Surface Effects on the Mechanical Elongation of AuCu Nanowires: De-alloying and the Formation of Mixed Suspended Atomic Chains

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Abstract

We report here an atomistic study of the mechanical deformation of Au$_x$Cu$_{1-x}$ atomic-size wires (NWs) by means of high resolution transmission electron microscopy (HRTEM) experiments. Molecular dynamics simulations were also carried out in order to obtain deeper insights on the dynamical properties of stretched NWs. The mechanical properties are significantly dependent on the chemical composition that evolves in time at the junction; some structures exhibit a remarkable de-alloying behavior. Also, our results represent the first experimental realization of mixed linear atomic chains (LACs) among transition and noble metals; in particular, surface energies induce chemical gradients on NW surfaces that can be exploited to control the relative LAC compositions (different number of gold and copper atoms). The implications of these results for nanocatalysis and spin transport of one-atom-thick metal wires are addressed.

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I. INTRODUCTION

Predicting the mechanical behavior of a strained nanoscale volume of matter is essential for many nanotechnological applications [1]. This has stimulated an intense study of mechanical elongation of atomic-size metal nanowires (NWs) [2]. In this range size (∼1-2 nm in diameter), surface energy plays a dominant role and, factors that are neglected in macroscopic theory, such as size and shape, determine deformation mechanisms. For example, surface energy can induce strengthening and asymmetrical mechanical response [3–6]. In addition, the high surface/volume ratio (SVR) may lead to the generation of anomalous helicoidal or tubular nanostructures during deformation [7, 8].

Alloying or doping are routinely utilized to improve the mechanical resistance of metals (solute strengthening) [9]. However, these manipulations are very difficult to apply to nanosystems due to the huge SVR, which may promote composition gradients, or even the expelling of impurities [10–12]. In addition, most of our knowledge on metal alloy nanosystem is associated with heterogeneous catalysts, where the use of alloy nanoparticles (NPs) represents an active research field. Nanoscale mechanical deformation of alloys represents a quite complex topic, as the constant injection of elastic energy into the system may be relaxed through a wide variety of structural, physical and chemical mechanisms. We also must keep in mind that the analysis of compositional gradients and segregation in alloy metal nanoparticles in heterogeneous catalysis still represent a question that awaits for a reliable answer. A recent cutting edge study reports the analyses of composition gradients in metal nanoparticles exploiting X ray TEM tomography [13]. In this way, this research remains quite challenging, as we must analyze the complex interplay between elastic, electronic and surface energy contributions.

Here, we present a detailed study of atomic structure evolution of Au-Cu alloy NWs following tensile deformation by means of high resolution electron microscopy (HRTEM). Molecular dynamics simulations were also carried out in order to analyze the dynamics of atomistic processes involved in the nanoalloy physical and chemical modifications.
II. METHODOLOGY

We have generated metal NWs from alloy bimetallic films \((\text{Au}_x\text{Cu}_{1-x})\) \((0 < x < 1)\) following the experimental procedure introduced by Takayanagi’s group \([15]\). Initially, holes are opened at several points in a self-supported metal film by focusing the microscope electron beam \((300\text{A/cm}^2)\); in this manner, nanometric constrictions (bridges) are formed between them. Then, the microscope beam current density is reduced to standard operation values \((1030\text{ A/cm}^2)\) for image acquisition; in this range of beam current density, the HRTEM sample temperature is estimated to be within \(300 - 350\text{ K}\) \([16]\). The spontaneous elongation and rupture of the nanowires is acquired using a high-sensitive TV camera (Gatan 622SC, 30 frames/s) and a standard video/DVD recorder. It is important to emphasize that this experimental procedure allows the acquisition of time-resolved atomic-resolution-imaging of NWs with a remarkable quality; nevertheless, it is not possible neither to measure the force being applied nor to control the stretching direction \([17]\). Usually the NW stretching and fracture occurs with average displacement rates of 0.1-1 nm per second. The worked described here used bimetallic \(\text{Au}_x\text{Cu}_{1-x}\) alloy thin films as initial sample to generate NWs in situ in the HRTEM (JEM-3010 URP 300 kV, 0.17 nm point resolution).

