Nanostructures

One-Step SnO₂ Nanotree Growth
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Abstract: A comparison between Au, TiO₂ and self-catalysed growth of SnO₂ nanostructures using chemical vapour deposition is reported. TiO₂ enables growth of a nanonetwork of SnO₂, whereas self-catalysed growth results in nanoclusters. Using Au catalyst, single-crystalline SnO₂ nanowire trees can be grown in a one-step process. Two types of trees are identified that differ in size, presence of a catalytic tip, and degree of branching. The growth mechanism of these nanotrees is based on branch-splitting and self-seeding by the catalytic tip, facilitating at least three levels of branching, namely trunk, branch and leaf.

Metal-oxide semiconductors have served as functional materials for gas sensors for decades.[1] The sensitivity, response speed and power efficiency scale with the size of the sensing element. Nanostructures of SnO₂ are ideal building blocks for realising improved sensors, for example, for inflammable gases such as CO.[2–4] Branched nanowires are particularly suited to provide high sensitivity at low cost without the need of spatial resolution, that is, it is of minor importance where a single gas molecule is detected on the sensor as long as it can be detected. The most economical method to fabricate such structures is chemical vapour transport.[5] This method employs a catalysed growth process in which nanowires are seeded by Au nanoparticles.[6] Branches are typically added by multi-step catalyst seeding.[7] Here, we demonstrate the efficient one-step growth of SnO₂ nanotrees using Au catalyst nanoparticles and explain the growth mechanism.

Experimental Methods

Samples were synthesised in a horizontal tube furnace (Nabertherm B180, 25 mm-diameter, heated length 600 mm) from Sn granules placed in a quartz boat. N₂ at atmospheric pressure was used as a carrier gas, with a typical flow rate of 300–900 sccm. The boats were physically connected to magnetic trans-
seen in Figure 1c. The catalyst particles only accumulate material and grow into spheres at too low flux but nucleation is not initiated owing to limited precursor supply. There is a threshold between 300 and 500 sccm at 800 °C where nucleation starts to occur. The large diameter of the nanowires grown at 500 sccm means that the axial growth rate is very slow and that radial growth dominates. It is the supersaturation of the catalytic tip that drives the growth at high enough flux (900 sccm). The resulting nanowires are thin and long. Many catalyst sites can be activated such that the substrate coverage, that is, the yield, is very high.

All nanowires that were grown using the Au catalyst have a catalytic tip larger than the 5 nm diameter of the initial Au nanoparticle. This is evidence of the fact that the catalyst forms an alloy with incoming precursor vapour and grows in size until supersaturation is reached, consistent with the vapour–liquid–solid (VLS) model.

TiO₂, however, is not liquid during the growth because of its higher melting point, so the VLS mechanism can be ruled out. It either forms a solid solution with SnO₂, or facilitates the formation of Sn droplets that allow for self-catalysed growth. TiO₂ catalyst particles should be found at the tip if a solid solution was formed to lead to tip-catalysed growth. TEM EDS does not detect any TiO₂ in the tip of the nanowire. Therefore, it can be concluded that the TiO₂ nanoparticles are located at the base of the wire. Growth is initiated by the formation of Sn droplets on their surface. Figure 1a shows such droplets as dark spots on TiO₂ patches. The nanowires are amorphous.

Au-catalysed nanowires grow longer and straight compared to the TiO₂ catalysed nanowires (c.f. Figure 2b) and branch occasionally, as also reported by Jin et al. for catalyst-free growth. In the centre of the catalytic tip is a dark spot, which is roughly the size of a single Au nanoparticle or a cluster of several nanoparticles, indicative of VLS growth. Whereas most of the nanostructures are composed of SnO₂, a small fraction of pure Sn is also found (see high resolution image of the tip area in Figure 2c). The growth direction of this short wire is [200], consistent with Sn nanowire growth. The SnO₂ nanowires have a tetragonal crystal structure and grow along the [101]-direction, as shown in Figure 2d.

Growth at 950 °C and a flux of 700 sccm lead to trees with and without catalytic tips. A nanowire tree without catalytic tip is tapered, as shown in Figure 3a and b. It consists of three components: trunk, branches and leaves. The components grow perpendicular to each other and become gradually smaller. Branches at the bottom are longer than branches at the top. The nanotree in Figure 3c is larger and consists of a trunk with 50 nm diameter and fine branches with 30 nm diameter. The branches are curled up and have grown denser than the tree in Figure 3a.

The four sections of the trunk are described in the following. The tip is formed by a droplet as usually observed for VLS-grown nanowires. The next section is marked by the appearance of short branches. The number and length of branches increases towards the main section. Branches are connected to the tree by extruding bases (Figure 3c(II), inset). The fourth section near the base is completely free of branches. An amorphous structure on the root indicates the first tens of nanometres where the growth started.
What is the mechanism for the formation of branches? A low magnification TEM image in Figure 4a gives an overview over the density and length of the branches. The density is especially increased by separation. In Figure 4b a nanobranch splits in the centre and separates into two branches. This can be induced by a crystal defect or, more likely, a split catalyst particle resulting in an additional growth direction. High magnification images reveal 5 nm-sized particles on the walls of the trunk, as shown in Figure 4c. The size is identical to the particle size of our commercial Au nanoparticles. There are three scenarios that can explain this feature: 1) Au nanoparticles migrate over the SnO2 surface and deposit onto the trunk. The 5 nm particle shown in Figure 4c could be such a nanoparticle. However, this would require an exceptional mobility of Au on SnO2 at this temperature; 2) Sn precursor atoms spontaneously form droplets on the SnO2 trunk and branches grow self-catalysed following the VLS mechanism. This also explains the observation in Figure 4c and the formation of leaves grown from smaller droplets on the branches; 3) the main catalyst particle at the tip gives off smaller particles if the size exceeds a certain threshold, as depicted in Figure 4c. These particles can even migrate several micrometers along the trunk. During the first few micrometers of growth the size does not exceed the threshold yet. Hence, fewer branches grow at the bottom. Branch length is a function of height, measured from the trunk, with the longest branches towards the root and short branches, or even seeds, at the top. Leaves grow also mediated by catalyst migration on branches. Some of the nanotrees do not have a catalyst particle attached to the tip, and it can be assumed that it has been consumed during growth, yet they still grow branches.

![Figure 3. SEM images of SnO2 nanotrees. a) Vertical trees at the substrate edge. The inset shows a trunk (brown arrow) with branches (yellow arrow) and leaves (green arrow). b) Sketch of a nanowire tree. The tip is tapered with no catalyst particle visible at the top (upper inset). Both trunk and branches, as well as branches and leaves are perpendicular to each other (lower inset). c) Horizontally oriented nanotree on the surface. There are four sections (I-IV) with distinctive features: tip (I), little branch growth with extrusion in the inset (II), middle of the trunk (III) and root (IV).](Image)

In conclusion, Au and TiO2 nanoparticles are efficient catalysts for SnO2 nanowire growth. Networks of polycrystalline and bent or single-crystalline, straight and branched nanowires can be grown by tuning the growth parameters. TiO2 catalyst particles facilitate the self-catalysed growth by enabling Sn droplet formation on the catalyst surface. At 950 °C Au nanoparticles diffuse over the substrate and the grown structures. This leads to a preferential growth on nanowires forming trunks, branches and leaves. These nanotrees are excellent building blocks for highly-sensitive gas detectors and other nanoelectronic devices.

**Keywords:** crystal growth · nanocatalysis · nanogrowth · nanostructures · physical vapor deposition

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