 h in nitrogen ambient and Cu-Chemical Mechanical Polishing (CMP). In the process, a very thin 2 nm-thick Ti-based barrier successfully formed between Cu and an insulator film. Ti atoms segregated only at the barrier region, as shown in **Fig. 13**.

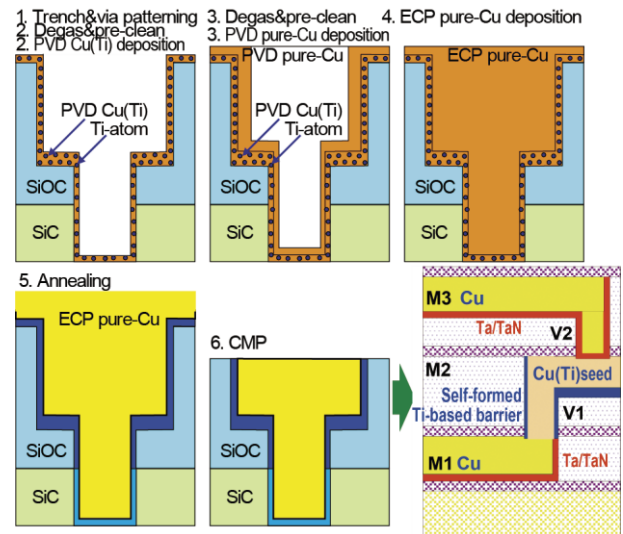


Fig. 12 Schematics of process flow for Ti-based SFB: after Cu(Ti) and pure-Cu seed deposition, after ECP-Cu deposition, after annealing, after CMP, and interconnect structure after fabrication.

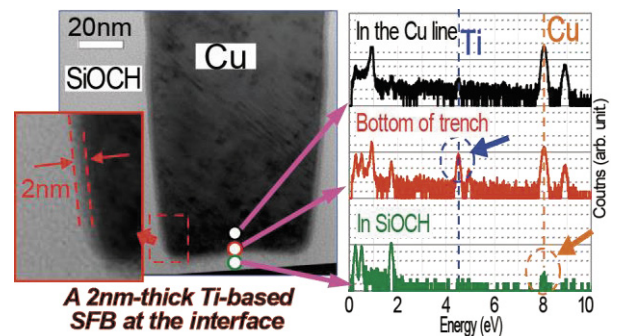


Fig. 13 Cross-sectional TEM image of trench filled with ECP-Cu using Cu(Ti) alloy seed and EDX analysis around the bottom of trench at the ECP-Cu¹⁶.

To evaluate influence of the Ti-based SFB on electrical properties of the dual-damascene interconnects, via and line resistance measurements were conducted. Distribution of via resistances using the Ti-based SFB is shown together with that using a conventional Ta/TaN barrier (**Fig. 14(a)**). The average of via resistance using the Ti-based SFB was reduced to about 30% of that using a Ta/TaN barrier, and its distribution drastically improved. The low via resistance and its narrow distribution was explained by no barrier layer formation at the bottom of via using the Ti-based SFB (**Fig. 14(b)**). In contrast, high resistance barrier formed at the bottom of via using a Ta/TaN barrier (**Fig. 14(c)**). The line resistances with a 1000nm-wide trench using the Ti-based SFB were almost similar to that using a Ta/TaN barrier (**Fig. 15(a)**), but those with 76nm-wide trench using the Ti-based SFB were reduced to about 70% of that using a Ta/TaN barrier (**Fig. 15(b)**). This indicates that line resistance is dependent on thickness of barrier layers for a narrow line, but this is not for a wide line. **Figure 15 (c) and (d)** shows cross-sectional TEM images of Cu trenches using

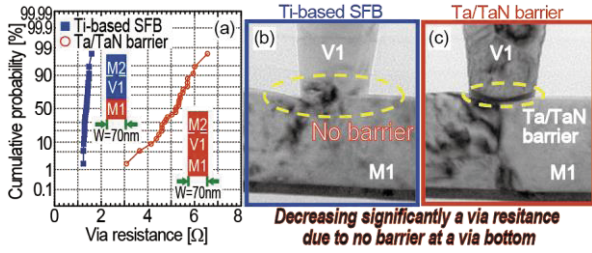


Fig. 14 (a) Cumulative via resistance distribution with Kelvin structures. Cross-sectional TEM images of via bottom using (b) the Ti-based SFB and (c) the Ta/TaN barrier²⁵⁾.