Polycrystalline \(\text{Au}_x\text{Cu}_{1-x}\) lms \((30 - 50\text{ nm in thickness})\) have been prepared by thermal co-evaporation in a standard vacuum evaporator \((10-7\text{ mbar})\). A quartz crystal monitor was used to set the evaporation rate of each individual metal source and, subsequently, to measure the equivalent thickness of the film. Owing to higher cooling rates associated with the film deposition the bimetallic films are expected to consist of a solid solution with random distribution of gold and copper atoms \([18]\), what was confirmed by micro-electron diffraction results. To prevent possible oxidation by exposure to ambient conditions, the bi-metallic films were sandwiched between two \((3\text{-nm-thick})\) amorphous carbon thin layers. Before generating the NWs, the carbon layers are removed by strong electron irradiation \([19]\) inside the HRTEM. The structural characterization has been performed by means micro-electron diffraction (JEM 2100 ARP, operated at 200 kV). In our experiments, the electron diffraction patterns (DP) were acquired from a region of \(800\text{ nm in diameter}\) and recorded using a CCD camera (Gatan ES500W) (more details see Figure S1 in the Supplementary Material). We have also measured the chemical composition of synthesized alloy films using Energy-Dispersive X-ray Spectroscopy (EDS); the \(\text{Au}_x\text{Cu}_{1-x}\) alloy films were supported
over conventional molybdenum TEM grids to avoid spurious x-ray signal. In particular, we analyzed several localized regions inside the initial illuminated area used for electron diffraction studies and, the observed atomic composition variations were within the typical composition error bar (5%, using Cliff-Lorimer method without absorption correction [16]). Also, the measured compositions were in very good agreement with electron diffraction estimation using Vegards law. However, we have observed significant composition changes when comparing measurements performed in pristine alloy thin films and, after the in-situ formation of NWs (i.e. after intense electron beam irradiation during several hours). It is important to highlight that the electron irradiation necessary to prepare the metal alloy film for a NW study requires a several hours long electron beam irradiation. In contrast, the final NW elongation and rupture processes recorded by the experimental videos last, at most, $3 - 5$ minutes at a much lower electron beam intensity (see description of electron microscopy works in Section 1.a of Supplementary Materials); then, it is reasonable to think that no significant chemical composition change occurs during the nanowires imaging study. In this sense, we have assumed that the final EDS estimated concentration is a good value to describe the NW composition, and it has been used to describe the nanowires.

We have also carried out molecular dynamics simulations to gather deeper insights on the atomistic processes occurring during the alloy NW elongation. A tight-binding molecular dynamics methodology based on the second-moment approximation (TB-SMA) [20, 21] was used to analyze the elongation structural evolution. The theoretical methodology has already been described in detail by Sato et al. [22]; this approach has proved to be very effective for the study of Au and Cu NWs [19, 23, 24].

III. RESULTS

The in-situ HRTEM experiments indicate that Au$_x$Cu$_{(1-x)}$ alloy NWs can deform along only three [111], [100] and [110] crystallographic directions. NWs elongated along [111] and [100] directions generate by-pyramidal constrictions that always evolve into an atomic contact or linear atomic chains (LACs); this behavior is identical to pure Au or Cu wires [3, 24]. In contrast, alloy NWs stretched along the [110] direction (hereafter noted as [110] NWs) display a concentration dependent structural behavior. While [110] Au NWs display rod-like morphology and break abruptly when formed by 3-4 atomic layer thick [3, 23], alloy
NWs (both Au$_{0.55}$Cu$_{0.45}$ and Au$_{0.2}$Cu$_{0.8}$) display a structural sequence typical of pure Cu wires [19, 24]: 1) rod-like wire; 2) by-pyramidal constrictions and, 3) a final one-atom-thick contact or LAC formation (Figures 1b,c, and corresponding videos in the SM). This indicates that a Cu content of $\sim 45\%$ is enough to trigger typical copper behavior, consequently modifying the rupture mode from brittle to ductile. These results are quite different from the Au$_x$Ag$_{(1-x)}$ NW case, where a much higher Ag content ($\sim 80\%$) was necessary to reveal silver nanowire characteristics [11].

