Inverted Perovskite Photovoltaics Using Flame Spray Pyrolysis Solution Based CuAlO2/Cu−O Hole-Selective Contact

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Supporting Information

ABSTRACT: We present the functionalization process of a conductive and transparent CuAlO2/Cu−O hole-transporting layer (HTL). The CuAlO2/Cu−O powders were developed by flame spray pyrolysis and their stabilized dispersions were treated by sonication and centrifugation methods. We show that when the supernatant part of the treated CuAlO2/Cu−O dispersions is used for the development of CuAlO2/Cu−O HTLs the corresponding inverted perovskite-based solar cells show improved functionality and power conversion efficiency of up to 16.3% with negligible hysteresis effect.

KEYWORDS: CuAlO2/Cu−O alloying, delafossite oxides, hole-transporting layers, perovskites solar cells, flame spray pyrolysis

1. INTRODUCTION

The discovery of novel solution processed light-harvesting materials boosted the photovoltaic research toward innovative concepts, such as hybrid perovskite-based solar cells.1 Within less than 10 years of development, perovskite-based solar cells exhibit lab-scale power conversion efficiencies (PCEs) of over 22%.2−4 In addition, the substantial progress on novel perovskite solar cell architectures based on solution processing, marks the compatibility of these devices with large-scale printing production.5 Among many types of perovskite solar cell architectures,6 the planar heterojunction p−i−n architecture shows some relevant advantages to large scale production. For example, the fullerene processed on top of the perovskite as the n-contact has been found to eliminate the current−voltage (J−V) hysteresis, which is generally observed in the n−i−p device architecture.7 Furthermore, both the fullerene n-type contact and the p-type contact can be fabricated at low processing temperatures.8,9,10

Charge selective contacts are of high importance for the functionality of p−i−n perovskite solar cells.11,12 Poly(3,4-ethylenedioxythiophene)-poly(styrenesulfonate) (PEDOT:PSS) is the most commonly used p-type material acting as hole transporting layer (HTL) in p−i−n perovskite solar cells. However, despite its excellent optoelectronic properties, most of the PEDOT:PSS-based CH3NH3PbI3 solar cells were reported to have an inferior open-circuit voltage (VOC) (0.88−0.97 V)13 compared to that obtained by n−i−p CH3NH3PbI3-based solar cells (1.05−1.12 V).14 Therefore, it is critical to pursue materials and interface systems with minimal voltage losses for developing high-performance inverted perovskite solar cell devices.