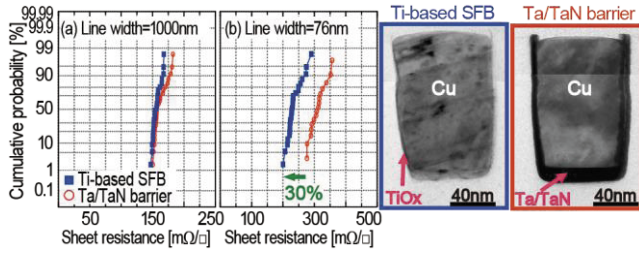


Fig. 15 Cumulative sheet resistance distribution of (a) a 1000 nm-wide line and (b) a 76 nm-wide line. Cross-sectional TEM images of trenches using (c) the Ti-based SFB and (d) the Ta/TaN barrier²⁵⁾.

the Ti-based SFB and a Ta/TaN barrier. Thickness of the Ti-based SFB was thinner than that of the Ta/TaN barrier. Also resistivity increase due to impurity Ti atoms in Cu trenches was negligible, since the Ti atoms in the Cu(Ti) seed layer were almost consumed to form the Ti-based SFB at the wall and bottom of the trenches. Thus, the thin Ti-based SFB contributed to decrease of via and line resistance of Cu dual-damascene interconnects.

Figure 16(a) shows relationship between capacitance and resistance of the Cu dual-damascene interconnects using both the Ti-based SFB and Ta/TaN barrier. There was almost no change in capacitance between them, but the resistance using the Ti-based SFB was lower than that using a Ta/TaN barrier. Breakdown voltages between Cu interconnects and time dependent dielectric breakdown (TDDB) endurance using a comb structure are shown in **Fig. 16(b)** and **(c)**, respectively. The breakdown voltages using the Ti-based SFB were a little bit higher than that using a Ta/TaN barrier, and TDDB endurance were similar. Those indicate that the Ti-based SFB has enough barrier property against Cu diffusion into dielectrics.

Figure 17 shows resistance shifts after thermal stress with two kinds of via chain patterns: a wide upper line and a wide lower line. The Ti-based SFB was applied to a wide line for both samples, and a Ta/TaN barrier was applied to other layers. Although the resistance using a Ta/TaN barrier increased after thermal stress for both samples, the resistance shift using the Ti-based SFB was very small. The stress induced voiding (SIV) endurance is well-known to be improved by taking Cu(X) (X is Ti, Al) alloy seeds. This was explained by grain boundary

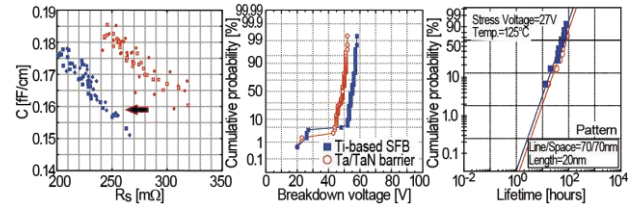


Fig. 16 (a) Relationship between capacitance and resistance of the Cu dual-damascene interconnects. Cumulative (b) breakdown voltage distribution and (c) failure distribution with a comb structure¹⁶⁾.