From a mechanical point of view, it is important to analyze the active deformation mechanisms of atomic-size alloy wires. Concerning bulk material, Au and Cu are Face Centered Cubic (FCC) metals and, plastic deformation occurs mostly by the gliding of compact (111) atomic planes along [112]-type directions. In particular, partial edge dislocations (PDs) are formed, which encapsulate a stacking fault (SF) ribbon. In tiny gold nanorods, where diameter ($\sim 1$ nm) is smaller than the SF ribbon width, $((d) \sim 2$-$3$ nm in bulk [9]), plastic deformation occurs by the formation of planar defects that generate a compact glide (block on a block) of the (111) planes by $(1/6)[112]$ [5, 25]. In these very tiny wires thermal energy at room temperature is enough to recombine these planar faults [5, 26] and, pure Au and pure Cu NWs stay defect free when stretched at 300 K [3, 24]. Concerning macroscopic alloys, it is well known that alloying influences drastically, the elastic modulus and yield strength [9]. On this basis we could expect that, in Au-Cu alloy NWs, energy barrier blocking planar defects should be higher. In fact, we have observed the formation of planar defects at room temperature in some NWs with Au$_{0.55}$Cu$_{0.45}$ composition (Figure 2a,b). Nevertheless, many alloy NWs also displayed defect free structures; this may be associated with subtle local variations of chemical composition inside the alloy thin film or, be even induced during the wire elongation [11]. Figures 2c and 2d show some interesting images of Au$_{0.20}$Cu$_{0.80}$ NWs. Note several darker dots at the NW apexes. This might be associated with the formation of several small gold clusters during the NW elongation (gold atoms are expected to generate darker dots in the images). Accordingly, our video recordings show that these clusters move slowly during the mechanical elongation, suggesting that they may be located on the NW surface (see Supplementary Material (SM)). In fact, several theoretical studies of Au-Cu nanoparticles have predicted the migration of gold atoms to the surface in order minimize surface energy [10, 29]. Certainly, the gold lower surface energy [30] and lower diffusion barrier drive gold atoms migration to the wire surface during the mechanical deformation.
Molecular dynamics simulations can provide additional insight into atom reorganization and redistribution during alloy NW elongation. Figure 3a shows a sequence of snapshots of the stretching of Au$_{0.5}$Cu$_{0.5}$ NW along [110] direction. Initially, a by-pyramidal constriction is formed, in good agreement with experimental observations; then, a long NW is generated. It is important to emphasize that most of the NW gold atoms are located on the NW surface, the inset shows a cross-sectional view where it is clear that Au atoms enclose a chain of Cu atoms. Finally, a one-atom-thick contact is formed before breaking. During the elongation, a clear gold enrichment of the narrowest wire regions can be measured (see also data in the SM). Figure 3b shows a similar Au surface migration effect in Au$_{0.2}$Cu$_{0.8}$ NW stretched along [110] axis. However, the behavior changes, because the initial Au content is rather low (20%) and, there are not enough available gold atoms to cover the whole surface [10]. Finally, Figure 3c illustrates an Au$_{0.2}$Cu$_{0.8}$ NW being elongated along the [100] direction, which becomes gradually thinner until forming an atomic contact. The formation of small gold clusters (3-5 atoms) on the NW surface can be clearly observed. The segregated gold clusters remain coalesced and sometimes diffuse slowly on the NW surface.

Briefly, the simulations have revealed two effects associated with local composition changes: (i) surface segregation of Au atoms and, (ii) gold clustering. These effects are in excellent agreement with the experimental observations displayed in Figures 1 and 2. However, we must keep in mind that electron beam induced effects that may also influence atom diffusion or segregation. We think that these effects are negligible in our experiments due to the rather short duration of the experiments (less than 1 minute for the elongation and rupture of the 1-nm-wide alloy wires, see a detailed discussion in the SM).

