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# Low-loss contacts on textured substrates for inverted perovskite solar cells

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#### 1 Low-loss contacts on textured substrates for inverted perovskite solar cells

- 2 So Min Park<sup>1,2,10</sup>, Mingyang Wei<sup>3,10</sup>, Nikolaos Lempesis<sup>4,10</sup>, Wenjin Yu<sup>5</sup>, Tareq Hossain<sup>6</sup>, Lorenzo
- 3 Agosta<sup>4</sup>, Virginia Carnevali<sup>4</sup>, Harindi R. Atapattu<sup>6</sup>, Peter Serles<sup>7</sup>, Felix T. Eickemeyer<sup>3</sup>, Heejong
- 4 Shin<sup>1</sup>, Maral Vafaie<sup>2</sup>, Deokjae Choi<sup>1</sup>, Kasra Darabi<sup>8</sup>, Eui Dae Jung<sup>2</sup>, Yi Yang<sup>1</sup>, Da Bin Kim<sup>2</sup>, Shaik
- 5 M. Zakeeruddin<sup>3</sup>, Bin Chen<sup>1</sup>, Aram Amassian<sup>8</sup>, Tobin Filleter<sup>7</sup>, Mercouri G. Kanatzidis<sup>1</sup>, Kenneth
- 6 R. Graham<sup>6</sup>, Lixin Xiao<sup>5</sup>, Ursula Rothlisberger<sup>4</sup>, Michael Grätzel<sup>3</sup><sup>†</sup> & Edward H. Sargent<sup>1,2,9</sup><sup>†</sup>
- <sup>7</sup> <sup>1</sup>Department of Chemistry, Northwestern University, Evanston, Illinois 60208, United States.
- 8 <sup>2</sup>Department of Electrical and Computer Engineering, University of Toronto, Toronto, Ontario,
- 9 M5S 3G4, Canada.
- <sup>3</sup>Laboratory of Photonics and Interfaces, Ecole Polytechnique Fédérale de Lausanne, Lausanne,
   1015, Switzerland.
- <sup>4</sup>Laboratory of Computational Chemistry and Biochemistry, Ecole Polytechnique Fédérale de
   Lausanne, Lausanne, 1015, Switzerland.
- <sup>5</sup>State Key Laboratory for Artificial Microstructure and Mesoscopic Physics, Department of
- 15 Physics, Peking University, Beijing 100871, P. R. China.
- <sup>6</sup>Department of Chemistry, University of Kentucky, Lexington, KY, 40506, United States.
- <sup>7</sup>Department of Mechanical and Industrial Engineering, University of Toronto, Toronto, Ontario,
  M5S 3G8, Canada.
- 19 <sup>8</sup>Department of Materials Science and Engineering, and Organic and Carbon Electronics
- 20 Laboratories (ORaCEL), North Carolina State University, Raleigh, NC, 27695, United States.
- <sup>9</sup>Department of Electrical and Computer Engineering, Northwestern University, Evanston, Illinois
- 22 60208, United States.

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<sup>10</sup>These authors contributed equally to this work: So Min Park, Mingyang Wei, Nikolaos Lempesis
<sup>†</sup>e-mail: <u>ted.sargent@northwestern.edu</u>; <u>michael.graetzel@epfl.ch</u>

Inverted perovskite solar cells (PSCs) promise enhanced operating stability compared to 26 their normal-structure counterparts $^{1-3}$ . To improve efficiency further, it is crucial to 27 combine effective light management with low interfacial losses<sup>4,5</sup>. Here we develop a 28 conformal self-assembled monolayer (SAM) as the hole-selective contact on light-managing 29 textured substrates. Molecular dynamics simulations indicate cluster formation during 30 phosphonic acid adsorption leads to incomplete SAM coverage. We devise a co-adsorbent 31 strategy that disassembles high-order clusters, thus homogenizing the distribution of 32 phosphonic acid molecules, thereby minimizing interfacial recombination and improving 33 electronic structures. We report a lab-measured power-conversion efficiency (PCE) of 25.3% 34 and a certified quasi-steady-state PCE of 24.8% for inverted PSCs, with a photocurrent 35 approaching 95% of the Shockley-Queisser maximum. An encapsulated device having a PCE 36 of 24.6% at room temperature retains 95% of its peak performance when stressed at 65°C 37 and 50% relative humidity following >1000 hours of maximum power point tracking under 38 1-sun illumination. This represents one of the most stable PSCs subjected to accelerated 39 ageing - achieved with a PCE surpassing 24%. The engineering of phosphonic acid 40 adsorption on textured substrates offers a promising avenue for efficient and stable PSCs. It 41 is also anticipated to benefit other optoelectronic devices that require light management. 42

