High-Throughput Direct Writing of Metallic Micro- and Nano-Structures by Focused Ga\textsuperscript{+} Beam Irradiation of Palladium Acetate Films

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ABSTRACT: Metallic nanopatterns are ubiquitous in applications that exploit the electrical conduction at the nanoscale, including interconnects, electrical nanocontacts, and small gaps between metallic pads. These metallic nanopatterns can be designed to show additional physical properties (optical transparency, plasmonic effects, ferromagnetism, superconductivity, heat evacuation, etc.). For these reasons, an intense search for novel lithography methods using uncomplicated processes represents a key on-going issue in the achievement of metallic nanopatterns with high resolution and high throughput. In this contribution, we introduce a simple methodology for the efficient decomposition of Pd\textsubscript{3}(OAc)\textsubscript{6} spin-coated thin films by means of a focused Ga\textsuperscript{+} beam, which results in metallic-enriched Pd nanostructures. Remarkably, the usage of a charge dose as low as 30 $\mu$C/cm\textsuperscript{2} is sufficient to fabricate structures with a metallic Pd content above 50\% (at.) exhibiting low electrical resistivity (70 $\mu$Ω·cm). Binary-collision-approximation simulations provide theoretical support to this experimental finding. Such notable behavior is used to provide three proof-of-concept applications: (i) creation of electrical contacts to nanowires, (ii) fabrication of small (40 nm) gaps between large metallic contact pads, and (iii) fabrication of large-area metallic meshes. The impact across several fields of the direct decomposition of spin-coated organometallic films by focused ion beams is discussed.

KEYWORDS: focused ion beams, spin-coated organometallic films, electrical contacts, nanogap electrodes, large-area meshes

INTRODUCTION

Metallic nanopatterns are important building blocks in modern technology with fundamental relevance in the construction of interconnects for microelectronics\textsuperscript{1}, conductive nanomeshes for organic photovoltaics\textsuperscript{2} and for transparent conductive electrodes\textsuperscript{3}, organic light-emitting diodes for flexible electronics\textsuperscript{4}, plasmonic nanostructures for sensing\textsuperscript{5} and for lithography\textsuperscript{6}, nanogap electrodes for optical applications\textsuperscript{7} and for molecular electronics\textsuperscript{8}, electrical contacts to nanowires for sensing\textsuperscript{9} and for quantum technologies,\textsuperscript{10} and so forth.

The existing fabrication methods of these metallic nanopatterns are based either on chemical or physical approaches or on hybrid physical/chemical methodologies, with each technique leading to different features in terms of resolution, throughput, cost, and so forth.\textsuperscript{11,12} Whereas bottom-up chemical methods (such as those relying on self-assembly and self-organization) generally excel in throughput and cost,\textsuperscript{13,14} top-down lithography techniques based on physical techniques (such as optical/electron/ion beam lithography in combination with thin-film techniques) allow for the control of the dimensions of the nanopattern with great accuracy.\textsuperscript{15,16}

Importantly, hybrid approaches (combining chemical and physical techniques) can benefit from the virtues of both, with two examples being nanosphere lithography and directed block-copolymer lithography.\textsuperscript{17,18} Without pretending to be comprehensive, a few paradigmatic examples of advanced nanofabrication methods to create metallic nanopatterns can be discussed. For instance, mature multi-step techniques based on resists, such as optical lithography,\textsuperscript{19} electron-beam lithography (EBL),\textsuperscript{20} and nanoimprint lithography,\textsuperscript{21} are capable of patterning metals with a high resolution and throughput. Direct patterning of metals using a focused ion beam (FIB)\textsuperscript{22} or by scanning probe lithography\textsuperscript{23} has demonstrated high-resolution capabilities, albeit at the expense of low throughput. Moreover, less conventional approaches
have been used in specific applications, such as the fabrication of ultra-small gaps through sketch-and-peel lithography combined with transfer printing24 and three-dimensional metallic nanoarchitectures by means of two-photon lithography25 or focused electron beam-induced deposition.26