From a more fundamental point of view, the formation of one-atom-thick nanowires containing different metal atomic species, represent one of the most interesting nanosystems to study 1-D quantum physics. So far, mixed suspended linear atom chains (LACs) were experimentally produced only with gold and silver [11]. Mixed LACs containing gold and transition metal atoms open the possibility to address excellent physical and chemical questions such as nanomagnetism, spin transport, s-d bonding in low dimensional systems, etc. From this perspective, the one-atom thick wire generated from the Au-Cu alloy nanowires observed in our study represent an excellent case study. Our HRTEM results indicate that suspended chains display variations in the intensity/contrast at atomic positions (see Figure 1c and SM), suggesting that they should be formed by both gold and copper atoms.
A quantitative comparison between experimental and simulated HRTEM image intensities confirms this interpretation. The interatomic distances in the LAC are in the [0.25-0.32] nm range, what agrees with impurity-free gold and copper chains. Light possible impurity atoms such as C, O, N should produce a much lower contrast than the signal noise ratio experimentally observed (see details in SM).

A natural question arises, can we control the LAC chemical composition by selecting the proper alloy, wire shape, elongation direction, etc.? Previous theoretical studies have shown that most of the atoms composing suspended chains come from the outermost layers for Au nanowires. With this idea in mind, we have analyzed our theoretical simulations and looked at the chemical composition (pure Au or Cu, or alloyed chains) of LACs generated along different stretching directions for different alloy compositions (Au$_{0.5}$Cu$_{0.5}$ and Au$_{0.2}$Cu$_{0.8}$). Most of suspended atomic chains were composed by two hanging atoms, while seldom rather long chains (4-5 atoms) were observed. Although, the rather low available statistics, a clear alloy composition and elongation direction dependence shows up (see results in Table 1). LACs generated from [100] NWs show the tendency to be either pure Au or pure Cu depending on the alloy mixture. In contrast, mixed LACs dominate the occurrence along [111] axis for both studied alloys. Finally, [110] alloy nanowires show a slight tendency to produce pure Au chains, followed by alloyed chains.

In order to understand these simulation results, we must first consider that alloy composition influences NW morphologies through changes in the surface energy of the different crystallographic facets. A gold particle should be a cuboctahedra with regular hexagonal facets (minimal energy planes are \{111\}\[10\], schema at left in Figure 4a)), while Cu nanoparticle have a morphology dominated by \{100\} facets (center and right octahedra in Figure 4a). In first approximation, an alloy nanosystem behavior must be somewhat in between these two extrema. In addition, as diffusion and migration is enhanced in nanosystems, it is reasonable to make the hypothesis that the wires will have a spontaneous tendency to have composition gradient in the volume and on the surfaces. In particular, we can assume that \{hkl\} facets will accommodate more atoms of the chemical species that minimize the facets surface energy, what would lead to Au (Cu) rich \{111\} (\{100\}) facets in an Au-Cu alloy NW (or NP). A gold-rich-alloy NW along [100] should have a pyramidal shape of square base defined by four triangular \{111\} facets (see left side in Figure 4b). These facets will become Au rich during elongation, what finally it will enhance the for-
formation of pure Au LACs. In contrast, a Cu rich [100] NW should display a rod-like shape with a square cross section, the surface being formed by four \{100\} facet (see right side in Figure 4b), then having a tendency to generate mostly pure Cu LACS. Wires along [110] direction will have a hexagonal cross section formed by both \{111\} and \{100\} facets, with a relative weight that varies from Au to Cu; this explain why pure Au and mixed chains are formed along [110] elongation axis. Finally, alloy NWs formed along [111] will generate at some moment a triangular cross section (see for example the \{111\} facet shape of octahedra at the center of image 4a). This triangular facet is surrounded by 3 \{100\} (Cu rich) facets and the tip of 3 \{111\} (Au rich) facets, what can provide both Au and Cu atoms. This kind of wire must be expected to generate mostly mixed LAC’s, as in fact, we observed in the simulations.

