Area-Selective Atomic Layer Deposition of In$_2$O$_3$:H Using a $\mu$-Plasma Printer for Local Area Activation

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Supporting Information

Electronic device fabrication generally requires several lithographic steps to obtain well-aligned features. This is becoming ever more challenging in the semiconductor industry as feature dimensions of state-of-the-art devices keep shrinking.1−4 It is, however, not only the semiconductor industry that faces such challenges. Some of the challenges in patterning for nanoelectronics share common ground with those of microscale patterning as used in large-area electronics. Furthermore, removing material by etching can be detrimental in terms of materials damage and contamination.5 Moreover, for large-area electronics cost per unit and throughput considerations also limit the use of expensive and iterative “litho-etch” steps. This holds, for example, for patterned transparent conductive oxides (TCOs).6−9 TCO films are ubiquitous in many devices such as displays,10−12 solar cells,13 and solid state lighting and sensors,14 including those based on emerging organic light-emitting devices (OLEDs)15 and on halide perovskite materials. To date, methods used for patterning TCOs such as such as rapid laser patterning16 often have limited uniformity and resolution or suffer from material degradation during the etch processes. Furthermore, more than six steps are typically required to fabricate patterned TCOs.17−19 Hence, several applications that require millimeter and submillimeter patterned TCO films would benefit from the availability of alternative patterning methods.20−12

Area-selective atomic layer deposition (AS-ALD) has emerged as a promising solution to push device manufacturing toward new frontiers. Significant efforts, in academia and industry, are currently devoted toward the development of AS-ALD techniques that can be incorporated within existing fabrication schemes. To date, most of these efforts have been inspired by the need to prepare nanoscale features in the semiconductor industry as an alternative or a possible solution for extremely complex lithography schemes. Although the number of viable methods is still very limited, the approaches for achieving AS-ALD can be categorized as (1) area deactivation, in which a part of the surface is rendered inert toward a specific ALD process, and (2) area activation, where conversely, an inert surface is locally activated for a specific type of ALD chemistry.21−23 A prototype example of the first case is area deactivation of a surface by blocking it with self-assembled monolayers.21,22 A particular case of the latter approach is the direct activation of a surface region without any subtractive steps. This can be referred to as direct-write ALD21,23 and it enables patterning and ALD to be combined into a truly bottom-up process drastically reducing the number of required fabrication steps. One embodiment of direct-write ALD was developed by Mackus et al. by exploiting the catalytic activity of a Pt seed layer grown locally using electron beam induced deposition (EBID). After EBID, ALD was applied for selectively thickening the Pt seed layers into high-quality Pt nanostructures.24,25

In this work, we present a novel method for AS-ALD which targets the preparation of microscale features relevant in large-area electronics. It is a direct-write ALD process of In$_2$O$_3$:H, a highly promising and relevant TCO material,26 which makes use of printing technology for surface activation. As schematically depicted in Figure 1, first the surface of H-terminated silicon materials is locally activated by a $\mu$-plasma printer in air or O$_2$, and subsequently In$_2$O$_3$:H is deposited selectively on the activated areas. The selectivity stems from the fact that ALD In$_2$O$_3$:H leads to very long nucleation delays on H-terminated silicon materials.27 In this work it is demonstrated that this method allows for preparing microscale In$_2$O$_3$:H features in a true bottom-up approach on surfaces of H-terminated Si(100), a-Si:H or a-SiN$_x$:H with the In$_2$O$_3$:H material quality being as high as for blanket films.28 This novel approach for direct-write ALD of In$_2$O$_3$:H holds therefore potential for applications in large-area electronics which make use of patterned TCOs such as displays and solar cells.

For the experiments an Oxford Instruments Opal ALD reactor was used to deposit the In$_2$O$_3$:H films. Indium cyclopentadienyl (InCp) served as indium precursor and a mixture of H$_2$O and O$_2$ as coreactants.27 A standard recipe was employed as detailed elsewhere.22,27 The reactor pressure was ∼150 mTorr throughout the cycle with a pressure spike of ∼200 mTorr during the 20 ms H$_2$O dose. All samples prepared in this study were deposited at 100 °C.