Without undermining the power of such advanced methods to fabricate metallic nanopatterns, it is worth exploring other nanofabrication methods that could combine virtues such as simplicity, high resolution, and high throughput. In this context, the direct decomposition of organometallic films by focused charged beams is an interesting route to explore. The advantage of this technique lies in the fact that after spin coating of the organometallic film and the subsequent electron or ion irradiation, the metallic (Pd, Ag, Au, Ir, etc.) nanopattern is readily revealed by dissolving non-irradiated areas in a cleaning solvent.27−31 Thus, no sacrificial resist layer is necessary and, in the end, the organometallic film precursor becomes the functional material. However, in all cases reported in the literature so far, a post-processing annealing step is required to obtain low electrical resistivity of the nanofabricated material.31−35

Recently, our group has shown that by irradiating a Pd₃(OAc)₆ thin film with a high electron dose, it is possible to produce metallic nanostructures without the need of post-processing annealing steps.36 Unfortunately, the high dose required (30,000 μC/cm²) limits the applications of this approach to small-area patterning or low-throughput device fabrication. Motivated by the search for a high-throughput and annealing-free patterning process of Pd₃(OAc)₆ thin films, we have investigated whether Ga⁺ FIB irradiation could be effective toward a decrease in the required charge dose without putting in jeopardy the low electrical resistivity of the nanostructure. An illustration of the work here presented is summarized in Figure 1.

Remarkably, we have found that only 30 μC/cm² Ga⁺ irradiation is required to directly obtain metallic structures with a resistivity of 70 μΩ·cm. Importantly, this finding represents a substantial improvement with respect to all previous existing results and opens the route to large-area and high-throughput device applications. In order to put this result in context, it is worth mentioning that standard EBL processes make use of PMMA resists that need irradiation doses up to a few hundred μC/cm²,37 and FIB-induced deposition of metals (Pt, W, and Co) requires irradiation doses in the range of 10,000 μC/cm².38

For clarification purposes, this article is organized as follows. First, the optimization of the ion irradiation procedure employed to obtain functional Pd nanostructures with low electrical resistivity will be presented. Second, simulations of the Ga⁺ irradiation process that decomposes the palladium acetate film will be provided. In the last part of the article, three applications of this nanolithography technique will be shown: (a) fabrication of electrical contacts to nanowires; (b) fabrication of small gaps between metal contacts, which in turn will serve to test the resolution of the technique; and (c) fabrication of conductive large-area meshes.

**Experimental Details.** Ga⁺ FIB irradiations were carried out in a Helios NanoLab 650 (FEI company) on palladium acetate films with thicknesses in the 100−350 nm range; the palladium acetate films were previously deposited by spin coating onto Si/SiO₂ substrates36 (see the sketch in Figure 1). Spin-coated films with thicknesses in the range of 100 to 200 nm were prepared by spreading 10 μL of a 0.09 M Pd₃(OAc)₆ filtered solution in chloroform. Furthermore, those with thicknesses in the range of 200 to 350 nm were prepared by spreading 15 μL of 0.2 M filtered solution. The spin-coating process in both cases includes two sequential steps: (1) 10 s at 3000 rpm and (2) 40 s at 4000 rpm. If the spin-coated film is too thick, some lift off and compositional problems could appear due to the inefficient decomposition of the bottom part of the film. The optimized parameters were 30 kV of the ion beam voltage, an ion beam current of 1.1, 7.7, or 24 pA (depending on the irradiation area), 200 ns of dwell time, and 0% of overlap. The organometallic films were decomposed by applying Ga⁺ doses in the 2−100 μC/cm² range. After the developing step in chloroform, the resulting Pd nanostructures were characterized, and the optimal dose was chosen in terms of final application, which requires good electrical conductivities. Micro/nano pinholes have been found over the nanostructures. These have been associated with two
contributing effects: (1) possible small bubbles/holes forming during the spin-coating procedure due to CHCl₃ evaporation and (2) to the path created by the volatile components while being sublimated due to the beam exposure. Atomic force microscopy (AFM Veeco-Bruker Multimode 8) images of 25 μm² microstructures fabricated at 20 and 30 μC/cm². These images have been analyzed by using the Nanoscope V.1.40 software, obtaining a root-mean-square roughness value in the 0.9–1.7 nm range for pinhole-free smaller areas (see Figure S1).

