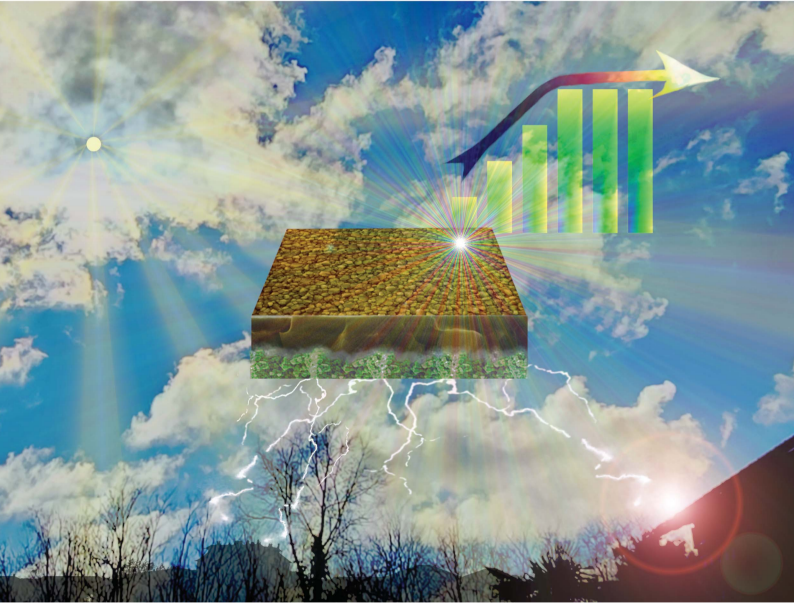
Volume 9 Number 6 21 March 2025 Pages 1391-1606

Sustainable Energy & Fuels

Interdisciplinary research for the development of sustainable energy technologies

rsc.li/sustainable-energy



ISSN 2398-4902



PAPER Vanira Trifiletti *et al.* Enhancing the stability of inverted perovskite solar cells through Cu_2ZnSnS_4 nanoparticles hole transporting material

Sustainable Energy & Fuels



Check for updates

Cite this: Sustainable Energy Fuels, 2025, 9, 1486

Received 20th September 2024 Accepted 8th January 2025

DOI: 10.1039/d4se01309c

rsc.li/sustainable-energy

1. Introduction

The relentless progression of climate change prompted the United Nations to set a new target for achieving a climateneutral world by the mid-21st century. This means maximum

Enhancing the stability of inverted perovskite solar cells through Cu₂ZnSnS₄ nanoparticles hole transporting material[†]

ROYAL SOCIETY OF **CHEMISTRY**

View Article Online

View Journal | View Issue

Elisa Fabbretti,^a Amin Hasan Husien,^a Rahul Patidar, ^b Karen Valadez-Villalobos,^b James McGettrick, ^b Andreia Amighini Alerhush,^a Ershad Parvazian,^b Matthew L. Davies, ^b Trystan Watson, ^b Alessandro Minotto, ^b ^a Giorgio Tseberlidis, ^b ^a Adele Sassella, ^b ^a Vanira Trifiletti ^b *^a and Simona Binetti^a

Perovskite solar cells (PSCs) have burst into photovoltaic (PV) research, revolutionising it and demonstrating that they can achieve performances comparable to technologies already on the market. Despite that, the major limitation of PSCs' commercialisation is their poor environmental stability. Inorganic carriers transporting materials have gained attention for enhancing PSCs' durability due to their superior stability and non-hygroscopic nature. Among them, kesterite Cu_2ZnSnS_4 (CZTS) is a promising inorganic hole transporting material (HTM) thanks to its optimal electronic properties, non-toxic, abundant elements, and simple solution deposition methods. Herein, the effect on the stability of inverted PSCs employing CZTS nanoparticles (NPs) as HTM is investigated by X-ray photoelectron and ultraviolet photoelectron spectroscopy, together with steady-state and time-resolved photoluminescence spectroscopies. Special attention was given to the alteration in the PV performances over time of the CZTS-based device compared to a control device with a common organic HTM. The enhanced stability of the CZTS NPsbased devices in opposition to degradation of the devices with a [2-(3,6-dimethoxy-9H-carbazol-9-yl) ethyl]phosphonic acid (MeO-2PACz) self-assembled monolayer as HTM was demonstrated through solar simulator measurements and external quantum efficiency. The efficiency of a CZTS-based p-i-n PSC increased by 34% after three weeks, while the efficiency of the organic HTM-based device decreased by 16% during the same period. Furthermore, CZTS-based PSCs showed a drop-in efficiency of 80% after four days under a humidity test at 70% RH, while the efficiency of the organic HTM-based devices dropped by 80% after two days under the same conditions. Moreover, analysis of the photoluminescence spectra reveals no modification of the CZTS-based PSCs.

> reduction in greenhouse gas emissions until they are equal to or lower than those naturally absorbed by the planet to achieve a net-zero emissions balance by 2050.1,2 This is necessary to comply with the provisions of the Paris Agreement to limit the temperature rise to 1.5 °C above pre-industrial levels.3,4 At present 73.2% of global greenhouse gases come from energy production from fossil fuels.5 Therefore, urgent and decisive action to transition towards sustainable energy solutions is required. The European Union (EU) has demonstrated a strong commitment to this transition through its European Green Deal, aiming for climate neutrality by 2050.6 Solar photovoltaic (PV) technology stands out due to its vast potential for decarbonised and inexhaustible energy production. In addition, given its scalability and declining cost, solar PV is poised to play a crucial role in meeting the EU objectives.⁷ In this context, emerging thin-film perovskite-based PV technology has garnered significant attention thanks to its impressive performance. The power conversion efficiency (PCE) of perovskite solar cells (PSCs) swiftly increased from about 4% in 2009 to

^aDepartment of Materials Science and Solar Energy Research Centre (MIB-SOLAR), University of Milano-Bicocca, Via Cozzi 55, 20126, Milan, Italy. E-mail: vanira. trifiletti@unimib.it

^bSPECIFIC IKC, Faculty of Science and Engineering, Swansea University, Fabian Way, Swansea, SA1 8EN, UK

[†] Electronic supplementary information (ESI) available: Which reports positions and identifications of the main peaks in XPS spectra for CZTS NPs film annealed at 350 °C, 300 °C and 250 °C; high-resolution XPS spectrum of Cu 2p of CZTS NPs powder, for Cu 2p, Zn 2p, Sn 3d and S 2p components of CZTS NPs film annealed at 300 °C and at 250 °C; XPS spectra of impurities containing N 1s and C 1s; UPS spectrum for CZTS NPs films annealed at 350 ° C, 300 °C and 250 °C; Kubelka–Munk plot; box charts indicating the trend of devices' performances over several days. See DOI: https://doi.org/10.1039/d4se01309c

over 26% in recent laboratory settings.8 This rapid advancement is largely due to the unique properties of metal halide perovskite materials, such as their high absorption coefficients (10⁵- 10^6 cm⁻¹), tuneable direct bandgap (from 1.1 eV to 2.0 eV), high carrier mobility (tens of $\text{cm}^2 \text{V}^{-1} \text{s}^{-1}$),⁹ and long carrier diffusion lengths.¹⁰ These properties enable efficient light absorption and charge transport, making PSCs highly competitive with other PV technologies. Moreover, the simple solution-based fabrication of PSCs offers several advantages, including low costs for raw materials and manufacturing, scalability, and the ability to use flexible substrates. This process involves depositing a series of layers: a transparent conducting oxide (TCO) on a glass substrate, a hole transport material (HTM), the perovskite lightabsorbing layer, an electron transport material (ETM), and a metal back contact. The device architectures are defined by the first layer deposited on the substrate: when the n-type semiconductor is the first on the TCO, the architecture is called conventional and identified as n-i-p.11-13 On the other hand, PSCs in p-i-n architecture, also known as inverted architecture, are increasingly favoured due to their numerous advantages such as enhanced stability and ease of fabrication.14,15 Additionally, the p-i-n structure facilitates efficient charge collection, resulting in higher open-circuit voltages and improved device performances.^{16,17} Also, the ability to process all the layers at sub 150 °C opens up the possibility of low-cost, high throughput manufacturing methods such as roll-to-roll manufacturing.^{18,19} Despite the impressive advancements, several challenges concerning stability remain in the commercial deployment of PSCs. The primary factors causing irreversible degradation of the perovskite photo-absorbing layer include continuous exposure to bias voltage, prolonged exposure to illumination and heat, and moisture.8,20-23 Among these degradation mechanisms, degradation due to humidity is the most detrimental one since metal halide perovskite consists of hygroscopic organic cations, such as methylammonium (MA, $CH_3NH_3^+$) and formamidinium (FA, $CH(NH_2)_2^+$), metal cations as lead (Pb_2^+) and tin (Sn_2^+) , and halide anions, typically iodine (I⁻), bromine (Br⁻), and chlorine (Cl⁻).^{24,25} When water molecules come in contact with the perovskite, reactions to form photoinactive phases occur, generating mainly lead halides and oxides.^{24,25} This issue is being tackled from two angles. First, researchers are developing various strategies to prevent moisture from reaching the perovskite layer, such as using encapsulants and improving the inner layers of PSCs.^{26,27} Second, efforts are focused on enhancing the inherent stability of PSCs in the presence of moisture.^{28,29} In the latter case, one attractive way is to employ inorganic carriers transporting materials which offer superior stability, easier synthesis, and lower cost compared to the commercial organic alternatives.^{21,30-32} Cu₂-ZnSnS₄ nanoparticles (CZTS NPs) emerged as a promising inorganic HTM due to their earth-abundant, non-toxic nature and excellent electronic properties, such as high hole mobility $(\sim 35 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1})$.³³⁻³⁵ Thanks to these features, kesterite has been largely employed as absorber material in thin-film solar cells to substitute Cu(In,Ga)(S,Se)2 and CdTe thin-film solar cells, constituted by rare and toxic elements.³⁶⁻³⁹ However, kesterite PV technology has always shown PCE values lower