As alternatives to PEDOT:PSS, other polymeric p-type materials, have been identified as suitable HTLs for high performance p−i−n perovskite solar cells15 and in most cases lead to devices with higher VOC values. However, the excessive cost of these materials, because of their synthetic procedure and high-purity requirement, is immensely hampering the future commercialization of perovskite solar cells. Low-cost, metal oxides have been identified as promising alternative HTLs in inverted perovskite solar cells.16,17 Solution processed CuO18,19 NiO19 and CuSCN20 have been reported as efficient...
HTLs providing high $V_{\infty}$ values (>1 V) for p–i–n perovskite solar cells. Despite their excellent progress, all the hole transporting layers usually exhibit low electrical conduction properties. By using metal oxide-based HTLs with relatively high electrical conductivity, thicker layers might be used without significant losses in device internal resistances.\textsuperscript{21} This is a widely known issue in organic photovoltaic (OPV) technology, where the use of thicker charge selective layers are essential for a reliable layer formation using roll-to-roll (R2R) large scale processing, as well as device reproducibility.\textsuperscript{22} Similar to OPV technology, the use of conductive and thicker charge selective layers results in a series of benefits concerning product development targets of printed perovskite solar cells.\textsuperscript{23} In contrast to solution processed n-type metal oxide charge selective layers, the realization of solution processed conductive and transparent p-type metal oxide thin films to be used as HTLs in inverted perovskite solar cells is a challenge. To this end, only few solution processed metal oxide materials which combine high transparency and electrical conductivity have been used as HTLs in p–i–n perovskite solar cells. Recently, Jung et al. showed a solution processed CuNiO$_x$\textsuperscript{24} as an efficient HTL for efficient p–i–n perovskite solar cells. Furthermore, Chen et al. demonstrated that the use of a conductive Ni$_x$Mg$_{1-x}$O HTL leads in an efficient extraction of positive carriers, and the corresponding perovskite-based p–i–n solar cells result in excellent fill factor (FF) values exceeding 80% and hysteresis-free behavior.\textsuperscript{25} Our search for highly conductive and transparent p-type oxide semiconductors for potential use as HTLs in perovskite solar cells, led us to the delafossite compounds, one of the few known categories of p-type materials that combine relatively high electrical conductivity and high transparency in the visible spectrum. Delafossites have the general formula of A$^{+1}$X$^{3+}$O$_2$, where A is mostly Cu (or Ag), X is often Al, Ga, Cr, and O is oxygen. Delafossites can be crystallized to either a rhombohedral 3R-(R3m) or hexagonal 2H-(P63/mmc) structure.\textsuperscript{26} Kawazoe et al. were the first to report that CuAlO$_2$ produced by laser ablation exhibits room temperature electrical conductivity of 1 S/cm,\textsuperscript{27} one of the highest reported room temperature electrical conductivity values of p-type metal oxide semiconductors. In addition, several studies show energy levels compatible with p-type electrodes of electronic devices, such as large optical band gap\textsuperscript{28} and deep work function in the range of 5.5–6.5 eV.\textsuperscript{29} However, despite their excellent potential, a major drawback for a wider use of these materials is the extreme amount of energy required to obtain pure delafossite functional films. For example, sol–gel processes can be used to fabricate delafossite thin films only by annealing the gel-films at temperatures over 1000 °C under inert atmospheres,\textsuperscript{30} which is unlikely to be used for the fabrication of solution based perovskite solar cells. Delafossite nanopowders can also be fabricated using hydrothermal methods or microwave assisted hydrothermal methods under extreme pressures (>300 MPa).\textsuperscript{31} Wang et al. demonstrated that solution-processed CuGaO$_2$ nanoplates is a promising HTL at the top electrode of inverted organic solar cells.\textsuperscript{32} However, the large size of CuGaO$_2$ nanoplates result in rough films and expected to affect device reproducibility. To overcome this issue, the same authors demonstrated a suspension of sub-10-nm CuCrO$_2$ NPs, which allows a smooth and transparent film to be fabricated from the dispersion and be used as an efficient HTL for P3HT:PCBM-based organic solar cells.\textsuperscript{33} Despite the impressive development on this promising category of p-type materials, only recently studies show a successful incorporation of delafossite materials as HTL in perovskite solar cells. Zhang et al. demonstrated an efficient CuGaO$_2$ nanoplate dispersion fabrication by microwave assisted hydrothermal method as a HTL at the top electrode of n–i–p CH$_3$NH$_3$PbI$_3$-based solar cells.\textsuperscript{34} In addition, Dunlap-Shohl et al. showed an efficient CuCrO$_2$ HTL in p–i–n perovskite solar cells.\textsuperscript{35} We have recently shown that ultrafine delafossite CuGaO$_2$ nanoparticles can be synthesized using a surfactant assisted hydrothermal reaction and be used efficiently as HTLs in p–i–n perovskite solar cells. Specifically, we have presented for the first time that the addition of Pluronic P123 surfactant is beneficial for the synthesis of CuGaO$_2$ NPs, as it can decrease the surface energy and temperature of crystallization and control NPs size (~5 nm).\textsuperscript{36} DMSO was subsequently used as a ligand and dispersing solvent for stabilizing the CuGaO$_2$ which were successfully used as HTL in p–i–n perovskite solar cells. The corresponding 15 nm CuGaO$_2$–HTL-based inverted perovskite solar cells show PCE of 15.3%.\textsuperscript{36} CuAlO$_2$ is one of the most promising delafossite materials for optoelectronic applications for several reasons. For example, it exhibits superior optical transparency in the visible spectrum compared to CuCrO$_2$ and at the same time maintains high electrical conductivity.\textsuperscript{37} In addition, it is comprised by earth abundant components (Cu and Al) promoting it in a better choice related with large scale use compared with the expensive CuGaO$_2$. To our knowledge there is no reported study on high performance crystalline delafossite CuAlO$_2$ HTLs for solar cell applications. One reason for this, is the complex aluminum chemistry that is prohibiting the efficient hydrothermal synthesis of CuAlO$_2$ nanopowders.\textsuperscript{38} Recently Igbari et al. used d.c. magnetron sputtering technique “amorphous” type CuAlO$_2$ (a: CuAlO$_2$) interfacial layer to modify the interface of ITO and PEDOT:PSS in planar perovskite solar cells.\textsuperscript{39} Experimental results of crystalline CuAlO$_2$ are limited because of the difficulty to synthesize pure phase delafossite CuAlO$_2$ thin films.\textsuperscript{31} It has been shown that p-type defects play a key role on the electrical properties of CuAlO$_2$\textsuperscript{40} and that CuAlO$_2$ thin films can be engineered by nonisovalent Cu–O alloying.\textsuperscript{36} Here, we report for the first time the application of a solution processed CuAlO$_2$/Cu–O HTL for p–i–n perovskite solar cells. The XPS peak analysis reported within section 2.1 estimated that the percentage of CuAlO$_2$ is 77% and that of Cu–O is 23% (within a ± 5% standard error). The CuAlO$_2$/Cu–O powder was developed by high temperature flame spray pyrolysis and the stabilized CuAlO$_2$/Cu–O dispersion sonicated and centrifuged as described in detail below and within the experimental section. Using the supernatant part of the treated CuAlO$_2$/Cu–O dispersions the ultralarge CuAlO$_2$/Cu–O agglomerates are avoided. The proposed treatment is essential to avoid shorting of the devices and to provide functional small area solution processed CuAlO$_2$/Cu–O films. A detail study of material structure, agglomerates/particle sizes, dispersion treatment process, properties, processing, and device parameters is presented. The photoluminescence spectroscopy measurements on perovskite films show a reduced electron–hole pair recombination for the treated CuAlO$_2$/Cu–O HTL device structure and the electroimpedance spectroscopy studies confirm their efficient hole extraction properties. The experimental results show the potential of hole extraction.
carrier selectivity and surface properties of the treated CuAlO₂/Cu−O HTL providing negligible hysteresis inverted perovskite PVs with best achieved PCE of 16.3%.