Current-versus-voltage (I-V) measurements were performed at room temperature inside a Helios NanoLab 650 Dual Beam instrument (FEI company) using electrical microprobes (Kleindiek Nanotechnik GmbH) placed inside the chamber and a 6221 DC current source/2182A nanovoltmeter (Keithley Instruments) connected to the microprobes via a chamber feedthrough.

High-angle annular dark-field (HAADF) imaging and energy-dispersive X-ray spectroscopy (EDS) measurements were carried out along the nanostructure thickness by transmission electron microscopy (TEM) in an analytical Titan low-base instrument (FEI Company). HAADF images were obtained at 300 keV, and the energy resolution of the EDS experiments was ∼125 eV using the scanning transmission electron microscope (STEM) mode.

X-ray photoelectron spectroscopy (XPS) was performed on a Kratos Axis Ultra DLD spectrophotometer with a monochromatic Al Kα X-ray source (1486.6 eV) and a pass energy of 20 eV. Data treatment and further analysis were performed using the CasaXPS v.2.3.15 software. Binding energies have been referenced to the C 1s peak at 284.9 eV. Peak fitting was conducted by using standard line shapes GL(30) and LA(1.9,7, 2) for the asymmetric signal associated with metallic palladium.

Simulations based on binary collision approximation (BCA) were carried out using the SDTrimSP code⁴⁹ to model the evolution of the precursor layer under Ga⁺ ion irradiation. This means that at a given time, only the interaction between two atoms is taken into account and many-body effects are neglected. The interatomic interactions were calculated using the KrC potential, and the electronic stopping was described by the Oen-Robinson model. Charges on atoms are not explicitly considered. The Gauss-Mehler method with 16 pivots was used for integration. The surface binding energy was calculated using the equation \( \text{sbe}(i, j) = 0.5 \times (E_i + E_j) \), where sbe is the surface binding energy of the current target, and \( E_i \) and \( E_j \) is the atomic surface binding energy of the chemical species i. The surface binding energy was calculated for any combination of two species used in this work, for example, gallium, silicon, palladium, carbon, hydrogen, and oxygen. The atomic densities of the different species were taken to be identical to the bulk values. Hence, the density of the simulated precursor layer might be above the value in the experiments. Neglecting the diffusion in SDTrimSP⁴⁹,⁵¹ led to largely overestimated concentrations for carbon, hydrogen, and oxygen (cf. Supporting Information). The Monte Carlo code does not include temperature; therefore the diffusion processes are implemented using Fick’s law, where diffusion is described as a displacement per ion impact and is calculated by taking the ion flux in ions × cm⁻² × s⁻¹ into account. Optimization of the diffusion coefficients resulted in values of \( 10^6 \) Å²/ion for hydrogen and oxygen and \( 2 \times 10^7 \) Å²/ion for carbon. The default displacement energies were used, that is, 12 eV for gallium, 13 eV for silicon, 26 eV for palladium, 25 eV for carbon, and 0.5 eV for hydrogen and oxygen. The precursor layer had an initial thickness of 110 nm. Ga⁺ irradiation was carried out at normal incidence with an impact energy of 30 keV. The maximum dose was set to 160 μC/cm². The generation of secondary electrons is not considered in this approach.
dependence is observed for ion doses higher than 6 \( \mu \text{C/cm}^2 \). According to these data, the palladium content is slightly lower at the dose of 30 \( \mu \text{C/cm}^2 \). Importantly, Ga\(^+\) implantation is negligible due to the low ion current. For the whole range studied except for the lowest doses, the electrical resistivity decreases sharply as the ion dose increases until it stabilizes at higher doses, that is, for the ion dose of 30 \( \mu \text{C/cm}^2 \) and above. In order to carry out structural characterization by transmission electron microscopy (TEM), cross-sectional lamellae of Pd nanostructures fabricated using the STEM mode. First, based on HAADF images (Figure 3a), the as-fabricated nanostructures exhibit very low roughness occurring upon Ga\(^+\) irradiation of the Pd\(_3\)(OAc)\(_6\) thin film. The dark layer on top of the Pd nanostructure corresponds to the interlayer formed between the Pd nanostructure and the Pt-FEBID deposit. The white rectangles indicate the area where the atomic composition was investigated by EDS.