than CIGS and CdTe solar cells.⁴⁰ Furthermore, using CZTS NPs as an absorber layer in solar cells does not work due to the excessive presence of grain boundaries.41 However, they have been explored as charge extractors in PSCs,42,43 and recently as electro/photo-catalysts.44,45 The employment of CZTS NPs as HTM has proved its worth since 2015 when Wu and colleagues reached a PCE of 12.8% with a conventional PSC architecture.43 The NPs were dispersed in 1-hexanediol and deposited by spin coating; the perovskite used was the methylammonium lead iodide (CH₃NH₃PbI₃, MAPbI₃). In 2016, Khanzada and colleagues obtained the record PCE of 15.4% with an inverted architecture and MAPbI3 as perovskite.46 The NPs layer was obtained by depositing a solution of ligand-stripped CZTS NPs in N-dimethylformamide (DMF) through spin coating. In 2018, Zuo and colleagues reached a PCE of 13.4% in n-i-p MAPbI₃ PSCs using a wurtzite CZTS NPs HTM.47 In 2019, Ashebir and colleagues prepared CZTS NPs compact films in situ on glass/ ITO substrates by spin coating a novel molecular precursor solution.48 In this way, they could better control the HTM's thickness, achieving a PCE of 6.0% with a 97 nm thick NPs layer. The device had an inverted architecture, and the perovskite was again the MAPbI₃. Although CZTS NPs are employed as HTM due to low-cost precursors, simple deposition methods and the stability they can give to PSCs, few articles in the literature show the effect on PSCs stability. Additionally, stability was always investigated over a small period of time. This is the case of Ashebir and colleagues; they showed that a CZTS NPs-based device, made as mentioned above, was able to retain 87% of its efficiency after 43 days stored in N2 atmosphere.48 Their control device, a p-i-n PSC with PEDOT:PSS as HTM, almost degraded after 35 days of storing. Another work showing the better stability of PSC given by CZTS NPs is the work of Patel and colleagues.49 They obtained n-i-p PSC а (CH₃NH₃Pb_{1-x}Sn_xI_{3-y}Cl_y) with CZTS NPs as HTM that outperformed the organic counterpart (Spiro-OMeTAD) by retaining the 75% of its initial efficiency (9.7%) after 168 h. Then, both the devices started to degrade at a similar rate.

This study explores the integration of CZTS NPs as an innovative HTM to increase the stability of inverted PSCs. The surface composition of CZTS NPs layers annealed at three different temperatures (350 °C, 300 °C, and 250 °C) was analysed. These layers were then incorporated into p-i-n PSCs to determine the optimal annealing temperature. Once optimised, CZTS NPs-based devices were compared to control p-i-n PSCs with the commercial self-assembled monolayer (SAM) of [2-(3,6dimethoxy-9*H*-carbazol-9-yl)ethyl]phosphonic acid (MeO-2PACz) as HTM. The introduction of CZTS NPs as an HTM showed promising results. The optimised CZTS NPs-based PSCs exhibited significant improvements in operational stability compared to the control devices using SAM as HTM. The CZTS NPs-based device demonstrated an increase in efficiency from 4.71% of the first measurement to 6.04% after 18 weeks. Meanwhile, for the SAM-based device, the efficiency decreased from 12.07%, the best value obtained one week after the first measurement, to 6.87% in 18 weeks. This improvement in PCE over time was further investigated and found to result from enhanced charge injection from the MAPbI₃ to the CZTS NPs layer. This suggests the interface between these two layers improves over time, enhancing overall performance.

2. Experimental details

2.1 Materials

Sulphur powder (>99.5%); copper(II) acetate monohydrate $(Cu(CH_3COO)_2 \cdot H_2O, >99\%)$; zinc(II) acetate dihydrate (Zn(CH₃- $COO_2 \cdot 2H_2O$, 99.99%); tin(II) chloride dihydrate (SnCl₂ $\cdot 2H_2O$, >98%); oleylamine (CH₃(CH₂)₇CH=CH(CH₂)₇CH₂NH₂, 98%); pxylene (>99.7%); and terpineol (mixture of isomers) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Ethyl cellulose (ethoxy content 48%, 22 cps) was purchased from Thermo Fisher Scientific (Waltham, MA, USA). Lead(II) iodide (PbI₂, 99.99%) was purchased from TCI (trace metals basis, Tokyo, Japan). Methylammonium iodide (CH₃NH₃I, MAI, 99.99%) was purchased from Greatcell Solar Materials (Australia). Methylamine (ca. 9% in acetonitrile, CH₃NH₂ in CH₃CN, MA in ACN) was purchased from TCI (Tokyo Chemical Industry UK Ltd). Anhydrous acetonitrile (CH₃CN, ACN, \geq 99.9%) was purchased from Fisher Scientific (part of Thermo Fisher Scientific, UK). Hydrochloric acid (HCl, 37%) was purchased from Sigma-Aldrich (St. Louis, MO, USA). MeO-2PACz ([2-(3,6dimethoxy-9H-carbazol-9-yl)ethyl]phosphonic acid, $C_{16}H_{18}NO_5P$, \geq 98.0%) was purchased from TCI (Tokyo Chemical Industry UK Ltd). Extra dry 2-propanol ((CH₃)₂CHOH, IPA, 99.8%) was purchased from Fisher Scientific (part of Thermo Fisher Scientific, UK). [6,6]-Phenyl C₆₁ butyric acid methyl ester (C₇₂H₁₄O₂, PC₆₁BM) was purchased from 1-Material Inc. (Canada). Anhydrous chlorobenzene (C₆H₅Cl, CB, 99.8%) was purchased from Sigma-Aldrich (Germany). Bathocuproine (C₂₆H₂₀N₂, BCP) was purchased from TCI (Tokyo Chemical Industry UK Ltd). Anhydrous ethanol (CH₃CH₂OH, EtOH) was purchased from Thermo Fisher (UK). Silver (Ag) pellets (99.99%) were purchased from Kurt J. Lesker Co. (US). Pre-patterned 15 ohm per sq. glass/ITO substrates (28 \times 28 \times 1.1 mm) were purchased from Kintec Co. (Hong Kong).

2.2 Synthesis of CZTS NPs

CZTS NPs were synthesised by hot injection technique, using a method well-described elsewhere in the literature with minor changes.46,50 The synthesis process begins with the preparation of two solutions. One solution, containing all metal precursor (Sol-M), was obtained by dissolving in 15 mL of oleylamine the reactants in the following order: 959.30 mg Cu $(CH_3COO)_2 \cdot H_2O$, 625.40 mg SnCl₂·2H₂O, and 628.90 mg Zn (CH₃COO)₂·2H₂O. The other solution was prepared by introducing 312 mg of sulphur powder in 10 mL of oleylamine (Sol-S). The two mixtures were prepared in two different three-neck flasks, put in heating mantles and connected to a vacuum/N2 Schlenk line. Sol-S was warmed at 60 °C and degassed for 1 hour under constant stirring. Sol-M underwent a three-step temperature process: (i) degassing at 100 °C under powerful stirring for 1 hour to extract dissolved oxygen and unwanted moisture; (ii) refilling of the flask with N2 while rising slowly the temperature to the injection temperature of 180 °C, during this step the

solution colour turned from bluish to yellowish; (iii) rapid injection of Sol-S into Sol-M after the changes in the Sol-M colour, leading to a further quick colour variation from pale yellow to dark brown. After the injection, the temperature increased to 280 °C, and the solution was left to react for 30 minutes. Then, the heating mantle was removed, and the flask was let to cool down to room temperature in an inert atmosphere. The final suspension was washed with a mixture of chloroform and ethanol in a 1:5 (v/v) ratio and centrifuged at 8000 rpm for 2 minutes. The supernatant was removed, and the washing procedure was repeated several times to remove the exceeding residues of oleylamine from the CZTS NPs surface, which may limit the particle-to-particle interactions and reduce the charge carrier in the HTM film. Then, the CZTS NPs were collected, dried under a vacuum and stored under an N2 atmosphere. In our recently published studies, it has been demonstrated that the synthesis temperature of CZTS NPs is of fundamental importance for the correct formation of the kesterite phase, as confirmed by XRD and TEM, and that purification with HCl effectively removes organic residues (such as oleylamine) as proven by the reported elsewhere FT-IR study.51,52

