Production of self-aligned metal nanocolumns embedded in an oxide matrix film

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Self-aligned silver nanocolumns embedded in amorphous Al2O3 matrix are produced by alternate pulsed laser deposition at room temperature in a single step process. The morphology and the spatial distribution have been investigated by standard, high resolution, and scanning transmission electron microscopy. Polycrystalline nanocolumns having diameters close to 3 nm and heights up to 7 nm are all oriented perpendicular to the substrate surface. The self-alignment of the nanocolumns is the consequence of the deposition of consecutive layers of Ag nanoparticles using the first layer as template for the ordered growth of the nanocolumns. Control over the density, diameter and height of the nanocolumns is achievable by tailoring the deposition sequence. © 2006 American Institute of Physics. [DOI: 10.1063/1.2180877]

Metallic nanostructures have a high potential for the production of nanoscale electronic, optoelectronic or magnetic devices. Examples of this potential are the use of nanostructures as building blocks for chemical sensors,1 substrates for surface enhanced Raman scattering,2–4 and contacts5 or single-electron transistors6 for nanoelectronics. In this respect, intense efforts have been focused on the production of metal nanorods and nanowires due not only to their essential role in the fabrication of new devices but also because they offer opportunities to study key fundamental aspects such as quantized conductance or new plasmonic effects. Bulk silver exhibits the highest electrical and thermal conductivity among all metals and silver nanoparticles of different sizes and shapes exhibit surface plasmon resonances that can be well resolved over interband transitions and tuned in the visible spectrum range. It is thus a very suitable metal case for studying both the fundamental and technological aspects of elongated and oriented nanostructures and there have thus been several reports on the production of silver nanorods or nanowires.7–14 Different methods have been developed to prepare such metal elongated nanostructures most of them using chemical routes (thermal,8 hydrothermal,10 or oxide reduction,1,7,11,12,14) or using templates (i.e., mesoporous silica8 or alumina3,4). They generally lead to randomly oriented nanowires that have a diameter smaller than 100 nm, lengths of a few micrometers or longer, and are in most cases in colloidal suspensions. However, for many of the envisaged applications, these elongated nanostructures need to be oriented and supported on substrates or embedded in matrices. Another important challenge that needs to be faced is the control of the shape and dimension dispersion in order to keep it low enough. There have been some works reporting contributions in this direction using multi-step approaches based on the use of electron beam lithography.

In this letter, we report a new single step method using pulsed laser deposition (PLD) to produce oriented Ag nanowires embedded in a matrix and having low dimension and shape dispersion. PLD has already been proven to be very suitable for the growth of complex oxide films.16 It is based on focusing nanosecond laser pulses from a laser onto a target material typically in the ultraviolet (UV) and having nanosecond pulse duration. The use of two independent targets of an oxide (for the matrix) and of a metal have successfully been applied to produce high quality nanocomposite films formed by metal nanoparticles (NPs) embedded in amorphous aluminium oxide (α-Al2O3).17,18 Figure 1 shows cross section schematics of one of those films that consist of “layers” of metal NPs of diameter D and whose spacing can easily be controlled through the deposition of the matrix, X being the equivalent thickness of the matrix layer. The technique for the deposit of the nanowires is an extension of this approach since it is based on the deposition of a limited amount of the matrix by decreasing X down to values close to D. The matrix acts then as a template that allows an oriented growth of the nanowires and the sequence is repeated a number of times n to lead to the situation sketched in Fig. 1. The resulting nanowires are oriented perpendicular to the substrate and the method allows to deliberately selecting their length through the number of cycles. Both features are not easily accessible via the chemical methods cited above or require a multistep approach such as in the case of lithography based techniques.

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FIG. 1. (Color online) Schematics for the films growth and deposition sequence to produce NCls.
An ArF excimer laser [full width at half maximum (FWHM) pulse duration=20 ns, repetition rate of 5 and 20 Hz for the Ag and the Al2O3 targets, respectively], with an average energy density of 1.9 J cm−2, was sequentially focused on the surface of high purity Al2O3 and Ag rotating targets. The films were grown in vacuum (10−6 mbar) on rotating substrates held at room temperature placed 32 mm away from the target surface. The films are produced on Si substrates covered by their native oxide held at room temperature. From the deposition rates of Ag and the Al2O3 required to produce NPs of a certain diameter and matrix layers of a defined thickness, already known from earlier works, we have selected our experimental conditions to produce nanowires with typical dimensions of an average diameter $D = 2 \text{nm}$ and a matrix layer thicknesses ranging from $X = 2D$ to $X = 0.5D$. The number of cycles was fixed to 5, and thus the produced nanostructures have a small aspect ratio when compared to the nanostructures typically referred to as nanowires in the literature. This feature together with the fact that they are all oriented along the direction perpendicular to the substrate, make the present nanostructures to be nanocolumns (NCls) rather than nanowires.