### RESULTS AND DISCUSSION

#### Electrical Characterization by the Four-Probe Technique

In order to optimize the ion dose in relation to the electrical resistivity, current-versus-voltage \((I-V)\) curves at room temperature were recorded. These measurements consist of applying a fixed current through the two outer electrodes while measuring the voltage drop across the two inner electrodes. For that purpose, Pd nanostructures patterned with the design illustrated in Figure 2a were fabricated from a spin-coated film of 110 nm in thickness. Two different ion beam currents were used to cover the full 2–100 \( \mu \text{C/cm}^2 \) range: 7.7 pA for 2, 4, 6, 8, and 10 \( \mu \text{C/cm}^2 \) and 24 pA for 20, 30, 40, 50, 60, 80, and 100 \( \mu \text{C/cm}^2 \). The thickness of each nanostructure was measured by profilometry and is represented in Figure 2b. As represented in Figure 2c, a linear \( I-V \) dependence is observed for ion doses higher than 6 \( \mu \text{C/cm}^2 \), that is, for the whole range studied except for the lowest doses, 2 and 4 \( \mu \text{C/cm}^2 \) (the linear \( I-V \) curve corresponding to the 6 \( \mu \text{C/cm}^2 \) ion dose is not included here for the sake of clarity, but it is shown in Figure S2). The linear fitting of the \( I-V \) curves indicates that the value of the electrical resistance decreases as the ion dose increases and then saturates at higher doses. Combining the electrical resistance together with the dimensions of the Pd nanostructures, the electrical resistivity was calculated and is represented in Figure 2d. This plot shows how the electrical resistivity decreases sharply as the ion dose increases until flattening out at a value of \((70 \pm 5) \mu\Omega\cdot\text{cm}\) for the dose of 30 \( \mu \text{C/cm}^2 \).

#### Compositional Analysis by STEM-EDS and the XPS Study

In order to carry out structural characterization by TEM, cross-sectional lamellae of Pd nanostructures fabricated under Ga\(^+\) doses of 8, 20, and 60 \( \mu \text{C/cm}^2 \) were extracted. HAADF images and EDS measurements were carried out by using the STEM mode. First, based on HAADF images (Figure 3) the as-fabricated nanostructures exhibit very low roughness and absence of voids at analyzed areas. Moreover, a comparison of these images indicates that higher doses result in brighter nanostructures, which is attributed here to the presence of more agglomerated metallic grains. Second, EDS experiments were performed in three different regions for each nanostructure (indicated with white rectangles in Figure 3) in order to investigate their atomic composition.

Atomic contents of Pd, C, and O are collected in Table 1. According to these data, the palladium content is slightly lower for the structure grown with an ion dose of 8 \( \mu \text{C/cm}^2 \), and higher for those grown through doses of 20 and 60 \( \mu \text{C/cm}^2 \). Importantly, Ga\(^+\) implantation is negligible due to the low ion doses used for the fabrication of these nanostructures.