2.3 Device fabrication

The devices under investigation were PSCs utilising methylammonium lead iodide (CH₃NH₃PbI₃, MAPbI₃) as the perovskite material, configured in a p-i-n architecture (glass/ITO/ CZTS NPs/MAPbI₃/PCBM/BCP/Ag). The control devices differed from the investigated PSCs only in the HTM. The commercial SAM MeO-2PACz was used (glass/ITO/MeO-2PACz/ MAPbI₃/PCBM/BCP/Ag). All the layers were deposited in air atmosphere at 30% RH. Before device fabrication, the patterned glass/ITO substrates $(28 \times 28 \text{ mm}^2)$ were cleaned by ultrasonication with Hellmanex in deionised water (1:40 v/v), pure deionised water, and IPA for 10, 15 and 20 minutes. The substrates were subjected to UV-ozone treatment for 20 minutes before HTM deposition. The CZTS NPs ink was prepared by dispersing 15 mg of NPs in 1 mL of ethyl cellulose in p-xylene solution (5 mg mL^{-1} concentration) in ambient conditions. Then 150 µL of terpineol were added to increase the solution viscosity. The resulting ink was sonicated for 30 minutes to achieve a homogeneous suspension. The CZTS NPs film was obtained by spin coating 70 µL of ink at 4000 rpm for 30 seconds on ITO substrates. Then, the obtained layer was annealed on a titanium hotplate at different temperatures, 350 °C, 300 °C and 250 °C for 2 hours. For the control device, the solution of MeO-2PACz was prepared in anhydrous IPA with 2 mg mL⁻¹ concentration and stirred at 60 °C for 2 hours in a GloveBox with N₂ environment. Before the deposition, it was filtered with a 0.20 µm PTFE filter. Then, 150 µL of solution was deposited on ITO, spin-coated at 4000 rpm for 35 seconds and annealed at 100 °C for 10 minutes. Once cooled, the MeO-2PACz layers were washed with IPA to remove molecules unbonded on the surface and then annealed at 100 °C for 5 minutes.53 MAPbI3 solution was prepared by dissolving 461 mg of PbI₂ and 159 mg of MAI in 1 mL of a mixture of ACN : MA in ACN (1.8 : 3.2 v/v) and stirred at 60 °C for 3 hours under N2 atmosphere in a Glove Box. Before

the coating, the solution was filtered with 0.20 μ m PTFE filter, and 7 μ L of HCl was added per millilitre of perovskite solution.⁵⁴ The perovskite film was obtained by dynamic spin coating of 120 μ L of the described solution at 2000 rpm for 59 seconds and by annealing the film at 110 °C for 10 minutes. For electron transporting material (ETM), PCBM solution was prepared in air by dissolving 40 mg in 1 mL of CB and stirring at 60 °C for 2 hours. The solution was filtered with the 0.20 μ m PTFE filter, and 75 μ L were deposited by spin coating at 3000 rpm for 45 seconds without further annealing at the hot plate. Following the coating of the PCBM layer, BCP (5 mg mL⁻¹ in EtOH) was coated by spin coating 200 μ L of the solution at 7000 rpm for 10 seconds. A 100 nm thick back metal contact of Ag was thermally evaporated to complete the device.

2.4 Characterization

X-ray photoelectron spectroscopy (XPS) analysis on glass/ITO/ CZTS NPs samples was carried out using a Kratos Axis Supra equipped with a monochromated Al Ka source at 225 W (15 kV, 20 mA). Charge calibration was to the C-C peak at 284.8 eV, and charge neutralisation was not required. High-resolution spectra at 40 eV pass energy, 1 s dwell time, and typically 5 sweeps were collected for all elements based on a survey scan at 160 eV pass energy. Ultraviolet photoelectron spectroscopy (UPS) was carried out on the Kratos Axis Supra using the He(1) line, and the electrons collected at a 10 eV pass energy, with the 55 µm aperture to protect the detector. XPS & UPS data were analysed in CasaXPS 2.3.24PR1.0 using the Kratos relative sensitivity factors. Synthetic components were fitted to the raw data using the default GL(30) lineshape. The UV-visible spectra of glass/ ITO/CZTS NPs samples were obtained using a Jasco V-570 UVvisible spectrophotometer in the 300-1300 nm range. A glass/ ITO substrate was used for the baseline. The thickness of CZTS NPs layers was measured with a Veeco Dektak 8 profilometer by applying a stylus with a 2.5 μ m radius and 4.9 μ N of force. A Keithley and an AAA solar simulator (Newport) characterised the complete device's PV performances. J-V characteristics of the devices were obtained under 100 mW cm^{-2} (1 sun calibrated with a reference photodiode with KG5 filter) in the range from 1.2 V to -0.2 V and with a sweep rate of 0.15 V s⁻¹. A metal mask employed during measurements gave the devices an active area of 0.09 cm². External Quantum Efficiency (EQE) spectra were obtained using a custom-built setup. This system included Xenon arc and Quartz halogen white light sources (Bentham) and a Bentham TMc300 monochromator, all controlled by Labview software. The photocurrent of the cell was measured with a trans-impedance amplifier (Femto), while the incident light intensity was calibrated using a silicon reference photodiode (Newport). The samples were scanned in the wavelength range of 300–850 nm. The active area was 0.05 cm^2 , defined by the incident light spot on the sample. After the solar simulator and EQE measurements, the samples were stored in a GloveBox with an N2 environment. A high humidity environment was simulated in a Temperature and Humidity Test Chamber by applying the standard condition of 70% RH at 30 $^{\circ}$ C. Steady-state photoluminescence (PL) measurements were

conducted using an FS5 Edinburgh Instruments spectrofluorometer, covering a wavelength range of 690 nm to 850 nm. The samples were excited at a wavelength of 500 nm, with a 520 nm bandpass filter in the excitation pathway and a 496 nm longpass filter in the emission pathway. Time-resolved photoluminescence (TRPL) measurements were performed using an FLS1000 Edinburgh Instruments photoluminescence spectrometer. The samples were excited with an EPL405 laser (λ exc = 405 nm), and the emission was recorded at 775 nm with a 30 nm emission slit width. The laser repetition period was set to 1 µs. The TRPL data were fitted using a bi-exponential function to determine the trap-mediated (τ_1) and radiative (τ_2) recombination lifetimes.55 The morphology of the samples was compared using scanning electron microscopy (SEM) imaging using a Gemini 500 Zeiss microscope. Moreover, AFM images were also collected on the samples over 5 μ m \times 5 μ m areas using a Bruker Multimode Nanoscope V in intermittent contact mode with silicon tips (spring constant = 40 Nm^{-1} , resonance frequency \approx 300 kHz) and with an image resolution of 512 \times 512 pixels.

3. Results and discussion

3.1 CZTS NPs layer

Stable and performing PSCs through inorganic HTMs can be achieved with an optimal compact and transparent HTM layer. Therefore, this study's preliminary stage focused on optimising ink suspension and CZTS NPs film deposition. At first, the idea was to avoid additives or stabilisers in the ink because they could have limited the charge extraction in the device. Unfortunately, the resulting films presented drawbacks such as voids, insufficient compactness, and poor adhesion to the substrate, even if the solvent, being nonpolar,56 was indicated to disperse the NPs. Ethyl cellulose was introduced in the CZTS NPs suspension as a small organic binder, while terpineol was added to increase the solution viscosity. In this way, it was possible to set the optimal spin coating parameters and obtain a compact, void-free, transparent CZTS NPs layer. The downside of adding ethyl cellulose was that this polymer thermally degrades from around 250.7 °C up to around 378.4 °C,57 leading to the need for the CZTS NPs layer annealing temperature higher than 250 °C. However, this was critical for ITO since the resistivity of this transparent conductive oxide rapidly increases above 250 °C, and the film begins to degrade,57 thermally reducing PSCs' operating performances. In our experiments, heat treatment at 350 °C for 2 hours likely compromised the conductivity of ITO, contributing to a reduction in device performance.58 One alternative is tin fluoride (FTO), known for its higher thermal stability or decreasing ink deposition temperature. To overcome these limitations, we are currently testing conductive inks processable at temperatures below 100 ° C, which could offer a solution to reduce the thermal load on the substrate during device fabrication. This paves the way for further studies on CZTS NPs layer deposition optimisation. The present work investigated the optimal annealing temperature at 350 °C, 300 °C and 250 °C to get a compromise between removing ethyl cellulose, the ITO degradation in terms of CZTS NPs layer composition and PSCs PV performances. Three samples in the configuration glass/ITO/CZTS NPs were prepared by annealing the kesterite layer at the abovementioned temperatures. From here on out, the CZTS NPs layers annealed at different temperatures, 350 °C, 300 °C and 250 °C, will be named 350-CZTS, 300-CZTS and 250-CZTS layers. The surface chemical composition of the films was investigated using XPS analysis. The band alignment of the three HTMs with the MAPbI₃ perovskite was investigated by getting the valence band maximum (VBM) by UPS analysis and by calculating the conduction band minimum (CBM) by extrapolating the bandgap from UV-vis spectra.

