Amorphous Silicon Nanowires Grown on Silicon Oxide Film by Annealing
Zhishan Yuan1,2*, Chengyong Wang1, Ke Chen2, Zhonghua Ni2 and Yunfei Chen2

Abstract
In this paper, amorphous silicon nanowires (α-SiNWs) were synthesized on (100) Si substrate with silicon oxide film by Cu catalyst-driven solid-liquid-solid mechanism (SLS) during annealing process (1080 °C for 30 min under Ar/H2 atmosphere). Micro size Cu pattern fabrication decided whether α-SiNWs can grow or not. Meanwhile, those micro size Cu patterns also controlled the position and density of wires. During the annealing process, Cu pattern reacted with SiO2 to form Cu silicide. More important, a diffusion channel was opened for Si atoms to synthesis α-SiNWs. What is more, the size of α-SiNWs was simply controlled by the annealing time. The length of wire was increased with annealing time. However, the diameter showed the opposite tendency. The room temperature resistivity of the nanowire was about 2.1 × 103 Ω·cm (84 nm diameter and 21 μm length). This simple fabrication method makes application of α-SiNWs become possible.

Keywords: α-SiNWs, Cu patterns, Annealing time, Resistivity

Background
Among the various classes of one-dimensional semiconductor nanostructure, silicon nanowire (SiNW) has been exhibited bright future in the fields of electronic, photovoltaic solar, photonic, battery, and sensor. [1–6] The SiNW manufacture method includes top-down and bottom-up approaches. Table 1 is the summary of different SiNW manufacture method. Top-down approach is usually realized by reactive ion etching (RIE) and metal-catalyzed electroless etching of silicon. In those methods, nanowire site is controlled in top-down approach by nanofabrication tools such as e-beam lithography, [7] nanoimprint lithography [8], or nanosize template such as PS sphere, [9] AAO mask [10]. Nanofabrication tools control the site, size, orientation, and numbers of wire well with high-cost and complex fabrication process. Nanosize template [9–11] is the low-cost method, but the fabrication process is more complex than nanofabrication tool method for template should be built and removed during the whole process. Therefore, template-free method shows good potential in future [12]. Another top-down approach uses MEMS technique to fabricate site controllable SiNWs [13], this fabrication process easily fabricate SiNW sensor devices. However, MEMS technique brings complex manufacture process with high cost.

In bottom-up approach, chemical vapor deposition (CVD) is an important approach to synthesis SiNWs with low-cost and simple fabrication process. And this approach can readily produce extremely small diameter and super long SiNWs (as recorded, the smallest diameter was 1 nm, and the longest was millimeters) [14–16]. Good quality SiNWs are always synthesized through vapor-liquid-solid (VLS) mechanism with the help of Au or other metals in this method [2]. However, those novel materials are prohibited in clean rooms for degrading the electrical and optical properties of semiconductors.

Catalyst free method is put forward to solve pollution problem which brought by novel catalysts in bottom-up approach. Oxide-assisted growth (OAG) method does not require any metal catalyst [17]. Unfortunately, the compatibility with Si-based integration technology is poor in this method. And products are always affected by other residual impurities easily [18]. Room temperature continuous wave laser ablation of Si is another way to synthesis SiNWs without using metal catalyst [19]. Nevertheless, high vacuum is needed. Even in the simple SiO evaporation technique, good size controllability is always hard to realize. Moreover, SiO powder is harmful to health [20].
New catalysts such as aluminum and copper are researched to open the door of complementary metal oxide semiconductor (CMOS) technology to SiNWs [21]. Aluminum is used to reduce the deep level impurities; it can also be a p-type dopant producing a shallow acceptor in Si. However, the high sensitivity to oxidation makes using aluminum as catalyst method becoming unpractical. Copper is a good conductor of heat and electricity and has been widely used in integrated circuits (ICs) and CMOS processing. So, copper is considered as the suitable catalyst for SiNW growth. The size and site of Si wires were well controlled by copper catalyst in Kayes et al. work [22]. In the works which copper was used as catalyst to synthesis SiNWs, SiH₄, Si₂H₆, or SiCl₄ gases were used as Si precursor [22–24].

In this paper, we present a simple and effective method to synthesis SiNWs on SiO₂ films by Cu catalyst-driven SLS mechanism during annealing process without using any toxic precursor gases. This method has two advantages. Firstly, the metal contamination of the SiNWs was decreased. Secondly, no toxic precursor gases were used.