Films of a-Si:H and a-SiN$_x$:H on p-type Si(100) served as substrates. These 10 nm thick films were prepared by inductively coupled plasma chemical vapor deposition (ICP-CVD) from SiH$_4$ and Ar at 50 °C and from SiH$_4$ and NH$_3$ at 80 °C for a-Si:H and a-SiN$_x$:H, respectively. Alternatively, H-terminated Si(100) was prepared by dipping the wafer in diluted (1 vol %) hydrofluoric acid (HF) for 2−3 min. Additionally, a 400 μm thick stainless steel foil with 20 nm of a-Si:H on top was used as a flexible substrate.

After preparation, the surfaces were locally activated by a $\mu$-plasma printer developed by the company InnoPhysics. The
print-head of this μ-plasma printer consists of actuated needle electrodes in a multi needle-to-plate dielectric barrier discharge configuration. It allows varying the patterning width by setting the number of actuated needles and achieving high throughput (1 cm² in less than 5 s). For this study air and O₂ plasmas at atmospheric pressure were used to locally activate the surface of the silicon based materials.

By carrying out the sequence of local plasma activation and the ALD cycles as illustrated in Figure 1, the area-selective deposition was demonstrated for a macroscopic pattern on a large area. Figure 2a shows a 4-in. wafer covered with 10 nm a-Si:H onto which a patterned film was prepared by μ-plasma printing and 400 ALD cycles. This resulted in ∼35 nm thick In₂O₃:H locally deposited as measured by ex situ spectroscopic ellipsometry (SE) at two different angles (65° and 75°, using an M-2000D system from J. Woollam Co. with photon energy range of 1.2 to 6.5 eV). The selectivity of the process was confirmed by XPS measurements using a Thermo Scientific K-Alpha system probing distinctive points inside and outside the patterned area. The In 3d₃/₂ and In 3d₅/₂ signals for two of such points are shown in Figure 2b. High signals were collected on the patterned area whereas a negligible signal was measured outside the pattern. This demonstrates the excellent selectivity reached, even after 400 ALD cycles. This finding was substantiated by I–V measurements within and between the different letters of Figure 2a (see Supporting Information, Figure S3).

Figure 1. Schematic representation of the area-selective ALD process of In₂O₃:H on H-terminated silicon materials. In the first step (1), microscale patterns are defined by activating the surface with a μ-plasma operated in air or O₂. In the second step (2), the In₂O₃:H is deposited selectively on the activated areas in a building step. The ALD process consists of two alternating half reactions: InCp precursor dosing in pulse A and a mixture of O₂ and H₂O dosing in pulse B. Note that in the case that conductive substrates are used, a thin dielectric membrane (Al₂O₃) is positioned between the needles and the substrate, as shown in Figure S1a.

Figure 2. (a) Photograph of a 4-in. Si(100) wafer covered with 10 nm of a-Si:H with the letters “TU/e” prepared using the direct-write ALD process of In₂O₃:H. The number of ALD cycles was 400 and the thickness of the In₂O₃:H was ∼35 nm. (b) XPS signals for the In 3d₃/₂ and In 3d₅/₂ binding energy measured for two distinctive points inside (black) and outside (red) the patterned area.

As mentioned at the beginning of this communication, the selectivity of the In₂O₃:H ALD process is stemming from the fact that very long nucleation delays are obtained on H-terminated...
surfaces of silicon materials whereas the nucleation delay on oxide surfaces is considerably shorter as shown in Figure 4.25 To elucidate the role of the surface groups during ALD of In$_2$O$_3$:H, first-principle calculations have been carried out using density functional theory (DFT) with the Perdew–Burke–Ernzerhof (PBE) exchange correlation functional. van der Waals interactions were also accounted for on an empirical basis. Additional computational details can be found in the Supporting Information Section 1.4. The structures and associated relative energies have been calculated for the stationary points along the anticipated reaction pathways for the chemisorption of the InCp precursor and proton transfer from the substrate to the Cp ligand of the precursor. This was done for (1) hydrogenated silicon and (2) hydroxylated silicon oxide surfaces, as these are expected to represent the H-terminated and plasma activated surfaces to a good approximation. The reactions considered are therefore

$$\text{Si–H}^* + \text{InCp} \rightarrow \text{Si–In}^* + \text{HCp} \quad (1)$$

$$\text{Si–OH}^* + \text{InCp} \rightarrow \text{Si–O–In}^* + \text{HCp} \quad (2)$$

where the asterisks indicate the surface species involved in the reactions.