XPS measurements confirmed the valence states of Cu¹⁺, Zn²⁺, Sn⁴⁺ and S²⁻ in the CZTS NPs film. Fig. 1 shows the highresolution spectra of Cu 2p, Zn 2p, Sn 3d and S 2p core levels in the 350-CZTS layer. The doublet reported in Fig. 1a corresponds to Cu 2p_{3/2} and Cu 2p_{1/2} peaks, respectively, at 932.9 eV and 952.7 eV. The $2p_{3/2}$ peak position and the split of the two Cu 2p spin-orbit components of 19.8 eV indicate the oxidation state +1 of Cu.59 However, a high binding energy shoulder and a series of satellite peaks between 940.0 and 945.0 eV can be seen, indicating the presence of Cu^{2+} and, consequently, of secondary species such as CuO or CuSO4.60,61 Also, the satellite peak at 962.7 eV comes from the presence of copper(II) oxide.62 Since in the XPS spectrum of Cu 2p of the CZTS NPs powder, the satellite peaks are not present (Fig. S1, ESI[†]), these are probably due to the oxidation of CZTS NPs film surface due to the high annealing temperature. The two peaks at 1022.2 eV and 1045.3 eV in Fig. 1b are assigned to the two core levels $2p_{3/2}$ and 2p_{1/2} of Zn 2p. Fig. 1c shows the Zn L₃M_{4.5}M_{4.5} Auger emission at a binding energy of 499.6 eV, or a kinetic energy of 986.9 eV, is consistent with a Zn²⁺ species.^{41,63} Fig. 1c also shows the Sn 3d_{5/2} and 3d_{3/2} doublets located at 486.9 eV and 495.3 eV, respectively, with a doublet separation of 8.4 eV that confirms Sn's oxidation state +4.64 The situation is more complex for the sulphur 2p region (Fig. 1d), where four peaks can be detected. At 161.5 eV and 162.6 eV, the first two correspond to S 2p3/2 and S 2p1/2, respectively, with a spin-orbit component split of 1.1 eV. This is consistent with the binding energy range between 160 and 164 eV expected for S in the sulphide phases.64,65 indicating sulphur in the S²⁻ oxidation state. The other two peaks located at higher energies are due to other bonding states of sulphur. Specifically, in the energy range between 168.0 and 171.0 eV, the peaks are related to sulphur's metal sulphate phases.⁶⁶ Therefore, the peak at 168.9 eV can be assigned to the 2p_{3/2} spin-orbit component of sulphur in the CuSO₄ compound,^{66,67} The peak at 170.0 eV corresponds to the $2p_{1/2}$ spin–orbit component of S in the same compound. The shoulder confirms the existence of the CuSO4 phase around 935.0 eV in the Cu 2p spectrum (Fig. 1a), which Buldu et al. have assigned to Cu bonded to the sulphate ion $SO_4^{2-.67}$ The formation of this secondary phase can be attributed to the annealing process of the CZTS NPs film since the Cu 2p spectrum of the nanoparticle powder does not report the CuSO₄ shoulder before annealing (Fig. S1[†]).

XPS spectra were also collected for 300-CZTS and 250-CZTS layers to study the effects of the annealing temperature on the layer composition. A comparison of the main XPS peak

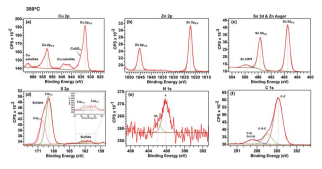


Fig. 1 High-resolution XPS spectra of (a) Cu 2p, (b) Zn 2p, (c) Sn 3d and (d) S 2p components of CZTS NPs film annealed at 350 °C, and XPS spectra of impurities containing (f) N 1s and (e) C 1s.

positions for the different analysed annealing temperatures is reported in Table S1.[†] Compared to the spectra of the 350-CZTS film in Fig. 1, the spectra related to the 300-CZTS sample (Fig. S2[†]) have the main peaks of Cu 2p, Zn 2p, Sn 3d and S 2p core levels in almost the same positions and with similar doublet splitting energies (Table S1[†]). This is also true for the other peaks as Cu satellites, CuSO₄ shoulder (Fig. S2a[†]), Zn Auger LMM (Fig. S2c[†]) and S 2p of sulphate (Fig. S2d[†]). The only differences between 350-CZTS and 300-CZTS samples are in the ratio of the sulphide and sulphate peaks: the relative sulphide peak area is higher in the film annealed at 300 °C, and in the relative sulphate peak area, which is reduced for the 300-CZTS layer. This can suggest that an annealing temperature lower than 300 °C can promote the presence of S^{2-} instead of SO_4^{2-} . This can be confirmed especially by the S 2p spectrum of the CZTS film annealed at 250 °C (Fig. S3d[†]), where the areas under the sulphide peaks are higher than for the sulphide peaks in the 300 °C and 350 °C spectra. Additional evidence that the 250 °C annealing temperature favours the presence of S in the sulphide phase can be found in the Cu 2p spectrum in Fig. S3a.† Here, the CuSO₄ shoulder at 935.0 eV has almost wholly disappeared compared to the 300 °C and 350 °C Cu 2p spectra (Fig. S2a† and 1a respectively). This can indicate a reduced formation of copper sulphate at low annealing temperatures. Considering again the Cu 2p spectrum at 250 °C, it is noticeable that the Cu satellite peaks, and thus most of Cu(II) secondary phases, disappear. All these facts together suggest that at low annealing temperatures, the oxidation of the film surface and the formation of secondary phases are reduced. Nevertheless, device performances are not increased using a low-temperature annealed film, as shown in the discussion about device characterisation by the solar simulator in Section 2.4 of the present study. The low operational performance of the 250-CZTS device can be related to the shoulder at 163.6 eV in the S 2p spectrum (Fig. S3d[†]). As Shanthi et al. points out, this shoulder can be related to a -S-C- bond, which can be due to a reaction between sulphur and oleylamine.68 This solvent has a boiling point between 348 and 350 °C. Therefore, it is possible that at 250 °C, the oleylamine is not entirely removed from the CZTS NPs film, reducing the solar performance of the devices. The hypothesis of the presence of oleylamine is reinforced by the N 1s core level

spectrum of the 250-CZTS layer, Fig. S3e,† which presents a peak around 400.0 eV related to the organic amine (A) group.⁶³ The high-resolution XPS for N 1s has shown the amine peak also for the 350-CZTS and 300-CZTS layers Fig. 1e and S2e,† respectively, suggesting the presence of oleylamine for these two layers. In the identical spectra, there is also a peak around 402.0 eV, which Hellgren et al.69 attributed to quaternary amine (QA), and in this case, it may be due to some byproduct of oleylamine degradation. Notably, it can be consistent with a range of partly oxidised organic amines due to oleylamine thermal degradation in the presence of oxygen and oxygencontaining molecules. The related A and QA peak positions are reported in Table S1.† Traces of oleylamine can also be detected in C 1s core level spectra of the 350-CZTS, 300-CZTS and 250-CZTS samples, Fig. 1f, S2f and S3f⁺ respectively. At high binding energies, all the spectra present a peak around 288.6 eV, which Ansari et al.70 attributed to a combination of C-N and O-C=O peaks due to oleylamine. These spectra also show a peak at 286.4 eV, referable to the C-O-C bonds and, thus, to ethyl cellulose.⁷¹ Unfortunately, the presence of both oleylamine and ethyl cellulose is confirmed for all three samples, indicating that the conductivity of 350-CZTS, 300-CZTS and 250-CZTS HTMs could be reduced by these nonconductive molecules, also limiting the PV performances of PSCs. The last peak, at 284.8 eV in these three spectra, is related to the C-C bond, which can still be linked to both oleylamine and ethyl cellulose since these molecules present this bond in their structure. However, it must be noticed that carbon contaminations are pretty frequent, and the large area under this peak may be due to the exposure of the layers to air during both deposition and annealing steps.

Once the surface chemical composition was known, we investigated how the three different annealing temperatures affected the band gap (E_{gap}) and the energy levels of the 350-CZTS, 300-CZTS and 250-CZTS NPs films. To this aim, XPS and UPS analysis were employed to obtain the VBM and the work function (WF) of the three layers, and UV-vis analysis was used to get the band gap values. VBM_{xps} values were found by extrapolating the linear portion of VB XPS spectra at lower energies and intersecting this with the spectral baseline, as reported in Fig. 2.⁷² These values are obtained *versus* the Fermi

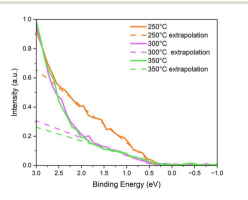


Fig. 2 Extrapolation of the VBM values from VB XPS spectra of CZTS NPs film annealed at 350 °C, 300 °C and 250 °C. The VBM values are 0.3 eV, 0.5 eV and 0.4 eV, respectively.