2. RESULTS AND DISCUSSION
The development of the proposed CuAlO₂/Cu−O flame spray pyrolysis solution-based HTL includes the following process: For flame spray pyrolysis, a metal organic precursor was prepared by a mixture of organic copper and aluminum salt dissolved in a 1:1 mixture of alkyl carboxylic acid and toluene. The metal organic precursors were filtered through a 0.45 mm PTFE syringe filter and fed (5 mL/min) to a spray nozzle, dispersed by oxygen (7 L/min), and ignited by a premixed methane−oxygen flame (CH₄, 1.2 L/min; O₂, 2.2 L/min). The off-gas was filtered through a glass fiber filter by a vacuum pump at about 20 m³/h. The obtained nanopowder was collected from the glass fiber filter. The flame spray pyrolysis powders undergo a high temperature (1100 °C for 30 min in air) post-treatment sintering process. However, the high temperature sintering step leads to agglomeration and to the formation of large agglomerates (rods/plates) which are shown in supplementary, Figure S1. The powders are stabilized in a 10 wt % 2-propoxyethanol (2-PPE) dispersion using an Avantama, Ltd., surfactant and the stabilized dispersions sonicated for over 2 h (with high frequency sonicator probe) following by centrifugation (at 3000 rpm) for 4 min. The supernatant of the treated CuAlO₂/Cu−O dispersion was collected and used for the processing of the functional HTLs. By application of high-energy sonication to the dispersions, although large particles are not eliminated other large agglomerates significantly reduced. During the centrifugation process the larger/heavier agglomerates remain at the bottom of the solution. Thus, by making devices from the top of the CuAlO₂/Cu−O dispersion (supernatant dispersion) although broad distribution of particle sizes is still existing large agglomerates are in most cases avoided (we discuss this issue in detail later within the text). To form the proposed CuAlO₂/Cu−O HTL, the supernatant part of the treated (sonicated and centrifuged) dispersions were dynamically spin coated at 3000 rpm for 60 s on the preheated ITO substrates followed by annealing on a hot plate at 400 °C for 20 min in ambient conditions. Thus, the following terminology is used within the text: Solution A (room temperature), named SolA@RT, is the CuAlO₂/Cu−O spray pyrolysis powders stabilized in a 10 wt % 2-propoxyethanol using an Avantama, Ltd., surfactant, solution B (room temperature), named SolB@RT, is solution A solution treated (sonicated and centrifuged) at room temperature (RT) and SolB@400°C is solution A treated (sonicated and centrifuged) and annealed at 400 °C for 20 min. The terms films A and B indicates that the measurements performing in films using the corresponding CuAlO₂/Cu−O solutions described above. The terms FilmA (@RT and @ 400 °C) and FilmB (@RT and @400 °C) shows the annealing conditions of the CuAlO₂/Cu−O films. We note that (FilmB@400°C) represents the identical processing conditions.
that used on the fabrication step of the CuAlO₂/Cu-O HTL within the reported functional inverted perovskite solar cells.

2.1. CuAlO₂/Cu−O Dispersions and Thin Films. Figure 1a shows the XRD patterns for the SolA@RT, SolB@RT, and SolB@400 °C samples described in detail above, which correspond to the CuAlO₂ phase with hexagonal structure (JCPDS 77-2493). These results suggest that the treatment process (sonication and centrifugation) as well as the thermal annealing (up to 400 °C for 20 min) of sample do not alter its crystallinity. However, it should be stressed that the XRD peaks of polycrystalline CuO are overlapping with the peaks of the CuAlO₂ phase and like previous studies, they are not resolved in the XRD patterns of CuAlO₂.40

Figure 21b shows the region of Cu 2p peak of the XPS spectra of the proposed CuAlO₂/Cu−O hole transporting layers. We note that the measurements performed on films (FilmB@400°C) using the identical conditions for the processing of the HTLs within the solar cells reported. The Cu 2p₃/₂ and Cu 2p₁/₂ core-level peaks are observed at 932.6 and 952.6 eV binding energy respectively, with a spin−orbiting splitting of about 20 eV. Between these two peaks a shape-up satellite exists, which is an indication that some of the Cu atoms are present in the CuO or/and Cu(OH)₂ form. The Cu 2p doublet is well fitted by using two peaks, which are assigned to the contributions from the Cu(I) ions (at 932.6 eV binding energy) of the CuAlO₂ phase and the Cu(II) ions (at 935.1 eV binding energy) of CuO and Cu(OH)₂.38,42 On the basis of the XPS peak analysis the estimated percentage of Cu(I) is 77% and that of Cu(II) is 23% (within a ±5% standard error). The main part of Cu(II) species is due to the CuO clusters or Cu−O dimers of hybridized CuAlO₂ lattice, while a minor contribution of Cu(OH)₂ phase cannot be excluded. However, because of the absence of CuO crystalline peaks in the XRD patterns of CuAlO₂ (Figure 1a), the major part of Cu(II) is attributed to be in the form of Cu−O dimers hybridized in CuAlO₂ lattice, which is consistent with previous reported results.40 Therefore, the proposed HTL for inverted perovskite solar cells is termed as CuAlO₂/Cu−O.

As shown in Figure 1c, we measured the UPS spectrum of the CuAlO₂/Cu−O HTL (FilmB@400°C). The work function 4.9 eV was calculated by subtracting from the energy of HeI the high binding energy cutoff, which was determined by linear extrapolation to zero of the secondary electrons edge. While the valence band maximum was resolved by linear extrapolation of the leading edge of the UPS spectra in the Fermi region, the ionization potential was calculated to be 6.45 eV. On the basis of computational results, it was suggested that the valence band region of CuAlO₂ is dominated mainly from the Cu 3d orbitals and to some extend from O 2p states.43,44

Figure 2. SEM images: (a) top view SEM measurement of CuAlO₂/Cu−O thin films (film A annealed at 400 °C for 20 min) with ~85.800× magnification and (b) top view SEM measurement of CuAlO₂/Cu−O thin films (film A annealed at 400 °C for 20 min) with ~12.000× magnification; inset size distributions of the CuAlO₂/Cu−O particles obtained from corresponding film SEM image. (c) Top view SEM measurement of CuAlO₂/Cu−O thin films (film B annealed at 400 °C for 20 min) with ~85.600× magnification and d) top view SEM measurement of CuAlO₂/Cu−O thin films (film B annealed at 400 °C for 20 min) with ~12.000× magnification, inset size distributions of the CuAlO₂/Cu−O particles obtained from corresponding film SEM image.
ionization potential and band gap estimated here were in the range of values reported for CuAlO\textsubscript{2} nonisovalent Cu–O alloysing films that have been used for the fabrication of top gate thin film transistors.\textsuperscript{41}

