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Comparing Lead Iodide and Lead Acetate Based Perovskite Absorber Layers by Aerosol-Assisted Chemical Vapor Deposition

Mohammad Afzaal,¹ Shreya Basak,² and Heather M. Yates³
¹Maths and Natural Sciences Division, Higher Colleges of Technology, P.O. Box: 7947, Sharjah, United Arab Emirates. Email: mafzaal@hct.ac.ae
²Department of Physical Sciences, Indian Institute of Science Education and Research Kolkata, Mohanpur, 741246, West Bengal, India
³Materials and Physics Research Centre, University of Salford, Salford, M5 4WT, United Kingdom

Abstract—Aerosol-assisted chemical vapor deposition at atmospheric conditions has been used to deposit optically active lead perovskite thin films. Lead iodide and lead acetate trihydrate precursors were used and the resulting films compared. X-ray powder diffraction measurements confirmed the room temperature stable tetragonal phase which converted to monohydrate or dihydrate phase of perovskite. The stability of the differently derived films were studied by X-ray photoelectron spectroscopy by a detailed probing of surface chemistry and composition of deposited materials before and after argon etching.

Keywords—Absorber layers; perovskite; chemical vapor deposition; Photoluminescence

I. INTRODUCTION

The last few years have seen an unprecedented interest in lead-based perovskite (PK) solar cells with stable efficiencies in excess of 24% [1]. Important characteristics, for example, high absorption coefficient, superior charge transport properties, ability to tune optical bandgap and long electron-hole diffusion lengths in PK layers are fundamental to improved photoconversion efficiencies in PK solar cells [2].

Before considering the technology as a serious contender to conventional silicon solar cells, low-cost processes capable of fabricating cell components on a large scale need exploring. Keeping this in mind, chemical vapor deposition (CVD) processes can be truly regarded as a commercially viable option as it is a choice of coating technology in, for example, the glass industry [3]. Many adaptations of CVD technologies are available which are helpful in circumventing issues, such as, the need for high vapor pressure precursors, deposition at reduced pressures, high operational costs and complex experiment design. Aerosol-assisted (AA) CVD process carried out at atmospheric pressure is highly attractive as long as precursor(s) are soluble in high boiling point solvents. This is necessary for generating an aerosol mist from a precursor solution, which is then transported to the heated substrate by a carrier gas. At the surface, the solvent evaporates resulting in the nucleation followed by thin film growth [4].

Several groups including ours have used AACVD to synthesize lead-based perovskite thin films. Herein, an attempt is made to systematically address the stabilities of lead iodide and lead acetate based perovskite thin films stored under different conditions. This is important from the point of view of device performance as any moisture along with air exposure has profound implications on the optical and morphological characteristics of perovskite thin films.

EXPERIMENTAL WORK

Precursor Solutions

Solutions of methylammonium lead iodide perovskite (CH₃NH₃PbI₃) were prepared under nitrogen (N₂) by stirring a dimethylformamide (DMF) solution of lead acetate trihydrate (Pb(Ac)₃.3H₂O) or lead (II) iodide (PbI₂) with methylammonium iodide (MAI). In a typical experiment, 12 mL of DMF was added to 1.466 g of MAI in a two-neck round bottom flask. After stirring for 1 hour, 1g of Pb(Ac)₃.3H₂O or 1.414g of PbI₂ was added and stirred for 15 mins before being stored under N₂. In all experiments, the molar ratio of MAI and PbI was fixed to 3:1 while that of MAI and Pb(Ac)₃.3H₂O was kept at 3.5:1 to compensate for the hydrated part.

A. Deposition of thin films

Before experiment, 1.1 mm borosilicate glass substrates (2.5 cm in width) were cleaned with detergent, water and propan-2-ol followed by air drying. The clean substrates were placed at the center (hot zone) of the AACVD chamber and heated to 140 (± 2) °C for 30 mins to remove any native oxide from the glass surface. After placing 12 ml of mother solution, the deposition was carried out at atmospheric pressure under a flow of nitrogen carrier gas (0.5 L/min) for 80 mins. To prevent any oxidation of deposited films, samples were cooled at room temperature under nitrogen before removing from the chamber. For a comparative study, deposited films were stored in a nitrogen-filled glove box (<1 ppm O₂, <2 ppm moisture), open-air (21°C and RH 75 %) and humid chamber (21°C and RH 90 %). The relative humidity (RH) readings were carried out using a handheld thermo-hygrometer.
B. Characterization of thin films
A Siemen D5000 instrument was used for X-ray powder diffraction (XRPD) measurements. The surface images were recorded on a Philips XL-30 FEG SEM. Photoluminescence measurements were performed with an excitation wavelength of 405 nm and an experimental set-up as described previously. X-ray photoelectron spectroscopy (XPS) was used to determine the surface composition of the samples using a Thermo Fisher Scientific K-alpha+ spectrometer. This used a micro-focused monochromatic Al X-ray source (72 W), along with a combination of low energy electrons and argon ions for charge neutralization.