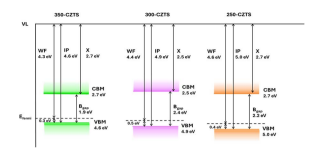


Fig. 3 Electronic structures for the 350-CZTS, 300-CZTS and 250-CZTS layers. The distances of VBM from E_{Fermi} were obtained thanks to XPS analysis; WFs were extrapolated from UPS spectra; and the B_{gap} values were obtained through UV-vis measurements. All the other values have been calculated.

energy, E_{Fermi} , which has to be considered at 0 eV, as shown in Fig. 2. The resulting values have been summed with the WF values to find the VBM level *versus* the vacuum level (VL), as reported in the band diagram in Fig. 3.

WFs of the 350-CZTS, 300-CZTS and 250-CZTS NPs films have been obtained by the formula:⁷³

$$WF = h\nu - E_{\rm cutoff} \tag{1}$$

Where $h\nu$ is the photon energy of the UV source (He–I, $h\nu = 21.2$ eV) and $E_{\rm cutoff}$ is the secondary electron energy cutoff at high binding energies. The linear extrapolation of the UPS spectra's high and low binding energies can determine this last energy. Fig. S4[†] shows UPS spectra and the $E_{\rm cutoff}$ extrapolation for the three CZTS NPs layers. By combining the VBM by XPS and the WF by UPS, it was possible to calculate also the ionisation potential (IP) through the formula IP = WF + VBM_{xps}, as reported in the electronic structure (Fig. 3). The band alignment was completed by extrapolating the optical $E_{\rm gap}$ for each sample through UV-vis analysis. Notably, from the reflectance (*R*) measurements of the samples, it was possible to estimate the Kubelka–Munk relation:

$$F(R) = \frac{(1-R)^2}{2R}$$
(2)

With F(R) named Kubelka–Munk function.⁷⁴ The Kubelka– Munk curves, Fig. S5,[†] were obtained by plotting $(F(R) h\nu)$,² where hv is the photon energy, as a function of hv. By extrapolating the linear part of the curves to the x-axis, the optical E_{gap} for each CZTS NPs HTM was determined to be 1.92 eV, 2.40 eV and 2.34 eV for the 350-CZTS, 300-CZTS and 250-CZTS films respectively. Finally, the conduction band minimum (CBM) and the electron affinity (χ) were estimated by the formula CBM = VBM $- B_{gap}$ and $\chi = IP - B_{gap}$ and reported in Fig. 3. The obtained B_{gap} values are large compared to the typical values reported in the literature around 1.4-1.6 eV.75 However, it must be noticed that the charge confinement and the electronic structure change in the nanoscale.76 Moreover, the thickness of these layers was on the nanoscale, while thicker layers are reported in the literature on the microscale, which again affects the absorption and bandgap of thin films.77,78

Table 1 Solar cell parameters for devices with CZTS NPs HTM annealed at 350 °C, 300 °C and 250 °C

Sample	$J_{ m sc}~({ m mA~cm}^{-2})$	$V_{\rm oc}$ (V)	FF (%)	PCE (%)
350 °C	14.64	0.77	41.58	4.71
300 °C	13.75	0.28	27.68	1.08
250 °C	17.56	0.41	29.21	2.12

3.2 Perovskite cell stability investigation

Here, the stability of PSCs was tested by comparing devices with two different HTMs: the inorganic CZTS NPs and the commercial organic MeO-2PACz. To this aim, complete devices were characterised through a solar simulator, External Quantum Efficiency (EQE) and moisture measurements over a prolonged time. JV measurements were first performed on devices with 350-CZTS, 300-CZTS and 250-CZTS HTMs to investigate the optimal temperature for the best performance. As reported in Table 1, the 350-CZTS device has shown the best performance compared to the other two, even though XPS measurements of the 250-CZTS layer showed the best phase composition with a reduced number of secondary phases (Fig. S3[†]). This is why it performed better than the 300-CZTS based device. However, the 250 °C and 300 °C layers may still contain oleylamine after film annealing, as suggested by the XPS analysis. Oleylamine, with a high boiling point of around 350 °C, is difficult to remove at lower temperatures. Since oleylamine is a non-conductive molecule, its presence can decrease the photovoltaic performance of the devices by hindering hole extraction. The high series resistance supports this observed in both the 250-CZTS and 300-CZTS devices, Fig. 4, which indicates poor charge extraction at the interfaces.

Compared to SAM HTM, PSCs with CZTS NPs HTM showed poor photovoltaic performance with low J_{sc} and FF. This may be due to the non-uniformity of the HTM layer, composed of semiconductive NPs interconnected by an insulating polymer, ethyl cellulose. A future challenge emerging from this work could be incorporating organic conductive molecules into the CZTS NPs layer. More importantly, SEM images of the SAM and CZTS NPs films reveal significant differences in surface morphology. The AFM and SEM analyses (Fig. S6†) show that

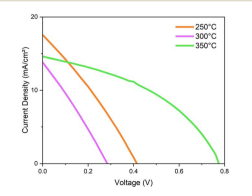


Fig. 4 Illuminated JV curves for the devices based on CZTS NPs HTM annealed at 350 $^\circ$ C, 300 $^\circ$ C and 250 $^\circ$ C.

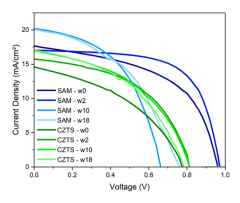


Fig. 5 JV curves under illuminated conditions of the CZTS-based and the control devices measured over days to investigate the stability of both the perovskite solar cells.

the SAM (MeO-2PACz) films offer uniform coverage at sub-µm and tens-of-µm scales. At a scale of 200 nm, regular structures associated with the molecular organisation of the MeO-2PACz are observed, with height variations substantially confined to less than 10 nm. In contrast, the CZTS films, while effectively covering the perovskite, exhibit a relatively much higher density of inhomogeneities. Namely, large agglomerates are detected, both by AFM and SEM, with lateral size and height exceeding 300 nm. The irregular structure of the CZTS film compromises the effective charge extraction since the agglomerates observed on the surface of the CZTS film could create localised points prone to short-circuiting the device. Efforts are underway to develop a novel deposition method to prevent their formation, thereby enhancing the uniformity of the CZTS NPs films.

Based on the above results, the 350-CZTS HTM was selected to investigate perovskite solar cell stability further. The performance of the CZTS-based device was compared to a control device using the common SAM HTM, as shown in Fig. 5. This figure presents the illuminated current-voltage curves of the two investigated devices, the CZTS-based and the control perovskite solar cells. Fig. 5, along with Tables 2 and 3, reveals a decline in performance for the control device, while the performance of the CZTS-based solar cell improves, indicating superior stability.

This trend is further confirmed by the box charts in Fig. S7 and S8,† which display the solar parameters of 64 additional devices. The enhanced stability of the CZTS-based device is also evident in the EQE spectra shown in the ESI (Fig. S9).† With

Table 2 Solar cell parameters of the CZTS-based device over weeks (w)

CZTS	$J_{ m sc}~({ m mA~cm^{-2}})$	$V_{\rm oc}$ (V)	FF (%)	PCE (%)
w0	14.64	0.77	41.58	4.71
w0 w1	16.55	0.70	47.59	5.89
w2	15.72	0.81	49.91	6.38
w3	16.57	0.83	45.30	6.24
w10	16.97	0.81	45.59	6.26
w16	16.18	0.79	48.89	6.24
w18	17.09	0.78	45.08	6.04

Table 3 Solar cell parameters of the SAM-based control device over weeks $\left(w\right)$

SAM	$J_{ m sc}~({ m mA~cm}^{-2})$	$V_{\rm oc}$ (V)	FF (%)	PCE (%)
w0	17.75	0.96	52.58	8.94
w1	18.24	0.96	68.60	12.07
w2	17.05	0.97	65.60	10.84
w3	17.60	0.97	49.38	8.44
w10	20.16	0.66	50.07	6.67
w16	16.87	0.79	53.51	7.11
w18	20.04	0.76	45.18	6.87

time, the control device decreases the overall quantum efficiency in the region below 600 nm and above. At the short wavelength, the dominant recombination occurs near the front contact interface (HTM/MAPbI₃), so EQE confirms the improvement with time of the charge extraction operated by CZTS NPs while highlighting the degradation of the SAM efficiency.12,79 EQE also justifies the overall lower efficiency of the CZTS-based device, showing that the parasitic absorption of the HTM heavily impacts the light that the device can exploit.12,79 Transmittance spectra (Fig. S10[†]) confirm this evaluation and highlight the reduced optical transmittance of the CZTS NPs layer compared to the MeO-2PACz one. This reduction in transmittance, particularly in the visible range, heavily impacts the light available for effective photoelectric conversion. This can be overcome by inverting the architecture to exploit the stabilising properties of CZTS NPs. Accordingly, it is increasingly urgent to design an ink that can be deposited at lower temperatures than those explored here.