Perovskite solar cells (PSCs) have recently attained a certified efficiency of 26.1%<sup>6</sup>; however, the very highest PCE devices have yet to meet operating stability under accelerated ageing tests<sup>5</sup>. The limited device stability is attributed to the presence of mobile and hygroscopic p-type dopants in hole-transporting layers, which undermine moisture and thermal stability<sup>7</sup>. Inverted PSCs present a solution by utilizing undoped hole-selective contacts<sup>8</sup>. Recent studies have shown PCEs surpassing 25% for inverted PSCs<sup>2</sup>. However, when adhering to the stringent quasi-steady-state (QSS) protocol, their certified efficiency (~24%) requires further improvement<sup>3,9</sup>.

Efforts to improve inverted PSC efficiency have focused on interface passivation<sup>2,9,10</sup>. This 50 approach suppresses non-radiative recombination and thus improves fill factors and 51 photovoltages<sup>11</sup>. Nevertheless, higher photocurrents are needed for efficiency gains. In normal-52 structure PSCs, this has been realized using textured fluorine-doped tin oxides (FTO) as the 53 transparent conductive oxides (TCO)<sup>4,5</sup>. The pyramidal grains on FTO minimize reflection losses 54 and extend the length of the average optical path<sup>4</sup>. By contrast, inverted PSCs, often built on 55 smooth indium tin oxides (ITO)<sup>9,10</sup>, face substantial optical losses due to the lack of light 56 management. 57

The discrepancy is ascribed to differences in transporting materials and deposition techniques. In normal-structure PSCs, inorganic hole-blocking layers are conformally deposited on textured substrates using spray pyrolysis and chemical bath deposition<sup>4,5</sup>. However, inverted PSCs typically employ ultra-thin (<5 nm) organic hole-selective contacts on substrates<sup>2,3</sup>, which can be challenging to deposit uniformly using solution processing<sup>12</sup>. Inhomogeneity leads to energy losses and insufficient carrier extraction<sup>13</sup>.

Self-assembled monolayers (SAMs), particularly those composed of phosphonic acid molecules with hole-selective tail groups, have shown promise in addressing this issue<sup>14</sup>. Phosphonic acids establish coordinative/covalent bonds with TCOs, allowing for sufficient SAM coverage on rough surfaces<sup>15</sup>. This could provide a low-loss interface, uniting light management with interface passivation. Indeed, photocurrent improvements have been observed when replacing SAMmodified ITO with FTO substrates in inverted PSCs<sup>16</sup>.

Despite the promise, achieving a high-density, closely packed SAM remains challenging, which
 leads to erratic interfacial properties<sup>17</sup>. For example, M. Liu et al. revealed that phosphonic acids
 failed to fully cover textured wafers, which affected the performance of tandem solar cells<sup>18</sup>. Even

on flat substrates, the formation of SAMs tends to be non-uniform, a problem usually attributed to
 the limited solubility of phosphonic acids<sup>19</sup> and their insufficient chemical bond formation with
 metal oxides<sup>20</sup>.

Several strategies have been proposed to ensure conformal SAM coverage, such as utilizing more reactive oxide surfaces (e.g., nickel oxides) to promote hetero condensation<sup>18</sup>, employing thermal evaporation for SAM deposition<sup>21</sup>, and designing highly soluble phosphonic acid molecules<sup>19</sup>. However, these methods can be laborious, requiring time-intensive molecule design, synthesis, and vacuum-based layer deposition. Additionally, the use of highly reactive oxides could compromise device stability due to redox reactions with ammonium halides<sup>22</sup>.