### Methods

#### Chip Fabrication

First, 300 nm SiO₂ film was grown on single side polished n-type silicon (100) wafers by thermal oxidation (Fig. 1a). Then, 400 nm copper film was deposited on SiO₂ by magnetron sputtering. After photolithographic process and ammonium persulfate solution (1:100 water) etching, Cu micron-size pattern array were fabricated on SiO₂ surface in target area (Fig. 1b). Subsequently, the wafer was diced into chips. And those chips were ultrasonically cleaned by ethanol and acetone in turn for 10 min. Afterwards, DI water was used for last clean process before blow-dry by N₂.

#### α-SiNW Growth

One thousand standard cubic centimeters per minute of Ar was used to exclude air in the tube for 10 min after chips were put on quartz boat and transferred into the center of the horizontal furnace.

Subsequently, five stages were used to synthesis SiNWs. The detailed annealing processing conditions are given in Fig. 2. In stage I, the temperature was increased from

---

Table 1 Summary of different SiNW manufacture method

| Manufacture method       | Advantages                                                                 | Disadvantages                                                                 | References |
|--------------------------|---------------------------------------------------------------------------|-------------------------------------------------------------------------------|------------|
| Top-down RIE             | The site and size of nanowires were well controlled.                       | Need nanofabrication tools.                                                   | [7–10]     |
| Metal-catalyzed electroless etching | The site and size of nanowires were well controlled, without any nanofabrication tools. | Template fabrication process was complex.                                      | [11]       |
| MEMS technique           | The site and size of nanowires were well controlled.                       | Complex fabrication process and time-consuming.                               | [13]       |
| Bottom-up CVD            | Simple and low cost and the quality of SiNW was good.                      | Novel metal materials were prohibited in clean rooms. Controllability is poor.| [14–16]    |
| OAG                      | Simple and low cost and no metal catalyst is needed.                       | Controllability is poor. The compatibility with Si-based integration technology was poor. | [17, 18] |
| Laser ablation           |                                                                          |                                                                               | [19]       |
| SiO evaporation          |                                                                          |                                                                               | [20]       |

---

---

Fig. 1 Schematic depiction of the fabrication process. a Thermal oxidation. b Cu micron-size pattern array fabrication. c Copper patterns changed into hemispheres. d Nanowire growth
room temperature to 400 °C in 1 h with the same Ar flow which is used to exclude air. In stage II, Ar flow was adjusted to 100 sccm, and 20 sccm H₂ was added. It took 2 h to reach 1080 °C. In this stage, copper patterns changed into hemispheres (Fig. 1c). Then, temperature was held for 30 min with 1000 sccm Ar and 40 sccm H₂ in stage III. After turning off the furnace, the fast cooling process only 10 min was taken as the IV stage and the flow was adjusted to 500 and 20 sccm respectively. In the last stage, slow cooling used to decrease the furnace temperature to room temperature with 100 sccm Ar and 20 sccm H₂. After the five stages, α-SiNWs were grown on the position of Cu patterns as shown in Fig. 1d.

**Characterization**

Scanning electron microscopy (SEM, Hitachi S-4800) and high-resolution transmission electron microscopy (TEM, JEM-2100F operating at 200 Kv) equipped with energy dispersive spectrometer (EDS) were employed for...
analyzing the morphology and composition of the nanowires. For TEM measurements, Mo grid was used to support nanowires. For FIB etching the root of the wire, a thin layer of Au was evaporated on the surface to protect the wire by electron-beam-induced deposition (EBID). Two-terminal device was used to measurement the resistivity of nanowire [25]. The wire was mechanically removed from the substrate by nano-operator equipped on focused ion beam (FIB) (FEI, QUANTA3D 600FIB System). Then, nanowire was weld on the two electrodes by Pt deposited with assisted electron beam. Finally, the resistivity of the nanowire was measured by Cascade Semi-automatic probe station HP 4156.

**Results and Discussion**

Figure 3 presents the SEM photos of two samples before and after annealing (sample I, the thick Cu film is 400 nm, sample II is the Cu pattern arrays with size of 400 nm thick and 1.9 μm diameter, and center-to-center pitch is 10 μm). It is obvious that the results of the two samples were quite different after annealed at 1080 °C for 30 min. For Cu film, shown in Fig. 3b, only Cu balls were scattered randomly on the surface of SiO₂. The inserted figure in Fig. 3b was the diameter distributions of Cu balls, and the average diameter of the ball was 4.4 μm. In-suit nanowire appeared in sample II after annealing in Fig. 3d. The length of nanowire can be as long as 20 μm, and the diameter of nanowire is about 57 nm as shown in the inserted image of Fig. 3d. It is clear that each pattern has grown one nanowire and the center-to-center distance equal to the value of Cu patterns. This means the density of nanowires can be controlled by number of Cu patterns simply. The phenomenon in Fig. 3 demonstrates that the micro size of Cu patterns are suitable for nanowire growth (in our case, the size of Cu pattern was 400-nm thick and 1.9 μm diameter). For
Cu film, dewetting effect happened at high temperature. In order to reduce the surface energy of Cu film, Cu balls were aggregated in random way (in Fig. 3b).