A discrepancy between the $J_{\rm sc}$ derived from the EQE integration and that obtained using the solar simulator (SS) was observed, with $J_{\rm sc-EQE}$ higher than $J_{\rm sc-SS}$. Although less common, this behaviour can be explained by considering two main factors.

First, the illuminated area during the J-V measurements (0.09 cm^2) is larger than that used for the EQE measurements (0.05 cm^2) . Perovskites exhibit a non-uniform chemical composition and defect concentration across the film, and a larger tested area can lead to lower PV performance.⁸⁰ Second, the solar simulator used is calibrated with a photodiode designed for organic cells, which partially filters the UV components of the spectrum.⁸¹ This spectral mismatch can affect the measurement of $J_{\text{sc-SS}}$ compared with the EQE, which uses a calibrated monochromatic source. The observed results are, therefore, consistent with the intrinsic characteristics of perovskites and with the experimental conditions adopted.

The superior performance of the CZTS-based cell is further illustrated by the IV characteristic measured in dark conditions (Fig. S11†), where the built-in potential is absent over different days for the CZTS-based device, in contrast to the control device. The minimum of the dark curves at zero voltage for the CZTS NPs-based devices suggests that the CZTS/MAPbI₃ interface is less prone to ion accumulation than that between SAM and MAPbI₃.⁸² However, the dark current in the CZTS-based PSCs is higher than that of the conventional devices. In the former, the

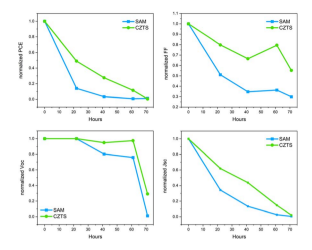


Fig. 6 PSCs parameters trends after various hours under moisture conditions.

dark current can be due to the non-uniformity of the CZTS NPs layer or thermal charge generation at the CZTS NPs/MAPbI₃ interface. Ollearo et al. demonstrated that the activation energy (E_{a}) of this thermal process is comparable to the energy difference φ between the CBM of the perovskite and the VBM (or HOMO level, depending if the HTM is organic or not) of the HTM.83 Therefore, they suggest that the VBM of the HTM also influences the dark current due to the thermally activated charge generation. In our case, a higher dark current for the CZTS NPs-based PSCs could suggest that the band alignment between CZTS and MAPbI3 is not ideal-however, the HOMO level for SAM MeO-2PACz reported by Ollearo et al.83 It is close to the VBM of the 350-CZTS layer reported in Fig. 3, around 5.2 eV. Thus, in the present case, the E_a may be due to oleylamine and ethyl cellulose residues affecting the interface between the CZTS NPs HTM and the MAPbI₃, potentially reducing the thermal charge generation $E_{\rm a}$ at this interface.

To further assess the stability enhancement by the inorganic HTM, a reference device and a CZTS-based device were subjected to highly humid conditions (70% relative humidity at 30 ° C) and periodically measured using a solar simulator. Fig. 6 presents the trends in photovoltaic parameters observed over various hours of exposure to high humidity. After 72 hours, both devices ceased to function, although the performance of the CZTS NPs HTM device was consistently better than that of the reference device. This suggests that the perovskite degradation was slower in the CZTS-based device despite a silver back contact, which typically accelerates perovskite degradation under humid conditions.

3.3 CZTS NPs/MAPbI₃ interface

Over time, the improved stability and increased PV performance of the CZTS NPs-based PSCs suggest a different interaction between the MAPbI₃ and the CZTS NPs than between the perovskite and the SAM. Steady-state photoluminescence (PL) and time-resolved photoluminescence (TRPL) analyses were conducted on fresh and aged samples to understand these

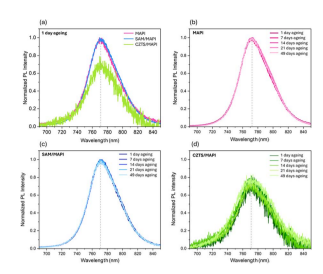


Fig. 7 Steady-state PL spectra of (a) MAPbl₃, SAM/MAPbl₃ and CZTS/ MAPbl₃ compared at day 1 of ageing. PL spectra from day 1 to day 49 of ageing of (b) MAPbl₃, (c) SAM/MAPbl₃ and (d) CZTS/MAPbl₃ samples.

interfaces better. These measurements were performed on the investigated CZTS NPs/MAPbI3 interface, the control SAM/ MAPbI₃ interface, and the bare MAPbI₃ layer. Fig. 7 shows the normalised PL spectra for the three samples, glass/ITO/MAPbI₃, glass/ITO/350-CZTS/MAPbI3 and glass/ITO/SAM/MAPbI3, and their evolution over time. The samples were stored in air atmosphere, in a box with silica gel absorbing part of the environmental moisture, to evaluate any change in PL spectra due to the deterioration of the HTM/MAPbI3 interfaces. From the spectra of day one, it is possible to see that all the peaks are centred around 770.2 nm, in line with the PL peak position of MAPbI₃, 770.0 nm.⁸⁴ Additionally, after days of ageing, the peak position remained consistent, suggesting the absence of phase segregation, typically evidenced by a shift in the peak position. The main difference among these spectra is the reduced emission of the MAPbI₃ on the CZTS NPs layer.

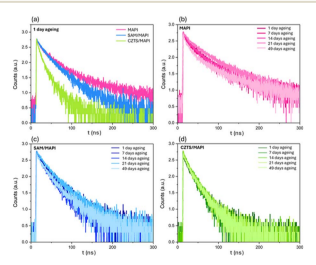


Fig. 8 TRPL spectra of (a) MAPbI₃, SAM/MAPbI₃ and CZTS/MAPbI₃ compared at day 1 of ageing. PL spectra from day 1 to day 49 of ageing of (b) MAPbI₃, (c) SAM/MAPbI₃ and (d) CZTS/MAPbI₃ samples.

Table 4 Trap-mediated recombination lifetime (τ_1) and the radiative recombination lifetime (τ_2) values of the MAPbI_3, SAM/MAPbI_3 and 350-CZTS/MAPbI_3 samples

τ (ns)	MAPbI ₃	SAM/MAPbI ₃	350-CZTS/MAPbI ₃
τ_1	19.33	17.16	12.67
τ_2	65.98	44.93	30.14

This result, along with the faster decay of the CZTS TRPL spectra compared to the SAM spectra, Fig. 8, suggests that the photogenerated holes are rapidly transferred from the perovskite to our HTM, reducing the number of radiatively recombining charge carriers. This results in more efficient hole extraction at the CZTS NPs/MAPbI3 interface.85 Table 4 shows the trapmediated recombination lifetime (τ_1) and the radiative recombination lifetime (τ_2) values of the three PL samples. The faster quenching of the PL emission in the CZTS NPs/MAPbI₃ sample indicates a more effective charge transfer at this interface again. The notable behaviour observed in the CZTS-based devices is the increase in efficiency can be attributed to a different interaction between the MAPbI₃ perovskite and the CZTS compared to the SAM, with the CZTS promoting a more efficient charge transfer. These results align with recent studies highlighting the effectiveness of chalcogenide materials in passivating lead-related defects (Pb2⁺ and Pb⁰), thus increasing the device stability.⁸⁶

4. Conclusions

This study successfully demonstrated the enhanced stability of inverted PSCs using CZTS NPs as an HTM. Incorporating CZTS NPs led to significantly improved device longevity compared to organic HTMs, such as MeO-2PACz. Devices with CZTS NPs showed more resilience to environmental degradation, particularly humidity, and increased PCE over time, suggesting that the CZTS NPs/MAPbI₃ interface matures, enhancing charge transfer efficiency. Specifically, the optimised CZTS-based devices retained higher performance after several weeks, outperforming their organic HTM counterparts. PL and TRPL analyses showed that CZTS NPs improved hole extraction efficiency from the perovskite layer, resulting in better charge transfer and reduced recombination than organic HTMs. Although challenges remain, such as improving the uniformity and conductivity of the CZTS layer and decreasing the treatment's temperature, this research paves the way for developing more robust, sustainable, and efficient perovskite solar cells by leveraging earth-abundant materials like CZTS, making PSCs more viable for commercial applications.