#### 82 Molecular dynamics simulations

We sought to deepen understanding of how phosphonic acids interact with textured TCOs. We 83 began by looking at the case of 2-(9H-carbazol-9-yl)ethyl)phosphonic acid (2PACz), an organic 84 molecule (Fig. 1a) capable of forming SAMs on TCOs<sup>14</sup>, and investigated its interactions with 85 FTO. 2PACz has found extensive applications as a hole-selective contact in PSCs in view of its 86 excellent defect passivation and deep highest occupied molecular orbital (HOMO) level<sup>21</sup>. The 87 surface properties of FTO were approximated using tin oxides (SnO<sub>2</sub>), given that commercial FTO 88 has a fluorine-doping level under 0.5% (atomic percentage)<sup>23</sup>. Flat vs. corrugated SnO<sub>2</sub> were 89 analysed to understand the effects of surface morphology (Supplementary Fig. 1). 90

Molecular dynamics (MD) simulations suggest that 2PACz molecules can agglomerate, forming 91 dimers, trimers, and tetramers (Fig. 1b and Supplementary Fig. 2). Within the initial 1 nanoseconds 92 of simulations, 90% of the final number of clusters has been reached (Fig. 1b), and dimers were 93 the most prevalent among clusters (Fig. 1c). Compared to flat surfaces, textured surfaces led to 94 more clusters (e.g., 7 vs. 0 for 20 2PACz molecules on 6-nm-wide SnO<sub>2</sub>) (Supplementary Note 1; 95 Supplementary Figs. 3 and 4). The cluster formation is closely linked to the inhomogeneity of 96 SAMs on FTO. For instance, when 162 2PACz molecules are positioned on top of textured SnO<sub>2</sub> 97 surfaces, despite their initial random distribution (Supplementary Video 1), the finial equilibrated 98 99 structures exhibit clear phase segregation (Fig. 1d), where almost 85% of the SnO<sub>2</sub> surface remains 100 uncovered by any 2PACz molecules (Supplementary Fig. 5). These observations were also verified in a larger system (22-nm width; 1134 2PACz molecules) (Supplementary Fig. 6). 101

102 Reasoning that a thiol group (-SH) could interact with phosphonic acids, and a carboxyl group (-COOH) attached to FTO, we introduced 3-mercaptopropionic acid (3-MPA) (Fig. 1a) in 103 104 simulations to break apart 2PACz clusters particularly on textured substrates. The approach is akin to the co-adsorbent strategy used in dye-sensitized solar cells to reduce dye aggregation on 105 titanium oxides<sup>24</sup>. Similar strategies have been employed to improve the quality of hole-selective 106 SAMs<sup>25,26</sup>. With 3-MPA at a molar ratio of 1:9 with 2PACz, we saw a relative 15% decrease in 107 the number of clusters on textured surfaces (Fig. 1b), and a slower formation rate ( $20 \text{ ns}^{-1} \text{ vs. } 28$ 108  $ns^{-1}$ ) (Supplementary Fig. 7). In particular, the higher-order clusters (trimers and tetramers) 109 decreased by 53% (Fig. 1c). As a result, the mixed system shows less phase segregation (Fig. 1e 110 and Supplementary Video 2), leading to extended surface coverage (67% vs. 15%) by 2PACz 111 molecules (Supplementary Fig. 5). Ab initio MD (AIMD) simulations (Supplementary Note 2 and 112 Supplementary Fig. 8) indicate that the presence of 3-MPA hinders the free movement of 113 individual 2PACz molecules by forming a supramolecular structure (Fig. 1f). This reduces 114 agglomeration with already-formed dimers, suppressing the formation of higher-order clusters 115 (Supplementary Video 3). Density functional theory (DFT) calculations revealed surface binding 116 energies of -3.2 eV and -2.6 eV for 2PACz and 3-MPA, respectively, which are much stronger 117 than the interaction energies of 2PACz clusters (-0.3 to -0.1 eV per molecule). This indicates 118 robust anchoring of 2PACz and 3-MPA molecules upon contact with substrate surfaces. 119

#### 120 Characterization of organic contacts

We deposited both 2PACz and a mixture of 2PACz:3-MPA on FTO substrates using solution processing, followed by annealing the film at 100°C and solvent washing (Methods). Micelles were detected in the processing solutions through dynamic light scattering (DLS) (Supplementary Fig. 9). However, due to their limited proportion in the volume distribution compared to 2PACz monomers, film formation primarily resulted from molecular diffusion rather than direct micellar adsorption<sup>27</sup>.