The UPS measurements have shown that the ionization potential is calculated at 6.45 eV and the Fermi level at 4.9 eV in respect to vacuum. Combining these results with the absorption measurements, where an optical band gap was calculated at \sim 3.5 eV, we can energetically position the conduction band minimum at \sim 2.95 eV. The presence of Cu–O dimers clearly affecting the energy levels and electronic properties of CuAlO\textsubscript{2}.

Such Cu(II) states within the bandgap have been observed at 3.9 eV, in respect to vacuum.\textsuperscript{45} On the other hand, hole carrier selectivity and transport is also influenced by the easily formed acceptor states in the band gap of the CuAlO\textsubscript{2} ascribed to Cu(II) and observed in the range of 23\% from the XPS measurements shown in Figure 1b and other publications.\textsuperscript{40,41} Such Cu(II) states within the bandgap have been observed at around 0.8 eV in respect to VB.\textsuperscript{46} The efficient photovoltaic performance results reported in section 2.3 suggest that these states can assist on the carrier selectivity and transport of the photogenerated holes at perovskite’s VB, which is located at \sim 3.9 eV in respect to vacuum.\textsuperscript{46}

\textbf{2.2. CuAlO\textsubscript{2}/Cu–O Dispersion Treatment, Particles Size Distribution and Surface Topography.} Figure 2 shows the SEM images on films (Film A@400 °C and Film B@400 °C). Figures 2a and 2b show the SEM studies on films fabricated from solution A (Film A@400°C). The SEM image of the films fabricated from solution A (Figure 2a with higher magnification \(\sim 85\ 800\times\)) reveals the presence of large agglomerates (dimensions in the range of 850 nm length and diameter 150 nm, as shown in Figure 2a). Despite the large CuAlO\textsubscript{2}/Cu–O agglomerates, the powders were successfully dispersed in a stable 10 wt % 2-PPE solution via the use of an Avantama surfactant (Sol A@RT).

Figure 2b shows the SEM studies on films fabricated from solution B (Film B@400°C) with lower magnification (\(\sim 12\ 000\times\)). The very large agglomerates dominated the film topography. A very rough surface, with sharp spikes over 700 nm is observed, which is shown in the profilometry measurements included within Figure S2. As a result, the devices made from the untreated CuAlO\textsubscript{2}/Cu–O dispersion (Film A@400°C) were shorted. A hero device fabricated by using the as prepared flame spray pyrolysis based CuAlO\textsubscript{2}/Cu–O dispersions resulted in poor device performance as shown in Figure S3.

To achieve functional CuAlO\textsubscript{2}/Cu–O HTLs the Sol A dispersions were treated with a sonicator probe at high frequencies for over 2 h. In total, more than 60 kJ of energy has been delivered to the dispersion, aiming to force some of the large agglomerates to break into smaller particles. Then, the treated dispersion was centrifuged at 3000 rpm for 4 min. By applying high-energy sonication to the dispersions (Sol B@RT), although large agglomerates are not eliminated (please see Figure 2c) other agglomerates significantly reduced in size. We believe that during the centrifugation process the larger/heavier agglomerates remain at the bottom of the solution and thus by using the top part of the treated dispersion (supernatant dispersion) to fabricated CuAlO\textsubscript{2}/Cu–O HTLs large agglomerates are in most cases avoided. Figure 2c and 2d shows the SEM surface topography of the treated (according to the above process) CuAlO\textsubscript{2}/Cu–O thin film annealed at 400 °C for 20 min in air (Film B@400°C) with higher (\(\sim 85\ 600\times\)) and lower (\(\sim 12\ 000\times\)) magnification respectively, showing that large agglomerates sizes have indeed been reduced. The profilometry measurements included within Figure S2 confirm that based on the above process CuAlO\textsubscript{2}/Cu–O HTL very large sharp spikes can be avoided.

The elimination of the ultralarge agglomerates leads to functional CuAlO\textsubscript{2}/Cu–O films. Comparing the particle size distribution from the SEM measurements on films (insets of Figure 2b and 2d), we see that indeed the proposed processing steps reduce the size of largest CuAlO\textsubscript{2}/Cu–O agglomerates from 1 μm (inset of Figure 2b) to 250 nm (inset of Figure 2d).