In summary, we have observed that alloy $\text{Au}_x\text{Cu}_{(1-x)}$ NWs show a strong concentration dependence mechanical behavior. Approximately a $\sim 45\%$ Cu content is required to trigger copper-alike mechanical behavior. For the tiny alloy NWs studied here ($\sim$nm in diameter) surface energy contribution is so important that can induce gold enrichment and, even gold surface segregation during elongation. The formation of suspended atom chains containing Au and Cu atoms was experimentally revealed. Molecular dynamics simulations suggest that it is possible to control the LAC chemical composition by choosing the appropriate alloy composition and NW elongation direction. This would exploit the spontaneous formation of chemical composition gradients (or preferential chemical enrichment of each family of crystallographic facets) on the NW surface. This phenomena can certainly be expected to also happen in alloy nanoparticles and, may modify significatively the reactivity and/or catalytic activity in heterogeneous catalysis. From another point of view, the possibility to generate in a reasonable controlled way alloy LACs containing Au and magnetic transition metals may open new opportunities for the study scattering and spin transport in one-atom-thick metal wires.

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TABLE I. Statistical analysis of LAC formation from the molecular dynamics simulations. The three numbers (X/Y/Z) indicate the number of LAC formed and composed only of gold, mix Au/Cu or copper, respectively.

|         | [100] | [110] | [111] |
|---------|-------|-------|-------|
| Au<sub>x</sub> Cu<sub>y</sub> | 50/50 | 6/3/0 | 6/4/0 | 2/6/2 |
|         | 80/20 | 2/2/7 | 4/3/2 | 3/7/1 |

FIG. 1. Sequence of atomic resolution images associated with the elongation and thinning of rod-like [110] Au-Cu NWs as a function of chemical composition (atom positions appear dark). (a) Au<sub>0.55</sub>Cu<sub>0.45</sub> and (b) Au<sub>0.2</sub>Cu<sub>0.8</sub> NWs form by-pyramidal constrictions and evolve into either suspended atomic chains or atomic contact. (c) closer view of suspended linear atomic chain generated from a Au<sub>0.55</sub>Cu<sub>0.45</sub> NW; we can see that the atom at the left of the LAC (arrowed) display a different contrast from the other suspended atoms suggesting the formation of mixed Au-Cu chains.

FIG. 2. Typical HRTEM images of Au<sub>0.55</sub>Cu<sub>0.45</sub> NW’s displaying planar defects (a) and twins (b). Defects are indicated by black arrows and, they can also be identified by a discontinuity in the atomic planes. (c,d) HRTEM images of stretched Au<sub>0.2</sub>Cu<sub>0.8</sub> NWs which indicate the formation of small gold clusters in the NW surface shown inside the ellipses. Atom positions appear dark.

FIG. 3. Sequence of snapshots associated with the theoretical simulations of bi-metallic Au-Cu nanowire elongation. Gold and copper atoms are represented by yellow and dark gray balls, respectively. Figures (a) and (b) correspond to the elongation along [110] axis of Au<sub>0.55</sub>Cu<sub>0.45</sub> and Au<sub>0.2</sub>Cu0.8 NWs, respectively. Note that in both cases gold atoms have a tendency to occupy the NW surface. Figure (c) illustrates the formation of small gold clusters (3-5 atoms) on the NW surface (indicated by arrows) during the elongation of a Au<sub>0.2</sub>Cu0.8 NW along [100] axis. The corresponding animations can be found in the SM.
FIG. 4. a) From left to right, shapes of cuboctahedral FCC nanoparticles increasing the relevance of \{100\} facets in relation to \{111\} facets. b) Scheme of the possible [100] alloy NW; a pyramidal shape containing only \{111\} surfaces, expected for a gold rich wire and, a rod-like wire of square cross-section expected for Cu rich NW (only \{100\} facets cover the surface). (c) Qualitative hexagonal cross-section for a pillar-like wire expected along [110] direction, note that this rod is formed by both \{111\} and \{100\} facets (see text for discussions).
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