Data availability

The data supporting this article have been included in the ESI.†

Author contributions

EF: data curation, formal analysis, funding acquisition, investigation, methodology, visualisation, writing – original draft,

writing – review & editing; AHH: investigation, methodology, writing – review & editing; RP: supervision, validation, writing – review & editing; KVV: investigation, resources, validation, visualisation, writing – original draft; JMG: investigation, resources, validation, visualisation, writing – original draft; AAA: data curation, formal analysis; EP: resources, supervision; MLD: resources, supervision; TW: funding acquisition, resources, supervision; AM: data curation, formal analysis, writing – review & editing; GT: methodology, validation, writing – review & editing; AS: funding acquisition, resources, validation, supervision; VT: conceptualisation, data curation, formal analysis, funding acquisition, investigation, methodology, project administration, supervision, validation, writing – original draft, writing – review & editing; SB: funding acquisition, project administration, resources, supervision.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors gratefully thank the VIPERLAB project funded by the European Union's Horizon 2020 research and innovation program under grant agreement No. 101006715. Thanks also to the University of Milan-Bicocca through the fund 2023-ATEQC-0078, Fondo di Ateneo Quota Competitiva. The authors also thank the MUSA (Multilayered Urban Sustainability Action) project, funded by the European Union - NextGenerationEU, under the National Recovery and Resilience Plan (NRRP) Mission 4 Component 2 Investment Line 1.5: Strengthening of research structures and creation of R&D "innovation ecosystems", set up of "territorial leaders in R&D". JM thanks the EPSRC Programme Grant ATIP (Application Targeted and Integrated Photovoltaics) (EP/T028513/1) and the Advanced Imaging of Materials (AIM) facility at Swansea University which was funded in part by the EPSRC (EP/M028267/1) and the European Regional Development Fund through the Welsh Government (80708).

References

- 1 W. L. Filho, T. Wall, A. L. Salvia, M. A. P. Dinis and M. Mifsud, *Sci. Rep.*, 2023, **13**, 20582.
- 2 N. Della Valle, G. Ulpiani and N. Vetters, *Humanities and Social Sciences Communications*, 2023, vol. 10, p. 440.
- 3 E. Karola Velten, C. Calipel, M. Duwe, N. Evans, C. Felthöfer, J. Gardiner, M. Hagemann, F. Hossfeld, C. Humphreys, L. Kahlen, S. Lalieu, M. Leśniak, P. Schöberlein, A. Śniegocki, A. Stefańczyk and J. Tarpey, *State of EU progress to climate neutrality*, European Climate Neutrality Observatory, 2024.
- 4 S. Matemilola, O. Fadeyi and T. Sijuade, in *Encyclopedia of Sustainable Management*, ed. S. Idowu, R. Schmidpeter, N. Capaldi, L. Zu, M. Del Baldo and R. Abreu, Springer International Publishing, Cham, 2020, ch. 516-1, pp. 1–5.
- 5 H. Ritchie and M. Roser, Our World in Data, 2024.

- 6 E. Commission, Research and Innovation for Climate Neutrality by 2050: Challenges, Opportunities and the Path Forward, Luxembourg, 2024.
- 7 I. Kougias, N. Taylor, G. Kakoulaki and A. Jäger-Waldau, *Renewable Sustainable Energy Rev.*, 2021, **144**, 111017.
- 8 R. Jose, T. M. Brown and J. K. Ling, *Perovskite Solar Cells: Prospects of Commercialization*, Elsevier Science, 2024.
- 9 L. M. Herz, ACS Energy Lett., 2017, 2, 1539-1548.
- 10 Y. Cheng, Y. Peng, A. K. Y. Jen and H.-L. Yip, *Sol. RRL*, 2021, 6, 2100545.
- 11 T. Webb, S. J. Sweeney and W. Zhang, *Adv. Funct. Mater.*, 2021, **31**, 2103121.
- 12 V. Trifiletti, N. Manfredi, A. Listorti, D. Altamura, C. Giannini, S. Colella, G. Gigli and A. Rizzo, *Adv. Mater. Interfaces*, 2016, **3**, 1600493.
- 13 V. Trifiletti, V. Roiati, S. Colella, R. Giannuzzi, L. De Marco, A. Rizzo, M. Manca, A. Listorti and G. Gigli, ACS Appl. Mater. Interfaces, 2015, 7, 4283–4289.
- 14 N. G. Park, Adv. Energy Mater., 2019, 10, 1903106.
- 15 M. R. Samantaray, Z. Wang, D. Hu, M. Yuan, H. Song, F. F. Li,
 G. Jia, L. Ji, X. Zou, H. H. Shen, B. Xi, Y. Tian, X. Q. Xu,
 D. J. Lee and H. Y. Hsu, *Sol. RRL*, 2024, 8, 2400235.
- 16 Z. Guo, A. K. Jena, G. M. Kim and T. Miyasaka, *Energy Environ. Sci.*, 2022, **15**, 3171–3222.
- 17 P. Caprioglio, J. A. Smith, R. D. J. Oliver, A. Dasgupta, S. Choudhary, M. D. Farrar, A. J. Ramadan, Y. H. Lin, M. G. Christoforo, J. M. Ball, J. Diekmann, J. Thiesbrummel, K. A. Zaininger, X. Shen, M. B. Johnston, D. Neher, M. Stolterfoht and H. J. Snaith, *Nat. Commun.*, 2023, 14, 932.
- 18 D. Burkitt, R. Patidar, P. Greenwood, K. Hooper, J. McGettrick, S. Dimitrov, M. Colombo, V. Stoichkov, D. Richards, D. Beynon, M. Davies and T. Watson, *Sustainable Energy Fuels*, 2020, 4, 3340–3351.
- X. Zhang, L. Shen, P. Baral, S. N. Vijayaraghavan, F. Yan, X. Gong and H. Wang, *Sol. Energy Mater. Sol. Cells*, 2022, 246, 111894.
- 20 Y. Ma, Z. Lu, X. Su, G. Zou and Q. Zhao, *Adv. Energy Sustainability Res.*, 2022, **4**, 2200133.
- 21 P. Zhu, C. Chen, J. Dai, Y. Zhang, R. Mao, S. Chen, J. Huang and J. Zhu, *Adv. Mater.*, 2024, **36**, e2307357.
- 22 Y. Ding, Q. Guo, Y. Geng, Z. Dai, Z. Wang, Z. Chen, Q. Guo,Z. Zheng, Y. Li and E. Zhou, *Nano Today*, 2022, 46, 101586.
- 23 C. Duan, A. Tang, Q. Guo, W. Zhang, L. Yang, Y. Ding, Z. Dai and E. Zhou, *Adv. Funct. Mater.*, 2024, **34**, 2313462.
- 24 K. Sun and P. Müller-Buschbaum, *Energy Technol.*, 2023, **11**, 2201475.
- 25 B. P. Kore, M. Jamshidi and J. M. Gardner, *Mater. Adv.*, 2024, 5, 2200–2217.
- 26 S. Baumann, G. E. Eperon, A. Virtuani, Q. Jeangros, D. B. Kern, D. Barrit, J. Schall, W. Nie, G. Oreski, M. Khenkin, C. Ulbrich, R. Peibst, J. S. Stein and M. Köntges, *Energy Environ. Sci.*, 2024, **17**, 7566–7599.
- 27 H. Zhang, J. W. Lee, G. Nasti, R. Handy, A. Abate, M. Gratzel and N. G. Park, *Nature*, 2023, **617**, 687–695.
- 28 F. Kong, E. Güzel and S. Sonmezoglu, *Mater. Today Energy*, 2023, **35**, 101324.

- 29 İ. C. Kaya, R. Ozdemir, H. Usta and S. Sonmezoglu, *J. Mater. Chem. A*, 2022, **10**, 12464–12472.
- 30 S. Sajid, S. Alzahmi, I. B. Salem, J. Park and I. M. Obaidat, *Mater. Today Energy*, 2023, **37**, 101378.
- 31 S. Akin, Y. Liu, M. I. Dar, S. M. Zakeeruddin, M. Grätzel, S. Turan and S. Sonmezoglu, *J. Mater. Chem. A*, 2018, 6, 20327–20337.
- 32 S. Akin, F. Sadegh, S. Turan and S. Sonmezoglu, *ACS Appl. Mater. Interfaces*, 2019, **11**, 45142–45149.
- 33 M. A. K. Sanhoury, in *Handbook of Energy Materials*, ed. R. Gupta, Springer Nature Singapore, Singapore, 2023, ch. 68-1, pp. 1–23.
- 34 J. J. Chaudhari and U. S. Joshi, *J. Mater. Sci.: Mater. Electron.*, 2020, **31**, 14411–14420.
- 35 A. Le Donne, V. Trifiletti and S. Binetti, *Front. Chem.*, 2019, 7, 297.
- 36 U. A. Shah, A. Wang, M. Irfan Ullah, M. Ishaq, I. A. Shah, Y. Zeng, M. S. Abbasi, M. A. Umair, U. Farooq, G. X. Liang and K. Sun, *Small*, 2024, **20**, e2310584.
- 37 G. Tseberlidis, C. Gobbo, V. Trifiletti, V. Di Palma and S. Binetti, *Sustainable Mater. Technol.*, 2024, **41**, e01003.
- 38 G. Tseberlidis, V. Trifiletti, A. Le Donne, L. Frioni, M. Acciarri and S. Binetti, *Sol. Energy*, 2020, 208, 532–538.
- 39 V. Trifiletti, S. Mostoni, F. Butrichi, M. Acciarri, S. Binetti and R. Scotti, *ChemistrySelect*, 2019, **4**, 4905–4912.
- 40 NREL, The National Renewable Energy Laboratory. Best Research-Cell Efficiency Chart, https://www.nrel.gov/pv/cellefficiency.html, accessed 24/11/2024.
- 41 B. Ananthoju, J. Mohapatra, D. Bahadur, N. V. Medhekar and M. Aslam, Sol. Energy Mater. Sol. Cells, 2019, 189, 125–132.
- 42 U. Syafiq, N. Ataollahi and P. Scardi, *Sol. Energy*, 2020, **196**, 399–408.
- 43 Q. Wu, C. Xue, Y. Li, P. Zhou, W. Liu, J. Zhu, S. Dai, C. Zhu and S. Yang, *ACS Appl. Mater. Interfaces*, 2015, 7, 28466– 28473.
- 44 A. S. Nazligul, M. Wang and K. L. Choy, *Sustainability*, 2020, **12**, 5138.
- 45 P. Semalti, V. Sharma and S. N. Sharma, *J. Cleaner Prod.*, 2022, **365**, 132750.
- 46 L. S. Khanzada, I. Levchuk, Y. Hou, H. Azimi, A. Osvet,
 R. Ahmad, M. Brandl, P. Herre, M. Distaso, R. Hock,
 W. Peukert, M. Batentschuk and C. J. Brabec, *Adv. Funct. Mater.*, 2016, 26, 8300–8306.
- 47 Y. Zuo, L. Chen, W. Jiang, B. Liu, C. Zeng, M. Li and X. Shi, *Mater. Tehnol.*, 2018, 52, 483–486.
- 48 G. Y. Ashebir, C. Dong, Z. Wan, J. Qi, J. Chen, Q. Zhao, W. Chen and M. Wang, *J. Phys. Chem. Solids*, 2019, **129**, 204–208.
- 49 S. B. Patel, A. H. Patel and J. V. Gohel, *CrystEngComm*, 2018, 20, 7677–7687.
- 50 C. Steinhagen, M. G. Panthani, V. Akhavan, B. Goodfellow,
 B. Koo and B. A. Korgel, *J. Am. Chem. Soc.*, 2009, 131, 12554–12555.
- 51 A. H. Husien, G. Tseberlidis, V. Trifiletti, E. Fabbretti, S. Mostoni, J. McGettrick, T. Watson, R. Po and S. Binetti, *Nanoscale Adv.*, 2024, 7, 250–260.