The film composition and its interaction with FTO substrates were investigated using X-ray photoelectron spectroscopy (XPS) and attenuated total reflectance Fourier transform infrared spectroscopy (ATR-FTIR), respectively. For the film comprising solely 2PACz molecules (the control), the presence of 2PACz was evidenced by the appearance of the phosphine (P) doublet peak (Fig. 2a) and the characteristic C-N peak and C-C/C-H peak (Supplementary Fig. 10) in the

XPS spectra<sup>21</sup>. In the case of mixed film, 3-MPA was indicated by the sulfur (S) 2p doublet peak 132 at around 163.8 eV (i.e., the thiol group) (Fig. 2a)<sup>28</sup>. From ATR-FTIR, bidentate or tridentate 133 134 binding was identified as the mode of 2PACz-FTO interactions, given the appearance of the symmetric  $PO_3^{2-}$  stretching (at 996 cm<sup>-1</sup> for the control sample)<sup>29,30</sup> and the disappearance of P-135 OH vibrations<sup>14</sup>. These also indicated that solvent washing removed unbound molecules<sup>14</sup>. With 136 3-MPA addition, a redshift of approximately 5 cm<sup>-1</sup> was observed for the PO<sub>3</sub><sup>2-</sup> vibrational mode, 137 indicating the enhanced surface binding of 2PACz - as confirmed by AIMD simulations 138 (Supplementary Note 3; Supplementary Figs. 11 and 12). ATR-FTIR of the FTO/3-MPA sample 139 (Fig. 2b) further showed carboxylate-related peaks and a reduction in the intensity of the C=O 140 peak, which correspond to the bidentate chelation of 3-MPA with FTO surfaces<sup>31</sup>. Combining the 141 results from XPS and ATR-FTIR, we reasoned that 3-MPA functioned as a co-adsorbent, 142 143 modulating the interaction of 2PACz with FTO substrates.

The resultant SAM uniformity was characterized by Kelvin probe force microscopy (KPFM). 144 Topography images revealed that, despite varying SAM modifications, the surface morphology of 145 FTO substrates was retained due to the ultra-thin nature of SAMs (Supplementary Fig. 13). From 146 the respective contact potential difference (CPD) maps (Fig. 2c), we recorded surface potential 147 variations on 2PACz-modified surfaces within a range of 120 mV. A Gaussian fit of the single 148 149 peak in the CPD distribution yielded the full-width half-maximum (FWHM) of 37 mV (Supplementary Fig. 14). Introducing 3-MPA enhanced the homogeneity of electronic properties 150 on SAM-modified surfaces, as reflected by a narrower CPD distribution with the FWHM of 22 151 152 mV.

To directly visualize the distribution of SAMs on FTO substrates, we performed high-angle 153 annular dark-field (HAADF) scanning transmission electron microscopy (STEM) measurements. 154 Fig. 2d shows cross-sectional HAADF-STEM images of both control and mixed samples. Because 155 of the contrast in the atomic number<sup>2</sup>, SAM was discernible as a dark layer sandwiched between 156 the brighter FTO and protective molybdenum oxide  $(MoO_x)$  layers. For the control SAM, we 157 158 observed thickness variations across the same FTO facet, with certain regions showing thicknesses 159 less than 1 nm. This suggests a low-density coverage and inhomogeneous distribution of 2PACz molecules<sup>32</sup>. The mixed sample exhibited significant improvements in terms of SAM uniformity 160 161 and coverage: A consistent thickness was recorded for the mixed SAM on the same facet, albeit with variations between 1 and 2 nm across different facets. Correspondingly, mixed SAMmodified FTO substrates exhibited higher hydrophobicity than control SAM-modified counterparts (Supplementary Fig. 15). Cyclic voltammetry (CV) measurements further determined the areal density of 2PACz in the mixed SAM to be  $3.9 \times 10^{13}$  molecules cm<sup>-2</sup>, a 70% increase from the  $2.3 \times 10^{13}$  molecules cm<sup>-2</sup> in the control SAM (Supplementary Fig. 16)<sup>16,27</sup>.

