Extremely low temperature selective epitaxial processes for advanced CMOS applications

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Abstract

Selective epitaxial growth of SiGe:B and SiGe:P layers at a very low growth temperature of 400°C in a commercial RPCVD epi tool is presented. The process was verified on device wafers to confirm their compatibility with advanced processes. A fine multilayered structure of the deposited layers reflecting the cyclic nature of the processes has been observed. High active concentrations of ~6e20 cm⁻³ and ~1e20 cm⁻³ for B and P are obtained, suitable for low temperature Source/Drain application of novel advanced device structures.

1. Introduction

One of the most obvious trends in group IV epitaxy is a systematic lowering of process temperature. This is governed by few reasons. On one hand novel device concepts which must fulfill scaling paradigm like gate all around FETs [1], complementary FETs [2], 3D stacked devices [3], etc. put stringent temperature requirements to epitaxy due to complex geometry and application of new approaches (complex high-k materials, buried power rail, etc.). On the other hand introduction of SiGe and Ge in the device’s channel or SiGe with high Ge content, Ge, GeSn, SiGe:Sn as source/drain (S/D) materials requires rather dramatic decrease of the epi process temperature in order to maintain layers properties.

In view of this trend use of high order Si and Ge precursors allowing development of epi processes at temperatures well below 500°C becomes very promising [4]. Combining such processes with Cl₂ which allows etching down to extremely low temperatures of ~400°C in the case of Si and ~250°C for Ge [5] open possibilities for development of selective SiGe processes at temperatures of 400°C and below. A visibility of such selective cyclic SiGe:B and SiGe:P at 400°C has been demonstrated on Shallow Trench Isolation (STI) wafers in [6].

In this work we elaborate further on integration of these processes in CMOS flow when full device comprising spacers, metal gates and hard masks is build. Morphological, electrical and structural properties are presented as well.

2. Experimental details

Epitaxial growth reported here was done on patterned wafers in the frame of 3D sequential stacked planar devices work with a typical processing flow reported in [3] using an ASM Intrepid XPTM epi tool, which is a horizontal cold wall, load-locked reduced pressure chemical vapor deposition system designed for production applications.

Low temperature selective growth of SiGe layers was done using high order precursors: disilane (Si₂H₆) and digermaine (Ge₂H₆) as Si and Ge precursors. B₂H₆ (B₂H₆ 1% in H₂) or PH₃ (PH₃ 15% in H₂) and Cl₂ were used as p and n type dopants and etching gases respectively. Peculiarities of the growth and selectivity tuning has been reported elsewhere [6].

A micro four-point-probe using the CIPTechTM M300 Micro Hall system from Capres A/S was used in order to measure active carrier concentration [7]. Scanning electron microscopy (SEM) was performed on EXPIDA1285 from FEI and eDR-7100 from KLA Tencor. Transmission electron microscopy (TEM) and high angle annular dark field scanning transmission electron microscopy (HAADF-STEM) was studied in Tecnai 30 F microscope operating at 300 kV.

3. Results and discussion

An application of the SiGe:B low temperature process developed in [6] to a device wafer for 3D stack integration resulted in a severe selectivity loss. It is well known that selectivity of a process depends on the type of oxides and nitrides used and usually must be tuned for each specific application.

Fig. 1. Selectivity tuning of cyclic SiGe:B : long deposition/short etch a), long deposition/long etch b), short deposition/short etch c), final tuned selective process d).
Since in our case we apply a cyclic process, the main tuning knob to recover selectivity is etch/deposition times. Fig. 1 presents results of four different cases: long deposition/short etch (a), long deposition/long etch, short deposition/short etch (c) and final tuned selective process.

Careful optimization of both deposition and etching allows to obtain a process which is selective to both oxides (Shallow Trench Isolation and hard masks on top of gates) and nitrides (spacers).

A cyclic nature of the process can be directly seen in transmission electron microscopy (TEM). Although Energy Dispersive X-Ray Spectroscopy (EDS) showed uniform over the layer thickness Ge concentration of ~ 40%, both high resolution TEM and high-angle annular dark-field imaging (HAADF) show that S/D consists of a number of SiGe layers which are separated with few atomic layers of SiGe with a darker contrast suggesting presence of SiGe with lower Ge concentration. These intermediate low Ge concentration layers are formed after an etching step and can be explained by Cl₂ preferential Ge pull out of the SiGe surface. Despite of this peculiarity SiGe:B layers demonstrated very good quality and high active B concentration of ~ 6e²⁰ cm⁻³ which is suitable for Si-based pMOS S/D application.

In the case of Ge-based devices selective SiGe:P is an interesting option for the n-MOS S/D since it typically offers higher active P concentration compared to selective Ge:P and tensile stress to the channel. In frame of this work we evaluated a similar growth strategy for SiGe:P. The selectivity tuning strategy was similar to SiGe:B (Fig.1) and described in [6]. A TEM image of the final SiGe:P layer with Ge concentration ~ 75% is shown in Fig. 3. A multi structure of SiGe:P similar to Fig.2 reflecting the process cyclic nature has been observed. Blanket experiments of the deposition part of this process showed that active P concentration of ~ 1e²⁰ cm⁻³ can be achieved when both Ge and P flows are optimized.

4. Conclusions

In this work we investigated selective growth of SiGe:B and SiGe:P at temperature of 400°C to be used in advanced device structures. Careful optimization of both deposition and etching allows to obtain processes which are selective to both oxides and nitrides present on the device wafers. A fine multilayered structure of the deposited layers is explained by the cyclic nature of the processes and Ge pull out effect during Cl₂ etching step. High active concentrations of ~ 6e²⁰ cm⁻³ and ~ 1e²⁰ cm⁻³ for B and P suitable for low temperature Source/Drain application of novel advanced device structures are obtained.

Acknowledgements

The authors express their sincere appreciation to the imec core CMOS program members, the European Commission, the TAKEM15 ECSEL project, local authorities and the imec pilot line. Air Liquide Advanced Materials is acknowledged for providing Ge:Hs.

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