- 52 G. Tseberlidis, V. Trifiletti, A. H. Husien, A. L'Altrella, S. Binetti and F. Gosetti, *Appl. Sci.*, 2024, **14**, 9923.
- 53 N. Vanni, R. Po, P. Biagini, G. Bravetti, S. Carallo, A. Giuri and A. Rizzo, *Nanomaterials*, 2024, **14**, 107.
- 54 D. Burkitt, R. Swartwout, J. McGettrick, P. Greenwood,
 D. Beynon, R. Brenes, V. Bulovic and T. Watson, *RSC Adv.*, 2019, 9, 37415–37423.
- 55 E. V. Pean, S. Dimitrov, C. S. De Castro and M. L. Davies, *Phys. Chem. Chem. Phys.*, 2020, **22**, 28345–28358.
- 56 P. Kush and S. Deka, Mater. Chem. Phys., 2015, 162, 608-616.
- 57 Y. A. Aggour, J. Mater. Sci., 2000, 35, 1623-1627.
- 58 Z. Xu, P. Chen, Z. Wu, F. Xu, G. Yang, B. Liu, C. Tan, L. Zhang, R. Zhang and Y. Zheng, *Mater. Sci. Semicond. Process.*, 2014, 26, 588–592.
- 59 J. Wang, P. Zhang, X. Song and L. Gao, *RSC Adv.*, 2014, 4, 27805–27810.
- 60 S. Ahmadi, N. Khemiri, A. Cantarero and M. Kanzari, *J. Alloys Compd.*, 2022, **925**, 166520.
- 61 C. Mongioví, G. Crini, X. Gabrion, V. Placet, V. Blondeau-Patissier, A. Krystianiak, S. Durand, J. Beaugrand, A. Dorlando, C. Rivard, L. Gautier, A. R. L. Ribeiro, D. Lacalamita, B. Martel, J.-N. Staelens, A. Ivanovska, M. Kostić, O. Heintz, C. Bradu, M. Raschetti and N. Morin-Crini, *Chem. Eng. J. Adv.*, 2022, **10**, 100282.
- 62 S.-Y. Ahn, K. Park, D. Choi, J. Park, Y. J. Kim and H.-S. Kim, *Electronics*, 2019, **8**, 1099.
- 63 A. V. Naumkin, A. Kraut-Vass, S. W. Gaarenstroom and C. J. Powell, NIST X-ray Photoelectron Spectroscopy Database, https://srdata.nist.gov/xps/, accessed 29/12/2024.
- 64 J. Chen, F. Wang, B. Yang, X. Peng, Q. Chen, J. Zou and X. Dou, *Coatings*, 2019, 9, 130.
- 65 W. Liu, B. Guo, X. Wu, F. Zhang, C. L. Mak and K. Wong, J. Mater. Chem. A, 2013, 1, 3182–3186.
- 66 J. Chastain and R. C. King Jr, Handbook of X-ray photoelectron spectroscopy, Perkin-Elmer Corporation, 1992, vol. 40, p. 221.
- 67 D. G. Buldu, A. Cantas, F. Turkoglu, F. G. Akca, E. Meric, M. Ozdemir, E. Tarhan, L. Ozyuzer and G. Aygun, *Phys. Scr.*, 2018, **93**, 024002.
- 68 P. M. Shanthi, P. J. Hanumantha, K. Ramalinga, B. Gattu, M. K. Datta and P. N. Kumta, J. Electrochem. Soc., 2019, 166, A1827–A1835.
- 69 N. Hellgren, R. T. Haasch, S. Schmidt, L. Hultman and I. Petrov, *Carbon*, 2016, **108**, 242–252.
- 70 S. M. Ansari, B. B. Sinha, D. Sen, P. U. Sastry, Y. D. Kolekar and C. V. Ramana, *Nanomaterials*, 2022, **12**, 3015.
- 71 J. Tang, S. Li, Y. Wang, H. Zhang, B. Lin and M. Sun, Prog. Org. Coat., 2022, 163, 106617.
- 72 C. Tamin, D. Chaumont, O. Heintz, A. Leray and M. Adnane, *EPJ Photovoltaics*, 2022, **13**, 24.
- 73 S. Pandharkar, S. Rondiya, A. Bhorde, S. Nair, R. Aher,
 P. Vairale, A. Waghmare, D. Naik, R. Waykar and
 Y. Jadhav, *J. Mater. Sci.: Mater. Electron.*, 2019, 30, 14781–14790.
- 74 Y. Tan, D. Fang, N. Cheng, F. Xiong, W. Ding, B. Mi and Z. Gao, *Mater. Sci. Semicond. Process.*, 2023, **166**, 107710.
- 75 R. Aruna-Devi, M. Latha, S. Velumani and J. A. Chávez-Carvayar, *Rare Met.*, 2019, 40, 2602–2609.

- 76 J. Jasieniak, M. Califano and S. E. Watkins, *ACS Nano*, 2011, 5, 5888–5902.
- 77 S. ğit Gezgin and H. Ş. Kiliç, Opt. Mater., 2020, 101, 109760.
- 78 S. Yiğit Gezgin and H. Ş. Kiliç, *Indian J. Phys.*, 2022, **97**, 779–796.
- 79 H. Fujiwara, M. Kato, M. Tamakoshi, T. Miyadera and M. Chikamatsu, *Phys. Status Solidi A*, 2018, **215**, 1700730.
- 80 G. Yan, Y. Yuan, M. Kaba and T. Kirchartz, *Adv. Energy Mater.*, 2024, 2403706.
- 81 S. Ye, B. Chen, Y. Cheng, M. Feng, H. Rao, Y. M. Lam and T. C. Sum, *J. Phys. Chem. Lett.*, 2020, **11**, 3782–3788.
- 82 J. Y. Huang, Y. W. Yang, W. H. Hsu, E. W. Chang, M. H. Chen and Y. R. Wu, *Sci. Rep.*, 2022, **12**, 7927.

- 83 R. Ollearo, J. Wang, M. J. Dyson, C. H. L. Weijtens, M. Fattori,
 B. T. van Gorkom, A. van Breemen, S. C. J. Meskers,
 R. A. J. Janssen and G. H. Gelinck, *Nat. Commun.*, 2021, 12, 7277.
- 84 J. I. Khan, F. H. Isikgor, E. Ugur, W. Raja, G. T. Harrison,
 E. Yengel, T. D. Anthopoulos, S. De Wolf and F. Laquai, ACS Energy Lett., 2021, 6, 4155–4164.
- 85 S. N. Habisreutinger, B. Wenger, H. J. Snaith and R. J. Nicholas, *ACS Energy Lett.*, 2017, **2**, 622–628.
- 86 A. Sadhu, Y. Guo, T. Salim, Q. Sun, S. G. Mhaisalkar,
 T. C. Sum and L. H. Wong, *Adv. Funct. Mater.*, 2023, 33, 2305215.