Using the above proposed treatment (high-energy sonication, centrifugation) and processing steps (using only the supernatant part of the treated CuAlO\textsubscript{2}/Cu–O dispersion) to avoid the presence of ultra large CuAlO\textsubscript{2}/Cu–O agglomerates we were able to achieve functional CuAlO\textsubscript{2}/Cu–O HTLs, which are fabricated using supernatant solution processing and temperature annealing at 400 °C for 20 min in air. The particle size distribution from the SEM measurements on films (Film B@400°C) fabricated with identical conditions for the preparation of the proposed CuAlO\textsubscript{2}/Cu–O HTLs (insets of Figure 2d) still shows a large distribution of particle sizes from 25 to 250 nm. Thus, even after following the above process 20\% of the small area devices reported (active layer of 0.09
fabricated from solution A (untreated solution), it is shown that due to the presence of large agglomerates, a nonuniform film ranging from 30 to 170 nm is formed. As discussed above using such not-treated CuAlO2/Cu−O HTLs (FilmA@400 °C) resulted to nonfunctional (shorted) perovskite PVs. On the other hand, Figure 4b shows the cross-section SEM image on CuAlO2/Cu−O film (FilmB@400 °C), fabricated from solution B treated solution (high-energy sonication, centrifugation, and processed using only the supernatant part of the treated CuAlO2/Cu−O dispersion). It can be seen clearly that the proposed treatment of CuAlO2/Cu−O dispersion forms a more uniform film (FilmB@400 °C) with smaller agglomerates ranging from 30 to 70 nm compared to the untreated film (FilmA@400 °C). As discussed in detail above and later within the text even though reliability issues are still present using the treated CuAlO2/Cu−O HTLs (FilmB@400 °C) resulted to functional perovskite PVs when the following perovskite active layer step is adequate to cover the particle distribution of the treated CuAlO2/Cu−O HTL (Figure S6, bottom plot).

2.3. CuAlO2/Cu−O Hole Transporting Layers for Inverted Perovskite Solar Cells. The previous paragraphs describe how we fabricate functional CuAlO2/Cu−O films. Within the following sections only the treated CuAlO2/Cu−O solution (high-energy sonication, centrifugation, and processed using only the supernatant part of the treated CuAlO2/Cu−O dispersion) is used. To implement those CuAlO2/Cu−O films in solar cell stack, several other requirements must be fulfilled. Figure 5 shows the cross-sectional view SEM image of the treated, CuAlO2/Cu−O HTL (FilmB@400 °C) perovskite solar cell device structure. The thickness variation due to the broad particle distribution of the treated CuAlO2/Cu−O HTL with estimated average thickness ~50 nm, is indicated within Figure 5 and is discussed in more details within the above sections. The perovskite active layer has a thickness of ~260 nm, while the PC70BM and the AZO films have a thickness of ~75 and ~50 nm, respectively. The perovskite solar cells have been fabricated on CuAlO2/Cu−O HTL (FilmB@400 °C) film coated ITO-glass substrates. Finally, 100 nm of aluminum were thermally evaporated to make an ohmic contact (not shown in Figure 5).

The device structure of the inverted perovskite PVs is based on ITO/[CuAlO2/Cu−O]/CH3NH3PbI3/PC70BM/AZO/Al and is schematically shown in Figure 6a. The bottom electrode should be transparent to guarantee that high amount of light reaches the active layer. Furthermore, because the bottom
electrode serves as the underlayer for the solution-processed perovskite active layer of the solar cell, its surface properties are of high importance.

As shown in Figure 6b, the transparency of the bottom electrode ITO/[CuAlO2/Cu−O] is high, above 80% in the entire visible spectrum (400–800 nm). The high transparency of the proposed ITO/[CuAlO2/Cu−O] bottom electrode guarantees maximum light penetration into the perovskite photoactive layer a parameter, which is important for the corresponding photovoltaic devices. Another important requirement for the bottom electrode of perovskite solar cells is its surface wetting properties. The surface free energy wetting envelopes of the two bottom electrodes under study are shown in Figure 6c. Surface energy wetting envelopes are closed curves that are obtained when the polar fraction of a solid is plotted against the disperse part. As shown in Figure 6c, the calculated surface tension of the CH3NH3PbI3 precursor solution lies well inside the surface energy wetting envelope of the ITO/[CuAlO2/Cu−O] bottom electrode and thus a good wetting is guaranteed. The contact angle of CH3NH3PbI3 precursor solution measured at 28° on top of ITO/[CuAlO2/Cu−O] and at 57° on top of ITO, which proves the efficient surface modification of the ITO surface by the CuAlO2/Cu−O HTLs. The digital pictures of the CH3NH3PbI3 solution droplets on top the layers ITO and ITO/[CuAlO2/Cu−O] are shown in Figure S7.

In addition, the bottom electrode has a major influence on the perovskite crystallization process as well as the hole collection properties of the p-i-n solar cell. To analyze these important aspects, we performed steady state photoluminescence (PL) on the glass/ITO/CH3NH3PbI3 and glass/ITO/[CuAlO2/Cu−O]/CH3NH3PbI3, as well as atomic force microscopy (AFM) measurements on glass/ITO/[CuAlO2/Cu−O]/CH3NH3PbI3 device structure as shown in Figure 7a and 7b, respectively.

As shown in Figure 7a, we compare the steady state PL measurements of CH3NH3PbI3 fabricated on top of glass/ITO and on top of glass/ITO/[CuAlO2/Cu−O]. The PL spectra of CH3NH3PbI3 fabricated on top of glass/ITO shows a strong peak at 764 nm. The strong PL peak at 764 nm is strongly quenched when the CH3NH3PbI3 layer is fabricated on top of ITO/[CuAlO2/Cu−O]. The strong PL quenching of the CH3NH3PbI3 PL peak at 764 nm clearly demonstrates efficient charge transfer of the photogenerated species between the CH3NH3PbI3 photoactive layer and ITO/[CuAlO2/Cu−O] bottom electrode.