As recently reported, the irradiation of Pd\(_3\)(OAc)\(_6\) thin films with an electron beam produces the decomposition of the organometallic precursor, varying the Pd-oxidation state and therefore the electrical resistivity associated with the resulting nanostructures by up to 2 orders of magnitude. In particular, the reduction of Pd\(^{2+}\) into metallic palladium, Pd\(^0\), occurs when applying a dose of 30,000 \( \mu \text{C/cm}^2 \). To quantify this conversion for Ga\(^+\) irradiated films, the Pd\(^{2+}\) conversion into Pd\(^0\) in ion-irradiated films, an XPS study was performed. The obtained results are gathered in Figure S3. Briefly, when irradiating Pd\(_3\)(OAc)\(_6\) thin films with a Ga\(^+\) beam by directly applying the smallest dose (8 \( \mu \text{C/cm}^2 \)), ~95% of the starting Pd\(^{2+}\) is reduced to Pd\(^0\), and the same applies to the other doses. Combining the information obtained from the electrical measurements and from the compositional analysis, the fabrication of Pd nanostructures for the applications presented below was performed by applying doses in between the optimal range of 20–40 \( \mu \text{C/cm}^2 \). The use of these low ions doses makes negligible potentially focused ion beam side effects such as thermal implications, implantation, or halo effects due to sputtering.

#### BCA-Based Simulations to Model the Evolution of the Precursor Layer

Theoretical calculations were performed to provide further understanding of the processes occurring upon Ga\(^+\) irradiation of the Pd\(_3\)(OAc)\(_6\) thin film. The SDTrimSP simulations show that the average Pd concentration (at. %) increases gradually with the Ga\(^+\) ion dose reaching a maximum value of 85% at the maximum dose of 160 \( \mu \text{C/cm}^2 \) (Figure 4). For a dose of 100 \( \mu \text{C/cm}^2 \), the Pd concentration is equal to 28.5% when considering all elements or equal to 40.4% when considering only Pd, C, and O (i.e., the elements that have also been detected by EDS, see Table S1). In the latter case, a good agreement between the simulations and the experimental results is achieved. The desorption of the volatile compounds and the sputtering of all compounds lead to a decrease in layer thickness. With increasing dose, the sputtering becomes dominated by the partial sputtering yield of Pd (Figure S4). It is also important to note that the definition of the interface position, that is, for a Si

![Figure 3. HAADF images of three selected samples that correspond to Pd nanostructures grown with Ga\(^+\) doses of (a) 8 \( \mu \text{C/cm}^2 \), (b) 20 \( \mu \text{C/cm}^2 \), and (c) 60 \( \mu \text{C/cm}^2 \). During the lamellae preparation, the Pd nanostructure was protected with Pt by Focused Electron Beam Induced Deposition (FEBID). The dark layer on top of the Pd nanostructure corresponds to the interlayer formed between the Pd nanostructure and the Pt-FEBID deposit. The white rectangles indicate the area where the atomic composition was investigated by EDS.](image)

| Ga\(^+\) dose [\( \mu \text{C/cm}^2 \)] | region | Pd [at. %] | C [at. %] | O [at. %] |
|---|---|---|---|---|
| 8 | 1 | 45 ± 0.2 | 50 ± 4 | 5 ± 3 |
| 2 | 46 ± 0.2 | 49 ± 4 | 5 ± 3 | 3 | 46 ± 0.2 | 49 ± 4 | 6 ± 3 | 20 | 1 | 65 ± 0.2 | 28 ± 11 | 7 ± 3 |
| 2 | 59 ± 0.2 | 36 ± 6 | 5 ± 3 | 3 | 45 ± 0.2 | 50 ± 4 | 5 ± 3 | 60 | 1 | 53 ± 0.2 | 41 ± 5 | 6 ± 3 |
| 2 | 52 ± 0.2 | 44 ± 4 | 4 ± 3 | 3 | 48 ± 0.2 | 50 ± 4 | 2 ± 7 |
concentration of 50%, leads to a maximum average concentration of about 20% in this layer. This can be explained by the mixing of other elements, especially H and O, into Si. More information on the composition of the layer and its evolution with irradiation dose can be seen in Figure S5. This value could be affected by the implementation of diffusion into the SdTrimSP code; that is, the diffusion of the light precursor elements into the silicon substrate could be overestimated. According to the layer thickness evolution, the thickness decreases when the Ga⁺ irradiation dose increases, which agrees with the experimental results observed during the characterization of our Pd nanostructures (Figure 2b).

