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Contacts for Monolithic 3D architecture: Study of Ni$_{0.9}$Co$_{0.1}$ Silicide Formation

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Abstract—In this work, we studied the solid-state reaction between a Ni$_{0.9}$Co$_{0.1}$ film and a silicon substrate. NiCo silicide is considered to substitute Ni- and NiPt-based silicides in 3D integration in order to extend the bottom transistor thermal stability. Thanks to the combined analysis of sheet resistance data, X-ray reflectivity spectra modelling, X-ray diffraction and wavelength dispersive X-ray fluorescence analyses on Ni$_{0.9}$Co$_{0.1}$/Si samples annealed at various temperatures, we were able to describe the phase sequence of the NiCo silicide formation.

Keywords—NiCo, silicide, solid-state reaction

I. INTRODUCTION - CONTEXT

In the past half-century, the integrated circuit (IC) industry has been growing rapidly, benefiting from the dimensional downsizing of transistors according to Moore’s law as well as larger wafer sizes. This downsizing results in higher performance, lower power consumption, more complex functionality, faster device speeds, and lower cost per transistor. 3D integration is regularly mentioned for its potential in decreasing costs, variability and delay in interconnections limiting nowadays ICs performance. The CoolCube™ 3D or sequential planar integration for future nodes [1], [2].

However this integration faces the challenge to realize a high performance transistor at the top level without degrading the electrical characteristics of the bottom one. Previous work highlighted that silicide is the main responsible of bottom transistor thermal stability degradation beyond 500 °C [3]. Even if addition of Pt in Ni-based silicides allows extending the monosilicide thermal stability (by increasing the barrier of nucleation of NiSi$_2$) [4], thermal budget involved for the elaboration of top level irremediably leads to morphological degradation and the subsequent deterioration of contact properties (i.e. increase of leakage or even shorts in the junctions) [3].

We investigated the addition of 10 % of Co in NiSi silicide in order to extend its thermal stability and to propose an alternative to Ni- and NiPt-based silicides. Indeed, if Pt is soluble in the monosilicide NiSi phase, Co is not. This element is thus expected to be mainly located at the surface and on grain boundaries within the silicide film and to affect the disilicide formation through other mechanisms than Pt (i.e. surface/interface energy effects) [5], [6]. In this work, we studied the solid-state reaction between a Ni$_{0.9}$Co$_{0.1}$ film and a silicon substrate after several annealing processes at temperatures ranging from 180 to 800 °C.

II. EXPERIMENTAL DETAILS

TiN (7 nm) / Ni$_{0.9}$Co$_{0.1}$ (7 nm) / 300 mm Si(100) substrate samples were prepared using RF-PVD chambers. Before metal deposition, Si substrates were treated by HF 0.5 % solution at around 18 °C and 260 °C, the sheet resistance is increasing from 180 °C to 800 °C for 30 s and under N$_2$ atmosphere. Finally, samples were treated in hot H$_2$O$_2$/H$_2$SO$_4$ solution followed by NH$_4$OH/H$_2$O$_2$ solution in a Screen SU-3100 single wafer cleaner in order to selectively remove metallic phases without impacting silicides.

The sheet resistance of samples was measured by using a fully automatic 4 point probe sheet resistance Napson WS-3000 tool, X-ray reflectivity (XRR) spectra were acquired on a Jordan Valley JVX5200 X-ray reflectometer, in plane X-ray diffraction (XRD) was performed on Rigaku SmartLab high-resolution X-ray diffractometer and Wavelength Dispersive X-ray Fluorescence (WDXRF) was realised on Rigaku AZX400 sequential WDXRF spectrometer.

III. RESULTS & DISCUSSION

Figure 1 plots the R$_{\text{sheet}}$ of the samples right after annealing (circles) and after selective etching (crosses) at temperatures ranging from 180 °C to 800 °C.

Right after annealing, four areas can be observed. Between 180 °C and 260 °C, the sheet resistance is increasing from 58 Ω/□ to 85 Ω/□. Then, between 280 °C and 320 °C a plateau around 87 Ω/□ is noticed. From 320 °C to 400 °C sheet resistance is decreasing until reaching a minimum value at around 18 Ω/□ which is nearly constant (a slight decrease is noticed) between 400 °C and 800 °C. The selective etching consists in the removal of TiN and unreacted NiCo layers, this leads to a modification of the sheet resistance behaviour and values, especially in the two first areas. Indeed, after selective etching, the sheet resistance is only governed by the silicide’s formation and its increase of thickness. Thus, sheet resistance variations can be explained by thickness measurements performed by XRR.

Figure 2 shows the modelling of XRR spectra after annealing (Fig. 2a) and after selective etching (Fig. 2b).
Fig. 1. Sheet resistance measured after annealing Ni\textsubscript{0.9}Co\textsubscript{0.1} (7 nm) on Si substrate at various temperatures for 30 s under N\textsubscript{2}. The sheet resistances of annealed samples with and without selective etching are indicated by crosses and circles, respectively.

Fig. 2. XRR spectra modelling without selective etching (a) and with selective etching (b) of TiN (7 nm) / Ni\textsubscript{0.9}Co\textsubscript{0.1} (7 nm) / Si samples at various annealing temperatures.