To further investigate the influence of the proposed bottom electrode, we conduct AFM measurements as shown within Figure 7b to study the surface topography of CH3NH3PbI3 on...
top of ITO/[CuAlO₂/Cu−O]. The grain size of the polycrystalline CH₃NH₃PbI₃ photoactive layers is relatively large with average grain size in the range of 500 nm. The perovskite active layer is found to be relatively smooth with RMS = 11.2 nm (Figure 7b). These observations show that the CH₃NH₃PbI₃ photoactive layer is well crystallized. The arrow within Figure 7b indicate the presence of perovskite pin-holes most likely due to the broad particle distribution and roughness of the treated CuAlO₂/Cu−O underlayer. In agreement with the AFM image of Figure 7b, the corresponding SEM image is also included within the Figure S8. To reduce the negative impact of perovskite active layer pin-holes aluminum-doped zinc oxide (AZO) electron transporting layer have been included within the inverted perovskite PV device architecture. The use of PC₇₀BM and AZO layers within the device architecture assist on reducing the negative effect of pinholes on the inverted perovskite PV device performance.²³

On the basis of the above experimental results, we have analyzed the nature of the synthesized particles their broad distribution and discuss the treatment and processing steps needed to fabricate functional CuAlO₂/Cu–O HTLs for p–i–n perovskite solar cells. Therefore, p–i–n perovskite photovoltaic devices composed of ITO/[CuAlO₂/Cu–O]/CH₃NH₃PbI₃/PC₇₀BM/AZO/Al were fabricated using the conditions described in detail above and within the experimental section and analyzed, by using current–voltage (J/V) and external quantum efficiency (EQE) measurements,
as well as impedance spectroscopy. As noted also above the treated supernatant CuAlO2/Cu–O HTL-based solar cell power conversion efficiency reproducibility and reliability critically depends on the parameters and accuracy of implementation of the delicate CuAlO2/Cu–O functionalization process described within the paper. For the device analysis provided below a perovskite PV from the best performing device run is used.

As shown in Figure 8a the p–i–n device exhibits negligible hysteresis in the J–V plots between the forward and reverse scan directions. The device exhibited $V_{oc} = 1.07$ V, $J_{sc} = 19.1$ mA/cm$^2$, high FF = 79.6%, and overall PCE = 16.3% in the forward scan with slightly lower overall PCE = 16% in the backward scan. The high open circuit $V_{oc}$ value (1.07 V) is similar to other reported metal oxide based HTLs and to the value we have achieved using monodispersed CuGaO2-based HTLs for inverted perovskite SCs. We note that in our previous work with solvothermal based synthesized CuO interfacial layers for inverted perovskite solar cells the thickness was in the range of 10 nm due to conductivity and transparency limitations. By the incorporation of the proposed treated CuAlO2/Cu–O HTL we have achieved rougher, nonuniform but thicker (in the range of 50 nm) HTLs. When the following perovskite active layer step is effectively covers the broad particle size distribution of the treated CuAlO2/Cu–O HTL the corresponding perovskite PVs exhibits FF values up to ~80%. This is due to the adequate electrical conductivity of CuAlO2/Cu–O HTL as well as the large grain sizes of CH3NH3PbI3 active layer. As a result, the parasitic resistances, known to limit the FF values of perovskite solar cells (i.e., interface and grain boundaries recombination) is suppressed. The latter is also verified with the measurements of the solar cell in dark conditions (Figure 8b), which shows very good diode behavior both in high negative voltages (parallel resistance regime) and in high positive voltages (series resistance regime). In addition, the solar cell performance show photostable performance at maximum power point as shown in Figure S9. Initial lifetime testing under the basic ISOS-D-1 protocol (RT, ambient humidity in the dark) is also performed for the encapsulated inverted PVs showing stable performance for the first 750 h (Figure S10). Despite the observation that devices show stable performance up to 750 h under the basic protocol ISOS-D-1 after this time most of the devices were not able to be measured due to Al contact failure (Al oxidation due to environmental conditions). Thus, the above conditions cannot be used to provide a clear understanding on the degradation mechanisms related to bottom electrode effects of the inverted PV solar cells reported within this paper. More detailed accelerated lifetime testing under light soaking will be presented in future publications. We have recently shown that under temperature accelerated lifetime conditions the interaction of the perovskite active layer with the top Al metal electrode through diffusion mechanisms cause a major thermal degradation pathway for inverted perovskite photovoltaics (PVs). We have proposed thick fullerene diffusion blocking layers to improve the thermal stability of inverted perovskite PVs.

We note that the treated dispersion supernatant batch-to-batch reproducibility critically depends on the sonication and centrifugation parameters used as well as the accuracy of using the top part of the solution after centrifugation process (supernatant dispersion). All the high-performance devices using the CuAlO2/Cu–O HTL reported in this paper achieved by following the implementation of this delicate treatment. We further analyze the device operation by measuring the external quantum efficiency (EQE) as a function of wavelength as shown in Figure 8c. The integrated $J_{sc}$ (18.5 mA/cm$^2$) value extracted from the EQE spectra is in accordance with the J–V measured $J_{sc}$ (19.1 mA/cm$^2$) with only 3% deviation is shown within Figure 68c. In addition, the EQE spectra is above 85% in the range 450–550 nm. In this area, the optical absorption of CH3NH3PbI3 photactive layer is almost 100%. Thus, the high EQE in this optical spectrum is a strong indication that almost all the absorbed photons are converted into charges and collected efficiently at the device electrodes. In the region of 550–700 nm the EQE percentage drops from 75% down to 55% which is most likely due to the drop-in absorption strength of the corresponding CH3NH3PbI3 photactive layers. The peak of the EQE spectra at 700–760 nm can be attributed to the use of AZO nanoparticulate layer at the top electrode resulting in light scattering and increased back electrode reflectivity.