At the beginning of the Ga⁺ irradiation, the composition of the precursor layer is homogeneous with respect to depth. However, the implantation depth of the 30 kV ion beam is not high enough to process the whole layer right from the beginning (Figure 5a), leading to some accumulation of Pd at the sample surface and some modification of the precursor composition at the precursor/Si interface, that is, some accumulation of Pd and C at the interface (Figure 5b). Theoretically, only roughly from a dose of 16 μC/cm², the energy of the Ga⁺ ions is deposited into the whole precursor layer. At the highest dose used in the simulation (Figure 5c), a significant amount of the Ga⁺ ion energy is deposited into the Si substrate. Due to the difference in density, the implantation depth of the Ga⁺ ions decreases with an increasing dose. Detailed information on the sample composition for the different fluences can be found in Figure S5. The higher Ga⁺ doses required to reach Pd concentrations in the 40% range in the simulations compared to the experimental values can be explained by underestimating the ejection of volatile components in SdTrimSP. This has been taken into account to some degree by applying certain diffusion coefficients to H, C, and O. Applying higher diffusion coefficients would not have solved the problem but would have produced even higher Pd concentrations at larger doses (see Table S1), which is contrary to the experimental results where Pd concentrations stay stable over a given dose range (Table 1). The inability of SdTrimSP to reproduce this behavior correctly can be attributed to the binary collision approximation, which takes into account only the interaction between a moving ion or recoil atom and a single target atom without including many-body effects that allow for the description of more complex chemical environments.

As a proof of concept, Pd microstructures were fabricated as top electrical contacts on a Pt nanowire. To achieve this objective, first, a Pt nanowire was directly grown by the FIBID (focused ion beam-induced deposition) technique on a Si/SiO₂ substrate, correctly positioned with respect to pre-patterned alignment marks. Second, after the spin-coating process, the organometallic film was irradiated following four patterns to be used as electrical contacts onto the nanowire.
Finally, non-exposed film areas were removed by immersing the sample in CHCl₃, and the nanowire was measured by the four-probe technique at room temperature. The Pt−C nanowire remained after the spin-coating process, film irradiation, and developing step without appreciating any change in either the morphology or the electrical properties.

Figure 6a shows one of the measured devices where the Pt-FIBID nanowire is artificially colored in orange (dimensions of 30 μm × 100 nm × 30 nm) and the Pd-based electrical contacts in blue. All Pt-FIBID nanowires were grown under the same conditions (30 kV of beam voltage and 1.1 pA of beam current) and electrically measured and are represented in Figure 6b. The four Pd-based electrical contacts were fabricated under optimized conditions by varying the thickness of the initial spin-coated film before the Ga⁺ irradiation. Two metallic Pd microstructures were horizontally placed facing each other and separated by only 40 nm. This gap value was chosen following the simulations on the ion trajectories shown in Figure 5b, which indicate ion lateral deviations within the ±20 nm range. The obtained resolution of 40 nm is comparable to other Ga-FIB-based processes, but it could be improved to sub-10 nm by using irradiation with focused He and Ne ions. In the latter case, the method would approach the best resolution achievable with more standard lithography techniques such as electron beam lithography. The structure shown in Figure 7,

Table 2. Growth Parameters for the Four Pd Electrical Contacts Here Studied Together With the Resistance and Resistivity Values for Each Pt Nanowire

| sample           | spin-coated initial thickness [nm] | Ga⁺ dose for Pd microstructures [μC/cm²] | resistance [kΩ] | resistivity [10⁳ μΩ·cm] |
|------------------|-----------------------------------|-----------------------------------------|----------------|------------------------|
| NW_Pt-FIBID_1    | 350                               | 30                                      | 235            | 9.2                    |
| NW_Pt-FIBID_2    | 200                               | 30                                      | 247            | 9.2                    |
| NW_Pt-FIBID_3    | 125                               | 30                                      | 147            | 5.4                    |
| NW_Pt-FIBID_4    | 125                               | 10                                      | 189            | 7.0                    |

“Value corresponding to the initial thickness of the spin-coated film before the Ga⁺ irradiation. b Value calculated considering that all nanowires have a width of 100 nm and a thickness of 30 nm.