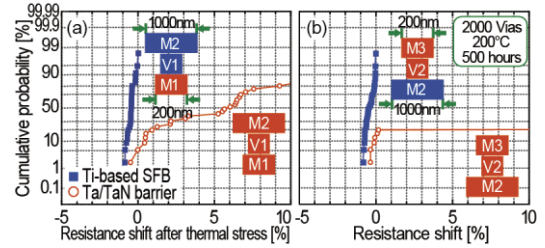


Fig. 17 Stress induced voiding tests of wide M2 via chain at 200°C for 500h; (a) M1=200 nm, V1=70 nm, M2=1000 nm and (b) M2=1000 nm, V2=70 nm, M3=200 nm¹⁶⁾.

(GB) segregation of X atoms, which prevented vacancy migration through GB²⁶⁻²⁸⁾. In a case of the Ti-based SFB, Ti atoms were detected in the ECP-Cu films by secondary ion mass spectroscopy (SIMS) analyses (not shown). This suggests that the Ti atoms must be segregated in GB, and improve stress migration reliability.

4. Cu interconnects on TFT-LCDs using Ti-based SFB

This process using Ti as an alloy element can be applied for Cu-interconnect formation in TFT-LCDs. Cu(Ti) alloy films were deposited on glass substrates and subsequently annealed in vacuum at 400°C for 3 h. Resistivity of the annealed Cu(Ti) alloy films was significantly reduced to about 2.8 μΩcm. Tensile strength of the Cu(Ti)/glass interface increased to about 60 MPa after annealing. The low resistivity and excellent adhesion resulted from Ti-based oxide layer formation at the Cu(Ti)/glass interface^{29, 30)}.

5. Conclusions

(1) Ti-based SFB technique was applied to Cu dual-damascene interconnects and its performances were evaluated. TEM and EDX analyses showed that a 2 mm-thick Ti-based SFB was formed along the interface between Cu(Ti) and dielectrics.

(2) Resistance of the narrow line and via decreased to about 70% and 30%, respectively. Also distributions of these resistances were drastically improved. The breakdown voltage and TDDB performance showed similar performance, indicating the Ti-based SFB has enough barrier property against Cu diffusion into dielectrics. The SIV reliability was improved by using Ti-based SFB technique.

(3) XPS analyses showed that the Ti-based SFB consisted of mainly amorphous Ti oxides. Formation of amorphous or crystalline Ti compounds such as TiC, TiN, TiSi was dependent on the kind of dielectric. Those amounts were lower than that of amorphous Ti oxides. RBS analyses showed that Ti-based SFB growth is controlled by a thermally activated process. The E and Z values decreased with decreasing C concentration in dielectrics, suggesting that the growth is controlled by a chemical reactions of Ti atoms with dielectrics.