To better understand the photophysical processes and the mechanisms of charge extraction, we conduct impedance spectroscopy measurements for the ITO/[CuAlO2/Cu–O]/CH3NH3PbI3/PC70BM/AZO/Al solar cells. The Nyquist plots were constructed, and the experimental data were fitted into an equivalent circuit model (ECM) as shown in Figure 8d. The ECM used as well as the physical meaning of its components can be found in more details elsewhere. The high frequency response is generally attributed to the charge transport resistance in the carrier selective contacts, whereas the low frequency arc is attributed to the charge recombination resistance of the device. Interestingly, the recombination resistance ($R_{rec}$) value is found to be at ~35 000 Ohm, which is considerably higher compared to the values reported for different HTLs using similar ECMs (e.g., PEDOT:PSS), indicating that there is a considerable decrease to the charge recombination events throughout the device interfaces as well as the individual layers themselves. In addition, the double layer capacitance ($C_{dl}$) is relatively small compared to other commonly used HTLs, indicative of reduced charge accumulation at the interfaces and thus faster hole extraction.

The above mechanisms are the major reasons for the large $V_{oc}$ and FF values measured from the J–V characteristics of the best performing inverted p–i–n perovskite solar cells under study (Figure 8a). Furthermore, to obtain good fits, we note that a constant phase element (CPE) was used for $C_{dl}$ which models the behavior of a nonideal capacitor. CPEs are often necessary to be used to compensate for variations in capacitance arising from phenomena such as nonuniform current distribution and variations in a layer thickness. In our case, we believe that it is a case of nonuniform space charge distribution at the metal-oxide based HTL electrode interfaces.

To summarize, the high temperature flame spray pyrolysis synthesized CuAlO2/Cu–O HTL (Film A annealed at 400 °C for 20 min) shows very large agglomerates, extremely rough surface, with ultrasharp spikes. As a result, devices made from the untreated CuAlO2/Cu–O HTLs were not functional (shorted). To achieve functional CuAlO2/Cu–O HTLs a delicate treatment process applied. The proposed CuAlO2/Cu–O dispersion treatment (high-energy sonication, centrifugation) and processing steps (using only the supernatant part of the treated CuAlO2/Cu–O dispersion) of the high
temperature (particles annealed at 1100 °C for 30 min in air) flame spray pyrolysis CuAlO2/Cu−O (Film B annealed at 400 °C for 20 min) provided functional HTLs for the small active area (0.09 mm²) perovskite PVs reported within this Article. Even though device reproducibility and reliability issues are still present because of delicate functionalization process applied, the treated CuAlO2/Cu−O HTL has adequate wetting properties relevant to CH3NH3PbI3 surface tension requirements and forms a functional ITO/[CuAlO2/Cu−O] bottom electrode as confirmed by the efficient quenching of the photoluminescence. The analysis of the impedance spectroscopy measurements indicates reduced interfacial charge recombination and faster charge collection properties. The optimized photoactive layer properties where the CH3NH3PbI3 fabricated on top of ITO/treated [CuAlO2/Cu−O] HTL shown a Jsc value of 19.1 mA/cm², while provided a high Voc of 1.07 V. The above combined with the FF of 79.6% resulted to champion PCE (forward scan) values of 16.3% and negligible hysteresis effect.

3. CONCLUSION

We presented a solution processed CuAlO2/Cu−O HTLs and discussed their functional operation in p-i-n perovskite solar cells. The CuAlO2/Cu−O powders are obtained by high temperature flame spray pyrolysis. However, we observe that the large agglomerate sizes of the as prepared CuAlO2/Cu−O dispersions are limiting the application of the corresponding HTL for perovskite solar cell applications (corresponding perovskite PVS were shorted). To achieve CuAlO2/Cu−O HTL functionality, the initial CuAlO2/Cu−O dispersion is treated with high frequency sonicator probe, which force some of the large agglomerates to break into smaller CuAlO2/Cu−O particles. Most of the remaining large particles are avoided from the dispersions by centrifugation and taking the top part of the solution (supernatant dispersion) for fabrication of the corresponding HTLs for the development of functional perovskite PVS.

Despite the broad particle sizes distribution by following the above treatment, we have presented the fabrication of transparent and electrically conductive CuAlO2/Cu−O electronic films. We have shown that by the proposed dispersion treatments and processing steps for the fabrication of the CuAlO2/Cu−O HTL functional small area (0.09 mm²) inverted perovskite-based solar cell can be achieved with PCE up to 16.3%. Identification of synthetic/chemical methods to reduce CuAlO2/Cu−O particle sizes and control the particle size distribution will be needed to further improve the device performance and reliability of inverted perovskite solar cells.

4. EXPERIMENTAL SECTION

4.1. Materials. Prepatterned glass-ITO substrates (sheet resistance 4 Ω/sq) were purchased from Piotic Ltd., PbI2 from Alfa Aesar, methylammonium iodide from Dyesol Ltd., Aluminum-doped zinc oxide (AZO) ink from Nanograde (product no N-20X), PC[70]BM from Solenne BV. All the other chemicals used in this study were purchased from Sigma Aldrich.

4.2. Preparation. 4.2.1. Dispersions. The CuAlO2/Cu−O powders were synthesized from Avantama Ltd. by flame spray pyrolysis. For flame spray pyrolysis a metal organic precursor was prepared by a mixture of organic copper and aluminum salt dissolved in a 1:1 mixture of alkyl carboxylic acid and toluene. The metal organic precursors were filtered through a 0.45 mm PTFE syringe filter and fed (5 mL/min) to a spray nozzle, dispersed by oxygen (7 L/min) and ignited by a premixed methane–oxygen flame (CH4, 1.2 L/min; O2, 2.2 L/min). The off-gas was filtered through a glass fiber filter (Schleicher & Schuell) by a vacuum pump (Busch, Seco SV1040CV) at about 20 m³/h. The obtained nanopowder was collected from the glass fiber filter. The as-prepared particles were annealed at 1100 °C for 30 min in air. The powders were stabilized in 2-PPE (10 wt %) using an undecilised Avantama Ltd. surfactant. The treatment of the CuAlO2/Cu−O dispersion by a sonicator probe and the following sonication step was performed at Cyprus University of Technology (CUT). More specifically, to reproduce the manuscript results the flame spray pyrolysis based CuAlO2/Cu−O dispersions must be treated with high frequency sonicator probe for over 2 h and centrifuged at 3000 rpm for 4 min. Importantly after the centrifuged step only the supernatant dispersion should be used to achieve functional devices as reported here.