The analysis of XRR spectra modelling exhibits a complex variation of phase sequence depending on the annealing temperature. At low temperature and without selective etching, the thickness of the denser phase (8.7 - 8.8) has been attributed to the unreacted NiCo (bulk theoretical density: 8.9). The NiCo thickness decrease with increasing temperature and no more NiCo is detected beyond 220 °C. In this range of temperature, the metal NiCo is consumed for the benefit of the formation of a metal rich intermetallic compound (NiCo)\textsubscript{x}Si\textsubscript{y} [7], [8] with a density of about 7 - 7.5. For simplification purpose, XRR modelling of metal rich silicide has been depicted as one phase on figure 2 but two layers with two distinct densities are necessary in order to obtain a good fitting of experimental XRR spectra. After selective etching, metallic layers (TiN and unreacted NiCo) are removed and a SiO\textsubscript{2} passivation layer is observed [9]. Spectra modelling exhibits that the thickness of (NiCo)\textsubscript{x}Si\textsubscript{y} layer increases with annealing temperature until reaching a plateau at around 280 - 300 °C. At higher temperature, typically 400 °C, modelling shows that a phase with a density around 6 - 6.5 is obtained. This phase has been attributed to the monosilicide (NiCo)Si thanks to XRD analyses.

Figure 3 shows the in-plane XRD spectra of samples after annealing at various temperatures and without selective etching.

Fig. 3. In-plane XRD spectra of TiN (7 nm) / Ni\textsubscript{0.9}Co\textsubscript{0.1} (7 nm) / Si samples annealed at various temperatures and without selective etching.

Thanks to JCPDS database and CaRIne Crystallography modelling, we were able to index most of the diffraction peaks of XRD spectra. TiN (200) and (220) peaks have been identified on all the XRD spectra. At 260 and 280 °C, two peaks have been attributed to a Ni\textsubscript{2}Si-like phase labelled (NiCo)\textsubscript{2}Si and corresponding to a metal rich phase. At 400 °C, numerous diffraction peaks appear, they have been indexed using an orthorhombic NiSi-like phase and related to the formation of the monosilicide (NiCo)Si phase. The indexation of peak located at 50.8° remains unclear.

The addition of 10 % of Co allows extending the thermal stability of (NiCo)Si by repelling the formation of the more resistive disilicide phase (NiCo)\textsubscript{2}Si. In order to specify the role of Co and its incorporation into silicides, WDXRF analyses have been performed. The evolution of Co rate after selective etching is depicted on figure 4.

Important fluctuations of Co rate are observed depending on annealing temperature. Indeed, from 180 °C to 220 °C, Co rate decrease with increasing temperature, then it tends to stabilize at around 6.5 %at. at 220 - 260 °C. Finally, for higher annealing temperatures, the Co rate increases until it reaches
observe that a small amount of Co is incorporated into the as-deposited film) at 400 °C. From the previous analyses, we observe that a small amount of Co is incorporated into the metal rich phase (i.e. (NiCo)Si). As a matter of fact, the incorporating Co rate is far less than 10 %at. Beyond 300 °C the monosilicide (NiCo)Si starts to form and the stabilized phase (since 400 °C) seems to tolerate the incorporation of Co at its nominal rate. The behaviour of Co rate at low temperatures is surprising, thickness of silicide is very low but the Co rate is higher than for annealing temperatures at around 250 °C.

Thanks to the combined analysis of sheet resistance data, XRR spectra modelling, XRD spectra analyses and WDXRF results, we can explain in details the phase sequence of NiCo silicide formation. At low temperature, at least two main phases coexist: NiCo and (NiCo)$_x$Si$_y$. NiCo phase is gradually consumed and leads to the concurrent formation of (NiCo)$_x$Si$_y$. In fact, regarding the XRR and WDXRF results, we assume the formation of two metal-rich silicide phases in which Co has not the same solubility. One of the two phases, probably the one in which the Co rate is the highest, is partially removed during the selective etching due to a poor Si %at. content. At 280 - 320 °C, the transformation is completed and the temperature is still low to initiate the formation of the monosilicide (NiCo)Si. Beyond 320 °C, the formation of (NiCo)Si is initiated until reaching 400 °C. At this point, the monosilicide (NiCo)Si is formed and it is stable until temperatures such as 800 °C.

IV. CONCLUSION

For 3D integration and in order to extend the thermal stability of the bottom transistor, we introduced the use of Ni$_{0.9}$Co$_{0.1}$ silicide. We studied the solid-state reaction between a 7 nm thick Ni$_{0.9}$Co$_{0.1}$ film and a silicon substrate after various annealing processes at temperatures ranging from 180 to 800 °C. Thanks to the combined analysis of sheet resistance data, X-ray reflectivity spectra modelling and X-ray diffraction analyses, we described the phase sequence of the NiCo silicide formation. The additional wavelength dispersive X-ray fluorescence analyses tend to exhibit the formation of two metal-rich phases at low temperatures. Beyond 320 °C, the formation of the monosilicide (NiCo)Si is initiated and this latter is completed at 400 °C. The (NiCo)Si phase appears to be stable until temperatures such as 800 °C.

From a material point of view, NiCo silicide seems to be promising in order to extend the thermal stability of the bottom transistor’s silicide in 3D integration. This material have been implemented on 14-FDSOI technology and its electrical performances have been studied [10].

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Fig. 4. Evolution of Co rate (%at.) measured by WDXRF of Ni$_{0.9}$Co$_{0.1}$ (7 nm) / Si samples annealed at various temperatures and after selective etching.

its nominal rate (10.5 %at., previously determined on NiCo as-deposited film) at 400 °C. From the previous analyses, we observe that a small amount of Co is incorporated into the metal rich phase (i.e. (NiCo)$_x$Si$_y$). As a matter of fact, the incorporating Co rate is far less than 10 %at. Beyond 300 °C the monosilicide (NiCo)Si starts to form and the stabilized phase (since 400 °C) seems to tolerate the incorporation of Co at its nominal rate. The behaviour of Co rate at low temperatures is surprising, thickness of silicide is very low but the Co rate is higher than for annealing temperatures at around 250 °C.

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