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References

- 1) M. T. Bohr, and Y. A. El-Mansy, IEEE T. Electron. Dev., **45** (1998) 620.
- 2) M. Moriyama, M. Shimada, H. Masuda, and M. Murakami, Trans. Mater. Res. Soc. Jpn., **29** (2004) 51.
- 3) M. Shimada, M. Moriyama, K. Ito, S. Tsukimoto, and M. Murakami, J. Vac. Sci. Technol. B, **24** (2006) 190.
- 4) K. Namba, T. Ishigami, M. Enomoto, S. Kondo, H. Shinriki, D. Jeong, A. Shimizu, N. Saitoh, W-M. Ki, S. Yamamoto, T. Kawasaki, T. Nakada, and N. Kobayashi, Proc. of AMC (2006), pp.269.
- 5) H. Wojcik, M. Friedemann, F. Feustelt, M. Albert, S.Ohsiekt, J. Metzger, J. Voss, J.W. Bartha, and C. Wenzel, Proc. of IITC (2007), pp.19.
- 6) D. Jeong, H. Inoue, and H. Shinriki, Proc. of IITC (2008), pp.95.
- 7) J. Koike and M. Wada, Appl. Phys. Lett., **87** (2005) 041911.
- 8) T. Usui, K. Tsumura, H. Nasu, Y. Hayashi, G. Minamihara, H. Toyoda, S. Ito, H. Miyajima, K. Watanabe, M. Shimada, A. Kojima, Y. Uozumi, and H. Shibata, Proc. of IITC (2005), pp.188.
- 9) S. Tsukimoto, T. Morita, M. Moriyama, K. Ito, and M. Murakami, J. Electron. Mater., **34** (2005) 592.
- 10) K. Ito, S. Tsukimoto, and M. Murakami, Jpn. J. Appl. Phys., **46** (2007) 1942.
- 11) K. Kohama, K. Ito, S. Tsukimoto, K. Mori, K. Maekawa, and M. Murakami, J. Electron. Mater., **37**, 1148 (2008).
- 12) K. Kohama, K. Ito, Y. Sonobayashi, K. Ohmori, K. Mori, K. Maekawa, Y. Shirai, and M. Murakami, Jpn. J. Appl. Phys., **50** (2011) 04DB03.
- 13) P.J. Ding, W.A. Lanford, S. Hymes, and S.P. Murarka, J. Appl. Phys., **75** (1994) 3627.
- 14) D. Adams, T.L. Alford, N.D. Theodore, S.W. Russell, R.L. Spreitzer, and J.W. Mayer, Thin Solid Films, **262** (1995) 199.
- 15) C.J. Liu and J.S. Chen, Appl. Phys. Lett., **80** (2002) 2678.
- 16) C.J. Liu, J.S. Jeng, J.S. Chen, and Y.K. Lin, J. Vac. Sci. Technol. B, **20** (2002) 2361.
- 17) M.J. Frederick, R. Goswami, and G. Ramanath, J. Appl. Phys., **93** (2003) 5966.
- 18) M.J. Frederick, and G. Ramanath, J. Appl. Phys., **95** (2004) 3202.
- 19) K. Hoshino, H. Yagi, and H. Tsuchikawa, Proc. 7th Int. VLSI Multilevel Interconnection Conf. (Piscataway, NJ; IEEE, 1990), pp.357.
- 20) J. Li, J.W. Mayer, and E.G. Colgan, J. Appl. Phys., **70** (1991) 2820.
- 21) C.-K. Hu, B. Luther, F.B. Kaufman, J. Hummel, C. Uzo, and D.J. Pearson, Thin Solid Films, **262** (1995) 84.
- 22) A.A. Galuska, J.C. Uht, and N. Marquez, J. Vac. Sci. Technol. A, **6** (1988) 110-122.
- 23) W. Gopel, J.A. Anderson, D. Frankel, M. Jaehnig, K. Phillips, J.A. Schafer, and G. Rucker, Surf. Sci., **139** (1984) 333-346.
- 24) K. Kohama, K. Ito, K. Mori, K. Maekawa, Y. Shirai and M. Murakami, J. Electron. Mater., **38**, (2008) 1913.
- 25) K. Ohmori, K. Mori, K. Maekawa, K. Kohama, K. Ito, T. Ohnishi, M. Mizuno, K. Asai, M. Murakami, and H. Miyatake, Jpn. J. Appl. Phys., **49** (2010) 05FD01.
- 26) K. Maekawa, K. Mori, K. Kobayashi, N. Kumar, S. Chu, S. Chen, G. Lai, D. Diehl, and M. Yoneda, Proc. of AMC (2004), pp.221.
- 27) K. Mori, K. Maekawa, N. Amou, D. Kodama, H. Miyazaki, N. Suzumura, K. Honda, Y. Hirose, K. Asai, and M. Yoneda, Proc. of AMC (2006), pp.467.
- 28) T. Tonegawa, M. Hiroi, K. Motoyama, K. Fujii, and M. Miyamoto, Proc. of IITC (2003), pp.216.
- 29) S. Uehara, K. Ito, K. Kohama, T. Onishi, Y. Shirai, and M. Murakami, Mater. Trans., **51** (2010) 1627.
- 30) S. Uehara, K. Ito, K. Kohama, T. Onishi, Y. Shirai, and M. Murakami, Mater. Trans., **52** (2011) 491.