II. RESULTS AND DISCUSSION

Figure 1: XRD patterns of CH₃NH₃PbI₃ thin films stored under different conditions.

Figure 2: SEM images of CH₃NH₃PbI₃ thin films (a) PbI₂-as deposited, (b), Pb(Ac)₂·3H₂O-as deposited, (c) PbI₂-air exposed, (d) Pb(Ac)₂·3H₂O-air exposed, and (e) PbI₂-humidity chamber.

Lead iodide and lead acetate based perovskite absorber layers prepared by AACVD experiments were exposed to different conditions for their structural and optical sensitivities. Films are grey and could be easily scratched with a sharp object. Figure 1 shows the changes in XRD patterns as the films are exposed to different environments. Initial XRD measurements on as-deposited perovskite films, for example, show a typical room stable tetragonal phase. As the films were exposed to different humidity levels either at room temperature or in a humidity chamber, a strong peak emerged at 10.6°, which is characteristic of monohydrate (CH₃NH₃·H₂O)PbI₃ [8]. It is also notable that the peak at 14.16°, responsible for methylammonium lead iodide (MAPI), observed for as-deposited films vanished after exposure and was replaced with more intense peaks at higher 2θ values. These peaks point to the formation of polycrystalline (CH₃NH₃)₂PbI₃·2H₂O within the perovskite films [8]. It is highly likely that the hydration of MAPI films occurs in two-steps. Initially, the crystal structure of MAPI is saturated with one water molecule per formula unit. With high humidity levels, a new structure containing two water molecules per formula is followed. A similar
observation is report earlier where authors have studied the hydration of MAPI structures [8].

Upon air exposure, the MAPI films turned from grey/black to yellow within 24 hours. Within a humidity chamber, films became patchy and translucent pale yellow within a few hours. The SEM images given in Figure 2 show how the morphologies of MAPI change. In Figure 2a, as-deposited films from PbI$_2$ are globular with almost full coverage. The films based on Pb(Ac)$_2$.3H$_2$O are somewhat denser and closely packed with evidence of some pinholes (Figure 2b). Upon storage under at room conditions, most of the film coverage is lost as a glass substrate can be clearly seen in the images.

Attempts to acquire reliable optical transmission and reflection data proved unfruitful, most likely a result of scattering effects from the rough surfaces. After exciting films with a blue laser (405 nm), PL response for as-deposited films was found at $\approx$ 776 and 784 nm for PbI$_2$ and Pb(Ac)$_2$.3H$_2$O, respectively. Perhaps more importantly, PL spectra of MAPI based on Pb(Ac)$_2$.3H$_2$O was much narrower (47.97 vs. 63.32 nm) than that of PbI$_2$ suggesting that the acetate related perovskite has a lower defect density. It is worth mentioning that second-order peak of the laser was removed during the handling of data. The air exposure resulted in a blue shift of 25.6 and 13.2 nm in perovskites formed from PbI$_2$ and Pb(Ac)$_2$.3H$_2$O, respectively due to the gradual formation of PbI$_2$. The air-exposed and humidity exposed PL peaks of films showed negligible difference in wavelength of PL maxima as well as FWHM. Considering the narrow PL responses, it can be suggested that Pb(Ac)$_2$.3H$_2$O is a better precursor than PbI$_2$ for CVD of perovskite thin films. The lower shift in PL overtime for the lead acetate derived perovskite suggests this may be slightly more stable.

The surface chemistry of deposited PK films was analysed by XPS. The survey scan confirmed the presence of I, N, Pb and C (Figure 4). High-resolution scans showed that the C1s consisted of two signals. The more intense signal at 286.7 eV associated with the carbon within the methylammonium cation [9] and a smaller one due to amorphous C (285.3 eV). Although not seen on the survey scan (apart from 2 specific samples which will be discussed later), a high-resolution O1s scan showed a very weak peak. The intensity of I and Pb increased after a 10 s Ar etch while the C and O decreased confirming that the latter related to environmental surface contaminates. Figure 4 shows the survey scan for a representative sample both before and after the Ar etch. The concentration of oxygen related species are too low to be seen on the survey scan so the related high resolution scans for O1s are also shown.