Figure 5 shows the corresponding energies obtained by the DFT calculations. The results reveal that Reaction 1 is endothermic and requires an energy input of 0.31 eV to proceed, with associated activation energy of 0.36 eV. For Reaction 2, the activation energy of 0.28 eV is slightly lower, and in contrast to Reaction 1, the overall process is exothermic.

Based on the Arrhenius equation, assuming the prefactor is similar in magnitude for both surface reactions, this difference in activation energies translates into an ~13-fold higher rate for Reaction 2 in comparison to Reaction 1 at the ALD temperature of 100 °C. Moreover, the proton transfer reaction for Reaction 2 is exothermic and releases energy (0.30 eV). By contrast, Reaction 1 requires an energy intake of 0.31 eV to take place, such that Reaction 2 likely proceeds with a much higher probability.

The predicted reaction energies therefore point to the thermodynamically favored nature of InCp binding on the hydroxylated SiO$_2$ surface as opposed to the H-terminated silicon. This can explain the high selectivity that can be reached.

Figure 3. (a) Photograph of In$_2$O$_3$:H lines being 3.0 and 0.8 mm wide as prepared by the direct-write ALD process with 400 cycles. XPS line scans for the patterns depicted in (a) showing the atomic percentages related to (b) In$_2$O$_3$ (In 3d$_{3/2}$ and O 1s) and (c) the Si substrate (Si 2p).

Figure 4. Film thickness measured by in situ SE as a function of the In$_2$O$_3$:H ALD cycles on SiO$_2$ (open triangles) and on a-Si:H (open circles). Ex situ SE measurements were taken only for 600 and 780 ALD cycles on a-Si:H. Above 600 ALD cycles the selectivity of the process appears to degrade.

Figure 5. Energy profiles computed by DFT method (PBE-D3) for the chemisorption of InCp on (a) hydrogenated silicon (Si–H termination) and (b) on hydroxylated silicon oxide (Si–OH termination).
for the area-selective ALD process of In$_2$O$_3$:H on activated and nonactivated surfaces of H-terminated silicon materials.

In conclusion, a novel direct-write ALD process has been demonstrated for In$_2$O$_3$:H, a highly relevant TCO material. By combining local activation of H-terminated surfaces of silicon materials by a μ-plasma printer and AS-ALD of In$_2$O$_3$:H, microscale patterns can be generated in a true bottom-up process. Excellent selectivity and low resistivity of the In$_2$O$_3$:H were demonstrated. The method therefore provides bright prospects for large-area applications because both the μ-plasma printing and the AS-ALD process are scalable. Large-area applications employing flexible substrates such as OLED displays and flexible photovoltaics are of particular interest due to the current trend toward printing technology and roll-to-roll ALD processes. Therefore, as a first demonstrator, the direct-write ALD process was employed on a flexible stainless steel foil with 20 nm thick a-Si:H as is shown in Figure 6. Transferring the technology to printing on flexible substrates combined with spatial ALD processes will be the next steps. Also the extension of the process to other substrate materials and ALD materials systems will be considered. Other ALD precursors that undergo similar surface reactions can possibly also be used to develop area-selective ALD in a few cases. Finally, we note that preliminary investigations using a focused electron beam have yielded promising results with respect to the extension of the current direct-write ALD process of In$_2$O$_3$:H toward nanoscale dimensions.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemmater.6b04469.

μ-plasma printer details, XPS line scans and I−V measurements, SEM images, and computational methods (PDF)

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