#### 167 Characterization of perovskite films

To investigate the impact of SAMs on the structural and optoelectronic properties of perovskites, 168 169 we fabricated triple-cation Cs0.05MA0.10FA0.85PbI3 perovskite thin films on different FTO/SAM substrates. From scanning electron microscopy (SEM), we observed no notable change in the 170 perovskite surface morphology when 3-MPA was incorporated into the SAM (Fig. 3a, b). Grazing 171 172 incidence wide-angle x-ray scattering (GIWAXS) measurements showed nearly identical crystal structures for perovskites deposited on both the control and mixed SAMs (Fig. 3c). Here, the main 173 constituent was photoactive  $\alpha$ -phase perovskites, with traces of the 4H hexagonal phase detected 174 at q = 0.82 Å<sup>-1</sup> and 0.92 Å<sup>-1</sup> as well as PbI<sub>2</sub> at q = 0.90 Å<sup>-1</sup>, respectively (Supplementary Fig. 14)<sup>33</sup>. 175 Time-of-flight secondary ion mass spectrometry (TOF-SIMS) further confirmed that both 3-MPA 176 and 2PACz molecules, due to their chemisorption on FTO surfaces, remained localized at the 177 perovskite/FTO interfaces (Fig. 3d, e). These findings suggest that 3-MPA is not likely to change 178 the perovskite formation process or contribute to the grain boundary or top surface passivation. 179

We conducted photoluminescence (PL) characterization by exciting perovskite thin films at a 1-180 sun-equivalent photon flux. We saw an average photoluminescence quantum yield (PLOY) of 0.13% 181  $\pm 0.02\%$  for perovskites directly deposited onto FTO substrates, corresponding to a quasi-Fermi 182 level splitting (QFLS) of 1.105 eV (Supplementary Table 1)<sup>34</sup>. 2PACz can passivate defects on 183 metal oxide surfaces and effectively block electron injection<sup>35</sup>, resulting in an approximately 50-184 fold increase in the PL intensity (Fig. 3f). We noted an average PLQY of  $7\% \pm 2\%$  for the 185 perovskite/2PACz/FTO stacks (Fig. 3g), corresponding to a QFLS of 1.207 eV. The incorporation 186 187 of 3-MPA further improved the average PLQY to  $10\% \pm 1\%$ . This we ascribed to better SAM coverage rather than 3-MPA passivation since the thiol group of 3-MPA was embedded within the 188 SAM (Fig. 1f). 189

We used ultraviolet and inverse photoelectron spectroscopy (UPS and IPES) to determine the workfunction and transport energy levels of perovskites and SAMs. The secondary electron cutoff in

192 the UPS spectra indicated an increase in the work function (WF) from 4.56 to 4.93 eV upon depositing the control SAM on the FTO substrate (Supplementary Fig. 18). 3-MPA led to an 193 194 additional WF shift of +100 meV. The ionization energy (IE) was 5.51 and 5.64 eV for the control and mixed SAMs, respectively. In contrast, perovskites exhibited similar conduction band 195 minimum (CBM) and valance band maximum (VBM) on different substrates (Fig. 3h). The 196 resultant energy level diagrams of perovskites and SAMs are depicted in Fig. 3i, j. A vacuum level 197 (VL) shift of 0.17 eV was recorded between the perovskite and the 2PACz bilayer, which further 198 increased to 0.32 eV for the perovskite/mixed-SAM bilayer<sup>36</sup>. A higher VL shift indicates an 199 enlarged built-in field<sup>37</sup>, which in PSCs can lead to enhanced carrier extraction<sup>38</sup>. Meanwhile, the 200 IE offset between the bilayer remained consistent for the control and mixed SAMs (referenced to 201 the Fermi level  $(E_F)$ ). 202