The morphology and structure of the films have been studied by transmission electron microscopy in a Philips Tecnai 20F FEG, transmission electron microscopy (TEM)/scanning TEM (STEM) analytical microscope operating at 200 kV with point-to-point resolution of 0.2 nm and equipped with an electron dispersive X-ray (EDX) detector, and a dark-field high angle annular detector for Z-contrast analysis. Composition mapping was performed in STEM-EDX mode. Both plan view and cross-sectional specimens have been prepared by mechanical polishing followed by ion milling in a liquid-nitrogen cooled holder at 3 kV and final angle of 8°. The silicon substrate in the cross section specimen has been used to perfectly orient the specimen perpendicular to the electron beam as well as to calibrate the scale of the images.

Figure 2 shows cross-section TEM images of films produced with different $X$ values. When $X = 2D$, no columnar growth is expected to occur as it is indeed seen in the image. Instead, it shows the excellent control achieved in the growth process by PLD since 5 dark “layers” of NPs can be well resolved. The average diameter of the NPs ($D = 1.9 \pm 0.2 \text{nm}$) as well as the spacing of the NPs “layers” ($X = 5.0 \pm 0.2 \text{nm}$) agrees well with what was intended. When $X = D$, elongated dark areas perpendicular to the substrate and thus oriented along the growth direction are observed. Figure 2 also shows that these elongated structures are well separated one from another for $X = D$ and oriented perpendicular to the substrate. The length or height $H$ of these elongated structures is plotted in Fig. 3 as a function of $X$, where it seems that $H$ undergoes a maximum of $7 \pm 0.2 \text{nm}$ for $X$ in the neighborhood of $D$. Figure 3 also shows a high resolution image of one of the NCls obtained for $X = D$ where lattice fringes are observed evidencing their crystalline structure. Fourier transform patterns of the NCls indicate that they are not single crystals. This is also seen in Fig. 3 where regions having fringes with different orientations along the NCIs are clearly visible.

Figure 4 shows plan view images of the specimen grown with $X = D$ taken in TEM and STEM mode with EDX analysis for X-rays and high angle annular detector for Z-contrast images, the latter showing the areas having large concentrations of Ag as bright areas. The good match of dark and bright areas in both images together with the X-ray fluorescence spectra confirms that isolated silver NCIs have been produced in the direction perpendicular to the substrate. The NCIs have an approximately round section with a very homogeneous diameter of $2.7 \pm 0.2 \text{nm}$ as it shown in the histogram included in Fig. 4.

When depositing a metal on an oxide surface, it is well known that it tends to follow the Volmer—Weber three-dimensional growth due to the lower surface free and interface energies of the metal with respect to that of the substrate. In addition, pulsed laser deposition (PLD) is a process involving a large ion to neutral species ratio and it is also well known that the presence of such charged species creates nucleation centers at the substrate. A high number density of NPs are thus homogeneously nucleated over the whole sub-
strate at the early stages leading to NPs dimensions close to 2 nm for the case of Ag (Ref. 17) or Au (Ref. 18) before coalescence starts. This stage is illustrated in Fig. 2 for X = 2D. To generate NCls, the production of this first “layer” of metal NPs is followed by the deposition of a certain amount of the matrix material, enough to fill-in the space among the NPs while leaving uncovered part of the metal NPs. The matrix thus acts as a mask defining a template where the next ablated metal self-organizes driven by the higher metal-metal than metal-oxide adhesion energy and the high mobility of the atoms at the surface due to the high kinetic energies involved in the PLD process. This self-organization process leads to the production of NCls when the process is repeated a number of times. The first layer of NPs constitutes thus the seeds for the NCls. Since it has been reported that the number density of NPs decreases as the amount of metal deposited per layer increases due to coalescence, the number density of NCls, their diameter and thus, their separation, can potentially be controlled by varying the amount of metal deposited in the first cycle (number of laser pulses in the metal target), the amount of metal deposited in subsequent cycles and the number of cycles, respectively. These results open new possibilities for material nano-engineering since they provide a simple route based on a single step process at room temperature to produce oriented metal nanocolumns with diameters typically <10 nm and with low dimension and shape dispersion, features that are not easily accessible for other techniques.

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by the number of cycles and thus only limited by the stability of the production system.

In summary, the preparation of parallel Ag nanocolumns embedded in an oxide host by thin-film deposition technologies has been demonstrated, these features being essential for applications in integrated nanoelectronics or photonics. The number density, diameter and height of the nanocolumns can potentially be controlled by varying the amount of metal deposited in the first cycle (number of laser pulses in the metal target), the amount of metal deposited in subsequent cycles and the number of cycles, respectively. These results open new possibilities for material nano-engineering since they provide a simple route based on a single step process at room temperature to produce oriented metal nanocolumns with diameters typically <10 nm and with low dimension and shape dispersion, features that are not easily accessible for other techniques.

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