The high-resolution transmission electron microscopy (TEM) image in Fig. 4a reveals that the nanowire has a smooth morphology at diameter of 50 nm in sample II. The highly diffusive ring pattern (inset) of the selected area electron diffraction (SAED) demonstrates that the nanowire was totally amorphous (in Fig. 4). Energy dispersive spectrometer (EDS) results in Fig. 4 indicate that the wire consists of Si and O with atomic ratio of 4, which is far from the ratio of Si dioxide and suggests that a trace amount of oxygen exists in the SiNWs. For reduction atmosphere which was composed of Ar and H₂ is maintained during nanowires growth process, so the light oxidation only happened during sample exposure to air after fabrication.

After FIB etching the root part of the wire and substrate, cross section of the wire root was characterized by SEM with sample holder rotated 45°. It is interesting to find that the nanowire grown from the boundary between Si and SiO₂ in Fig. 5. A long Si gap is also found at the Si/SiO₂ interface. Those observations demonstrate that the substrate was the only Si source for the wire. Meanwhile, no metal particle is found at the tip part of the wire. According to those results, a possible schematic illustration of α-SiNW growth is presented in Fig. 6 based on solid-liquid-solid mechanism. During the annealing process, Cu patterns (Fig. 6a) dewet to the center of the pattern (Fig. 6b) and react with SiO₂ to form Cu silicide (Fig. 6c). Then, Si atoms permeate into the Cu silicide. During this process, the different diffusion speed of Si atoms in the substrate which caused by the defect of substrate may induce the Si gap formation. When the dissolving Si atoms in silicide reached saturation, Si starts to precipitate to synthesis α-SiNWs (Fig. 6d).

It is clear that Cu has played a very important role in our study. Something like black particle can be found at the tip of the wire, although in most wires, this particle is not existed. The mapping results (Fig. 7) show that no metal particle exists at the tip of the wire. The particle seems like the misunderstanding by the angle of between the wire and holder, which was not suitable to observe. Unfortunately, no copper can be found at the root part of the wire (Fig. 5).
Cu diffused into Si substrates is the possible way that may give rise to this surprising result. It is well known that fast diffusion of Cu atoms in Si was tested at high temperature [26]. So, Cu atoms could diffuse into Si substrate in a few minutes after the window in SiO$_2$ was opened at high temperature.

To demonstrate the controllability of our method, different annealing time is carried out in our experiments. The SEM of nanowires synthesized in different annealing time with the same Cu pattern size (400-nm thick and 1.9 um diameter) and same SiO$_2$ thickness (300 nm) is shown in Fig. 8. Most nanowires have uniform diameter. It is interesting to find that the diameter decreased when the direction of the wire changed. As red arrow shows in Fig. 8c, the tip part diameter is 76 nm, and the root part is only 49 nm. This huge difference in diameter in the same wire may be caused by the variation of the energy per unit area for the nucleus [27]. And this phenomenon is seldom to see. Another interest finding is that the diameter of root part was the bigger part in the whole wire, and the tip part was smaller (red arrows shown in d–f). Comparing with the whole wire, the length of nonuniform part is very short. This result presents that α-SiNW had nonuniform diameter.

After SEM, the length and diameter of root part of α-SiNWs are calculated. The results in Fig. 9 show that the length of α-SiNW was increased with annealing time, as a function of the anneal time. The diffusion time of Si atoms is increased offering more atoms to synthesis nanowire. The length of α-SiNW increases to 24 μm while the annealing time increased to 30 min.

**Fig. 8 a–f** The SEM images of α-SiNWs grown in different annealing time

**Fig. 9** The nanowire diameter and length as a function of the anneal time. Blue triangles and red circles in figure are the date for length and diameter of nanowire in experiment, and blue and red lines in figure were the fitting line according to the experiment dates.

**Fig. 10** The electrical transport measurement of α-SiNW in Fig. 3. Two-terminal device was used to measure the resistivity of nanowire [25]. The wire was mechanically removed from the substrate by nano-operator equipped on focused ion beam (FIB) (FEI, QUANTA3D 600FIB System). Then, nanowire was weld on the two electrodes by Pt deposited with assisted electron beam. The resistivity of the nanowire was measured by Cascade Semi-automatic probe station HP 4156. Finally, the room temperature resistivity of the nanowire in Fig. 3 is $2.15 \times 10^3 \Omega \cdot cm$, measured by two-probe method.
The average growth rate of nanowire is approximately 1.1 μm/min, which was similar to the growth rate by annealing with block Si source [28]. The rapid speed of growth is leaving no time for Si atoms to stack themselves into crystalline order. Finally, amorphous nanowires instead of crystalline are synthesized.