Planar Nanogaps Using Pd Microstructures. The formation of planar nanogaps using Pd microstructures as electrodes has been undertaken. Two metallic Pd microstructures were horizontally placed facing each other and separated by only 40 nm. This gap value was chosen following the simulations on the ion trajectories shown in Figure 5b, which indicate ion lateral deviations within the ±20 nm range. The obtained resolution of 40 nm is comparable to other Ga-FIB-based processes, but it could be improved to sub-10 nm by using irradiation with focused He and Ne ions. The latter case, the method would approach the best resolution achievable with more standard lithography techniques such as electron beam lithography. The structure shown in Figure 7,

Figure 7. SEM micrograph of a planar nanogap device on a Si/SiO₂ substrate with the two microstructures separated by 40 nm and grown in 26 s of ion irradiation.

with a total area of 146 μm², was grown in only 26 s by means of an Ga⁺ beam current of 1.1 pA, which corresponds to an ion dose of 20 μC/cm². The initial spin-coating thickness was 215 nm, and the final thickness for this structure was 25 nm.

The high resolution achieved enables these structures to be used as nanogap electrodes for the fabrication of nanometer-size devices and circuits or to observe the electrical behavior of different nanomaterials. The main advantage of this fabrication method is the very low ion dose required to fabricate metal contacts and therefore the quick fabrication and
negligible ion-induced damage in the sample. Thus, this approach is a very good option to fabricate metallic nanopatterns with nanogaps that could be applied to place top or edge contacts on 2D semiconducting materials, neither inducing damage nor leaving resist residues as it occurs with resist-based fabrication methods. Furthermore, electrical nanogap devices for molecular electronics and biosensing have gained importance in recent years. A growing interest in inserting single molecules between electrodes exists due to the importance of understanding the transport behavior at the level of a single molecule to prepare entirely molecular integrated circuits.45,46 For biosensing, the biomolecules are trapped inside the gap between the electrodes and are detected by measuring their electrical behavior. Considering the range of our gaps (∼40 nm), these devices could, for example, be bridged to gold nanoparticles, which are used as labels for ultrasensitive electronic detection.47

Large-Area Mesh Fabrication and Electrical Characterization. The third application of this research consists in the growth of large-area metallic meshes, where the scale and shape of the patterns are changed without modification of the metallic nature of the Pd structures. To that end, square-based grids were fabricated on Si/SiO2 substrates covered by ∼270 nm-thick spin-coated films, bearing two wider pads at both sides of the grid in order to facilitate electrical measurements through the electrical microprobes, as shown in Figure 8a. Each mesh was composed of a total of 28 patterns with dimensions of 90 μm × 200 nm. The total irradiation time is 2 min, using an ion beam current of 1.1 pA, which corresponds to an ion dose of 30 μC/cm². The lateral electrical contacts, with dimensions of 75 μm × 10 μm, were fabricated using an ion beam current of 24 pA in only 30 s, which corresponds to an ion dose of 48 μC/cm². Considering the irradiation times, each electrical device (mesh and two electrical contacts) took only 2.5 min, resulting in a final thickness of (25 ± 5 nm). Figure 8b shows the I–V curves registered by the two-probe technique at room temperature. A linear Ohmic behavior is noticed in all devices fabricated under the same conditions. The electrical resistance values for these devices were in the 8–17 kΩ range.

With a view to expanding the scope of these metallic micro- and nano-structures toward their implementation in research areas in which optical transparency is required, we tested the fabrication of large-area meshes on indium tin oxide (ITO) substrates, obtaining promising results as shown in Figure S6.