4.2.2. Device Fabrication. The p-i-n solar cells under study had the structure ITO/[CuAlO2/Cu−O]/CH3NH3PbI3/PC70BM/AZO/Al. ITO substrates were sonicated in acetone and subsequently in isopropanol for 10 min and heated at 120 °C on a hot plate 10 min before use. To form a CuAlO2/Cu−O hole transporting layer, the CuAlO2/Cu−O dispersions used in this study were dynamically spin coated at 3000 rpm for 60 s on the preheated ITO substrates, followed by an annealing step on a hot plate at 400 °C for 20 min in ambient conditions. The perovskite precursor solution was prepared 1 h prior spin coating by mixing lead acetate: methylammonium iodide (1:3) at 36 wt % in DMF with the addition of 1.5 mol % (to PbAc) of methylammonium bromide. The precursor was filtered with 0.1 μm PTFE filters and deposited by static spin coating at 4000 rpm for 60 s followed by annealing for 5 min at 80 °C. The PC70BM solution, 20 mg/mL in chlorobenzene, was dynamically spin coated on the perovskite layer at 1000 rpm for 30 s. AZO ink was dynamically spin coated on top of PC70BM at 3000 rpm for 30 s. Finally, 100 nm Al layers were thermally evaporated through a shadow mask to finalize the devices with active of 0.09 mm². While an estimation of the other layer thicknesses can be performed by the cross-sectional SEM image included in Figure 5. The devices were encapsulated directly after the evaporation of the metal contact, in the glovebox using a glass coverslip and an encapsulation epoxy resin (Ossila E131) activated by 365 nm UV-irradiation.

4.3. Characterization. The thicknesses and the surface profile of the active layers were measured with a Veeeco Dektak 150 profilometer. XRD patterns were collected on a PANalytical X’pert Pro MPD powder diffractometer (40 kV, 45 mA) using Cu Kα radiation (λ = 1.5418 Å). The current density–voltage (J–V) characteristics were characterized with a Botest LIV Functionality Test System. Both forward (short circuit → open circuit) and reverse (open circuit → short circuit) scans were measured with 10 mV voltage steps and 40 ms of delay time. For illumination, a calibrated Newport Solar simulator equipped with a Xe lamp was used, providing an AM1.5G spectrum at 100 mW/cm² as measured by a certified oriel 91150 V calibration cell. A custom-made shadow mask was attached to each device prior to measurements to accurately define the corresponding device area. EQE measurements were performed by Newport System, Model 70356−70316NS. Impedance spectroscopy was performed using an Autolab PGSTAT 302N equipped with FRA32 M module. To extract the Nyquist plots, the devices were illuminated using a red LED at 625 nm and 100 mw/cm². A small AC perturbation voltage of 10 mV was applied, and the current output was measured using a frequency range of 1 MHz to 1 Hz. The steady-state DC bias was kept at 0 V. Transmittance and absorption measurements were performed with a Schimadzu UV−2700 UV−vis spectrophotometer. Steady-state photoluminescence experiments were performed in a Fluorolog3 Horiba Jobin Yvon spectrometer based on an iHR320 monochromator equipped with a visible photomultiplier tube (Horiba T90-04 module). The PL was nonresonantly excited at 400 nm by the line of 5 mW Oxxius laser diode. Atomic force microscopy (AFM) images were obtained using a Nanosurf easy scan 2 controller in tapping mode. Scanning Electron Microscope (SEM) measurements were performed using a Quanta 200 microscope (FEI, Hillsboro, Oregon, USA). X-ray photoelectron spectroscopy
spectra (XPS) and Ultraviolet Photoelectron Spectra (UPS) were recorded by using a Leybold EA-11 electron analyzer operating in constant energy mode at a pass energy of 100 eV and at a constant retardation of 4 eV for XPS and UPS, respectively. The spectrometer energy scale was calibrated by the Au 4f7/2 core level binding energy, BE, (84.0 ± 0.1 eV) and the energy scale of the UPS measurements was referenced to the Fermi level position of Au at a binding energy of 0 eV. All binding energies were referred to the C 1s peak at 284.8 eV of surface adventitious carbon. The X-ray source for all measurements was a nonmonochromatized Al Kα line at 1486.6 eV (12 keV with 20 mA anode current). For UPS measurements, the Hel (21.22 eV) excitation line was used. A negative bias of 12.22 V was applied to the samples during UPS measurements in order to separate secondary electrons originating from the sample and the spectrometer. The sample work function was determined by subtracting the high binding energy cutoff from the Hel excitation energy (21.22 eV). The position of the high-energy cutoff was determined by the intersection of a linear fit of the high binding portion of the spectrum with the background. Similarly, the valence band maximum is determined with respect to the Fermi level, from the linear extrapolation of the valence band edge to the background.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsaem.9b00070.

Additional details for materials characterization, surface properties, processing, and device performance (efficiency and initial lifetime studies) (PDF)

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Notes

The authors declare no competing financial interest.

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