The diameter of α-SiNW is decreased from 81 to 57 nm in annealing time increasing process. Usually, the length of SiNW depends on their diameter for Gibbs-Thomson effect in vapor-liquid-solid growth using silane as gaseous source and gold as catalyst. The length of SiNW increases when diameter increased for nanowires with diameter under 100 nm. Nevertheless, the result in our experiment shows the inverse conclusion that the diameter decreased with length. Long-time annealing gives more time for Cu atoms diffused into Si substrate, and the volume of silicide catalyst is also decreased. Meanwhile, the diffusion process of Si atoms is continued which made growth of α-SiNW all the time with catalyst particle size change. Therefore, the diameter of α-SiNW is decreased with anneal time.

Figure 10 shows the current (I) versus voltage (V) fitting curve with a near ohmic behavior. I-V measurement shows the room temperature resistivity; the measurement shows the room temperature resistivity; the nanowire in Fig. 3 is 2.15 × 10³ Ω-cm, measured by two-probe method. Comparing with M. Lieber’s work [29], the resistivity of α-SiNW in this work is ten times higher than the single crystal silicon wires without doping. The significantly higher electrical conductivity of the nanowires is probably due to the size effect.

Conclusions
In conclusion, α-SiNWs are grown directly on SiO₂ surface during annealing process in Ar/H₂ atmosphere via SLS mechanism without any toxic precursor gases. Cu patterns fabrication is the necessary condition for α-SiNW growth. Meanwhile, Cu patterns are used to control the density and the site of α-SiNWs. What is more, the annealing time is adjustable parameters to control the diameter and length of wire α-SiNWs. The room temperature resistivity of the nanowire is 2.15 × 10³ Ω-cm. This new growth method makes α-SiNWs candidate for potential applications in the future.

Acknowledgements
This work was supported by the National Natural Science Foundation of China (nos. 51505083 and 51505089).

Authors’ Contributions
ZY designed the experiments. ZY synthesized the silicon nanowires. KC and ZN conceived the study. ZY, CW, and YC analyzed the SEM data and wrote the initial drafts of the work. All authors discussed the results and commented on the manuscript. All authors read and approved the final manuscript.

Competing Interests
The authors declare that they have no competing interests.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Received: 5 April 2017 Accepted: 26 July 2017
Published online: 10 August 2017