The high resolution I 3d scan displayed only a single chemical state for the 3d$_{5/2}$ peak at 619.6 eV with a characteristic splitting of 11.5 eV [10]. The position of the N 1s at 402.7 eV is similar to that given by Zou et al. [11] and ascribed to N within the methylammonium ion. In all samples, the intense Pb 4f doublet (4f$_{7/2}$ at 137.6 eV and splitting of 4.88 eV) is in assigned to Pb within PK. Figure 5 shows high-resolution scans for both MAPI derived PbI$_2$ and Pb(Ac)$_2$.3H$_2$O. A few samples establish the presence of a lower intensity signal with 4f$_{7/2}$ at 137.1 eV related to metallic Pb. This is especially noticeable after the Ar etch. Metallic Pb within perovskite samples has been previously reported and has been attributed to decomposition during annealing [12], laser irradiation [13] or beam damage during measurements [10].

As the intensity of the peak noticeably increased after Ar etching the majority of the Pb presence is ascribed to beam damage. This was especially obvious in the lead acetate derived samples where the Pb$^+$ content approximately doubled on etching with, for example, rising from 6% to 13% of the total Pb$^+$ content in the samples stored within a dry glove box. See figure 5 for a comparison of the effect of etching on the differently derived MAPI samples. At the same time, this is mirrored with a reduction in the relative amount of N, which may indicate some PK decomposition. However, there is no obvious change in the very small high-resolution O 1s
signal, so establishing that no significant absorption of moisture took place, unlike that reported by Wang and co-workers [9].

Figure 5: High-resolution XPS scans of Pb 4f. (a) After argon etch and (b) as-deposited from Pb(Ac)2·3H2O. (c) After argon etch (d) as-deposited from PbI2.

A range of PK samples were analyzed (Figure 6) and the effects of initial precursor and storage conditions considered. Prior to calculation of PK stoichiometry, both the metallic Pb and amorphous C components were removed. Table 1 shows that the ratio of Pb:I:N:C is not that of a perfect PK samples (1:3:1:1). In particularly the N and especially C values obtained are not accurate due to interference from the Pb 4d signal for N1s, the low intensity of the peaks and poor signal/noise level for particularly the C 1s. Many papers report almost perfect stoichiometry for the perovskite as derived from XPS measurements. However, most of these have deposited the PK films in a vacuum with direct transfer to the XPS instrument [9, 11, 13], controlled low humidity (<30%) [14]. Our samples were prepared in an with limited control over the timescale between deposition and analysis. The latter was minimised by transporting the samples in sealed, evacuated bags. However, as all samples were treated similarly conclusions can still be reached from relative comparisons between the differently treated samples.

As the Ar etch led to damage of the films, the discussion will concentrate on the initial surface scan as more representative of the samples. Use of lead acetate rather than lead iodide lead to samples with similar stoichiometry, close to that of the perovskite. These samples stored in a glove box (with controlled low humidity and low light levels) showed limited decomposition. In comparison, the samples stored in the open lab environment with high humidity and room light showed a reduction in the amount of I to Pb due to the start of decomposition to PbI2 and the discharge of the HI and NH3 into the air [9]. The amount of decomposition is still low, as no sign of any yellow PbI2 and the samples retain their black coloration.

<table>
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Table 1. XPS analysis of perovskite thin films. *Third small peak at 288.7 eV, removed on etching, so probably surface contamination.

Figure 6. High-resolution scans of as-deposited film constituents resulting from Pb(Ac)2·3H2O (top) and PbI2 (bottom).
peaks for Pb(Ac)₂·3H₂O rather than PbI₂ derived films. A thorough surface analysis demonstrated different stoichiometries, stability, morphology and level of especially carbon contaminants.

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**Figure 7.** High-resolution spectra O 1s. (a) Pb(Ac)₂·3H₂O stored in open lab, (b) PbI₂ in humidity chamber, (c) Pb(Ac)₂·3H₂O in glovebox, (d) PbI₂ in open lab, and (e) PbI₂ in glovebox.
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