OUTLOOK AND CONCLUSIONS

Micro- and nano-lithography techniques for large-area patterning commonly rely on sacrificial spin-coated resists that are removed at an intermediate step. There are very few examples where the spin-coated film is functional and is not removed during the lithography process. One of these rare examples is the case of 3D polymer-based scaffold structures for cell growth proliferation48 or polymers structured to serve as mechanical cantilever resonators,49 both fabricated by two-photon lithography. Another example is that of 3D cryogenic electron-beam writing for optical applications.50 The advantage of an electrically functional spin-coated film like Pd3(OAc)6 is the simplicity of the overall process, consisting in only three steps: spin coating the substrate with a Pd precursor film, focused Ga+ irradiation, and film development, as illustrated in Figure 1. This is a significant improvement with respect to previous strategies using organometallic films, where post-annealing steps were always needed to reach a metallic behavior. Remarkably, the patterning resolution achievable with the focused Ga+ beam irradiation is very high: gaps of 40

Figure 8. (a) SEM micrograph of a device formed by a large-area mesh and two electrical contacts fabricated in one step on a Si/SiO2 substrate. (b) I–V measurements of five devices indicating electrical resistances in the 8–17 kΩ range.
nm have been shown here, but the potential for better resolution through further optimization of the working parameters or by means of lighter focused ions (He, Ne, etc.) is very high.\textsuperscript{31} Moreover, the fact that the required ion dose to create metallic patterns (with electrical resistivity values of 70 $\mu\Omega\cdot$cm) is very low, which is 30 $\mu$C/cm$^2$, makes this approach very efficient with respect to other direct-write lithography techniques. As a comparison, the growth of metallic structures by focused electron and ion beam-induced deposition requires charge doses several orders of magnitude higher.\textsuperscript{38} This translates into a much shorter processing time for growing these Pd structures. For example, the Pd microstructure shown in Figure 7, with a total area of 146 $\mu$m$^2$, only took 26 s of irradiation compared to several hours of irradiation if grown by focused electron/ion beam-induced deposition. Cryogenic-focused ion beam-induced deposition is another high-throughput direct-write lithography technique that uses a similar ion dose range as the one used for these films, but it requires lowering the temperature of the stage to cryogenic temperatures.\textsuperscript{52}

The results shown here are the first of their kind and are thought to have important implications. First, they pave the way for their straightforward use in various applications where metallic micro- and nano-patterns are needed, such as interconnects, conductive micro- and nano-meshes, metallic micro- and nano-templates, nanogaps, electrical contacts to nano-objects, and so forth. Second, these Pd$_3$(OAc)$_6$ films can also be decomposed by electron irradiation\textsuperscript{34,36} or light irradiation,\textsuperscript{53} offering the capability of applying on the same film several micro- and nano-lithography techniques that span a wide range of lateral sizes. Third, there exist other acetates and organometallic films that contain elements such as Au, Co, Fe, and so forth, and their patterning by focused ion beam irradiation could lead to structures exhibiting additional functional properties beyond electrical conductivity, such as plasmonic or magnetic behavior. Moreover, Al- and Cu-based resists with comparable behavior to Pd$_3$(OAc)$_6$ could be relevant for the growth of CMOS-compatible metallic structures.\textsuperscript{54}

In summary, an efficient direct-write lithography method based on focused Ga$^+$ irradiation to create Pd-rich metallic patterns out of spin-coated Pd$_3$(OAc)$_6$ films which does not involve any post-annealing step has been described here. The method exhibits high resolution, as demonstrated by the creation of structures with 40 nm gaps. The patterned structures, which present a low electrical resistivity value of (70 ± 5) $\mu\Omega\cdot$cm, have been used to contact a nanowire and measure its electrical properties as well as for creating proof-of-concept large-area metallic meshes. Theoretical simulations have shed light on the ion-induced processes leading to the growth of Pd-rich nanostructures.

\section*{Supporting Information}

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.2c05218.

Additional characterization details, AFM characterization, $I$–$V$ curves, thickness measurements, XPS studies, other SEM micrographs, and additional information about Monte Carlo simulations (PDF)
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