References
1. Yu P, Wu J, Liu S, et al. (2016) Design and fabrication of silicon nanowires towards efficient solar cells. Nano Today
2. Schmidt V, Wittemann JV, Serz S, Gosele U (2009) Silicon nanowires: a review on aspects of their growth and their electrical properties. Adv Mater 21(25–26):2681–2702
3. Peng KQ, Lee ST (2011) Silicon nanowires for photovoltaic solar energy conversion. Adv Mater 23(2):198–215
4. Lu W, Lieber CM (2007) Nanoelectronics from the bottom up. Nat Mater 6(11):841–850. doi:10.1038/Nmat2028
5. Wang YL, Wang YF, Da PM, Xu M, Wu H, Zheng GF (2013) Silicon nanowires for biosensing, energy storage, and conversion. Adv Mater 25(57):5177–5195
6. Dasgupta NP, Sun JW, Liu C, Brittain S, Andrews SC, Lim J, Gao HW, Yan RX, Yang PD (2014) 25th anniversary article: semiconductor nanowires synthesis, characterization, and applications. Adv Mater 26(14):2137–2184. doi:10.1002/adma.201305929
7. Park I, Li Z, Pisano AP et al. (2009) Top-down fabricated silicon nanowire sensors for real-time chemical detection. Nanotechnology 21(1):015501
8. Morton KJ, Nieberg G, Bai S et al (2008) Wafer-scale patterning of sub-40 nm diameter and high aspect ratio (>50:1) silicon pillar arrays by nanoimprint and etching. Nanotechnology 19(34):345301
9. Huang Z, Fang H, Zhu J (2007) Fabrication of silicon nanowire arrays with controlled diameter, length, and density. Adv Mater 19(53):744–748
10. Huang Z, Geyer N, Werner P et al (2011) Metal-assisted chemical etching of silicon: a review. Adv Mater 23(2):285–308
11. Chang SW, Chuang VP, Boles ST et al (2009) Densely packed arrays of ultra-high-aspect-ratio silicon nanowires fabricated using block-copolymer lithography and metalloc-assisted etching. Adv Funct Mater 19(15):2495–2500
12. Bai F, Li M, Huang R et al (2012) Template-free fabrication of silicon micropillar/nanowire composite structure by one-step etching. Nanoscale Res Lett 7(1):557
13. Yun SS, Yoo SK, Yang S et al (2008) Volume-producing fabrication of a silicon nanowire via crystalline wet etching of (1 1 0) silicon. J Micromech Microeng 18(9):095017
14. Ma DDD, Lee CS, Au FCK, Tong SY, Lee ST (2003) Small-diameter silicon nanowire surfaces. Science 299(5614):1874–1877. doi:10.1126/science.1080313
15. Shi WS, Peng HY, Zheng YF, Wang N, Shang NG, Pan ZW, Lee CS, Lee ST (2000) Synthesis of large areas of highly oriented, very long silicon nanowires. Adv Mater 12(18):1343–1345. doi:10.1002/1521-4095(20000912):18:1343<Aid-Adma1343>3.0.Co;2-Q
16. Park WI, Zheng GF, Jiang XC, Tian BZ, Lieber CM (2008) Controlled synthesis of millimeter-long silicon nanowires with uniform electronic properties. Nano Lett 8(9):3004–3009. doi:10.1021/Nl802063q
17. Wang N, Tang YH, Zhang YF, Lee CS, Bello I, Lee ST (1999) Si nanowires grown from silicon oxide. Chem Phys Lett 299(2):237–242. doi:10.1016/S0009-2614(99)01028-7
18. Xu XD, Wang YC, Liu ZF, Zhao RG (2007) A new route to large-scale synthesis of silicon nanowires in ultrahigh vacuum. Adv Funct Mater 17(11):1729–1734. doi:10.1002/adfm.20060658
19. Kokai F, Inoue S, Uchiyama K, Takahashi Y, Koshio A (2013) Catalyst-free growth of amorphous silicon nanowires by laser ablation. Appl Phys a-Mater 112(1):1–7. doi:10.1007/s00339-012-7169-y
20. Zheng RX, Yang QQ, Hirose A, Jari O, Mukhopadhyay I (2014) Catalyst-free synthesis of silicon nanowires by oxidation and reduction process. J Mater Sci 49(10):3592–3597
21. Arbiol J, Kalache B, i Cabarrocas PR, et al (2007) Influence of Cu as a catalyst on the properties of silicon nanowires synthesized by the vapour-solid–solid mechanism. Nanotechnology 18(39):305606. doi:10.1088/0957-4484/18/30/305606
22. Kayes BM, Filler MA, Putnam MC, et al (2007) Growth of vertically aligned Si wire arrays over large areas (>1 cm²) with Au and Cu catalysts. Appl Phys Lett 91(10):103110
23. Renard VT, Jublot M, Gergaud P, Chemis P, Rouchon D, Chabli A, Jousseauine V. (2009) Catalyst preparation for CMOS-compatible silicon nanowire synthesis. Nat Nanotechnol 4(10):654–657. doi:10.1038/nnano.2009.234

24. Wen CY, Reuter MC, Tersoff J, Stach EA, Ross FM (2010) Structure, growth kinetics, and ledge flow during vapor-solid-solid growth of copper-catalyzed silicon nanowires. Nano Lett 10(2):514–519. doi:10.1021/Nl903362y

25. Zeng HJ, Li T, Bartenwerfer M, Fatikow S, Wang YL (2013) In situ SEM electromechanical characterization of nanowire using an electrostatic tensile device. J Phys D Appl Phys 46(30):Art. 305501. doi:10.1088/0022-3727/46/30/305501

26. Wen, C. Y, Reuter, M. C.; Tersoff, J.; Stach, E. A.; Ross, F. M. Structure, growth kinetics, and ledge flow during vapor-solid-solid growth of copper-catalyzed silicon nanowires. Nano Lett 2010, 10, 514-519

27. Cao Y, Yang G (2012) Vertical or horizontal: understanding nanowire orientation and growth from substrates. J Phys Chem C 116(10):6233–6238

28. Yan HF, Xing YJ, Hang QL, Yu DP, Wang YP, Xu J, Xi ZH, Feng SQ (2000) Growth of amorphous silicon nanowires via a solid–liquid–solid mechanism. Chem Phys Lett 323:224–228

29. Cui Y, Duan XF, Hu JI, Lieber CM (2000) Doping and electrical transport in silicon nanowires. J Phys Chem B 104(22):5213–5216. doi:10.1021/jp0009305