#### 203 Solar cell efficiency and stability

We fabricated PSCs having the inverted structure: FTO/SAM/perovskites/3,4,5-trifluoroanilinium 204 (345FAn)/fullerene (C<sub>60</sub>)/bathocuproine (BCP)/Ag (Fig. 4a and Supplementary Fig. 19). 345FAn 205 was selected for interface engineering due to its thermal stability<sup>1</sup>. 2PACz (control) and 2PACz:3-206 MPA (mixed) were employed as the SAMs for hole-selective contact. The corresponding solar 207 cell parameters are summarized in Fig. 4b. We found that compared to control devices, the mixed 208 SAM led to improved performance (average PCE 25.0% compared with 24.1%). This came from 209 open-circuit voltage (Voc, from 1.141 V to 1.159 V) and fill factor (FF, from 82.3% to 84.1%) 210 (Supplementary Note 4). We noted that device reproducibility was comparable for both control 211 and mixed SAMs, likely due to SAM inhomogeneity occurring at the nanoscale<sup>18,27</sup>. The champion 212 device with the mixed SAM exhibited a PCE of 25.3% from the reverse J-V scan, which agrees 213 with the PCE obtained from the steady-state power output (Supplementary Fig. 20). The integrated 214 short-circuit current density ( $J_{sc}$ ) of 25.8 mA cm<sup>-2</sup> derived from the external quantum efficiency 215 (EQE) measurement matches well with that from the *J*-*V* sweep (Fig. 4c). 216

217 One mixed-SAM-based FTO device was characterized at Newport (Montana, USA); and produced 218 a QSS-certified PCE of 24.8%, with a  $V_{oc}$  of 1.150 V, a  $J_{sc}$  of 25.5 mA cm<sup>-2</sup>, and an FF of 84.5% 219 (Fig. 4d and Supplementary Fig. 21). Although there have been reports of inverted PSCs with 220 efficiencies surpassing 25%<sup>2,3</sup>, the certified PCEs measured under stabilized conditions (including 221 maximum power point (MPP) and QSS tracking) have yet to reach the same level (Supplementary 222 Table 2). The OSS efficiency reported herein represents a new record for inverted PSCs, improving upon the previous record OSS efficiency of 24.09% in literature (Fig. 4e). 223

For comparison, inverted PSCs were fabricated on smooth ITO substrates. We obtained an average 224

PCE of 23.4% for control ITO devices, which was lower than that of control FTO devices, despite 225 226

- their higher average  $V_{oc}$  and FF (Supplementary Fig. 22). This emphasizes the need to augment  $J_{sc}$ for inverted PSCs. The limited  $J_{sc}$  is ascribed to the smoothness of ITO substrates (Supplementary
- Fig. 23), leading to reduced light-scattering and thus insufficient light absorption in the full device 228
- (Supplementary Fig. 24). Interestingly, the performance of mixed-SAM-based ITO devices was 229
- only slightly improved, possibly due to the weak interactions of 3-MPA with ITO surfaces 230
- 231 (Supplementary Note 5; Supplementary Fig. 25).

We evaluated the operating stability of PSCs using International Summit on Organic Photovoltaic 232 Stability (ISOS)-L-3 protocols, in which the encapsulated device was subjected to continuous 1-233 sun equivalent, white-LED illumination (Supplementary Fig. 26) at 50% relative humidity and 234 heatsink temperature of 65°C. To impede ion and moisture diffusion, we replaced BCP with 235 atomic-layer-deposited SnO<sub>2</sub> as a buffer layer<sup>9</sup>. The resultant device delivered a PCE of 24.6% at 236 room temperature (Supplementary Fig. 27). During the ISOS-L-3 testing, the device initially 237 demonstrated a PCE of 22.6%, which increased to 23.1% after 1.6 hours of MPP tracking (Fig. 4f). 238 The lower PCE at 65°C, compared to room temperature, is ascribed to the negative temperature 239 coefficient of PSCs (-0.15% /°C)<sup>39</sup>. The PCE stabilized at 21.9% (95% of the peak PCE) until the 240 end of the test (1075 h), with the main degradation in the photocurrent (Supplementary Fig. 28). 241 The initial performance and operating stability reported herein are compared with other PSCs 242 subjected to ISOS-L-3 tests (Supplementary Table 3). 243

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#### 331 Figure Legends

- Fig.1|MD simulations of phosphonic acid adsorption with and without molecular additives. 332 **a**. Chemical structures of the phosphonic acid 2PACz and the bifunctional compound 3-MPA. **b**. 333 Total number of 2PACz clusters formed over a set period, in the absence (control) and presence 334 (mixed) of 3-MPA. c, Types of 2PACz clusters formed at equilibrium. d, e, Top views of 335 equilibrated molecular representations of control (d) and mixed (e) systems. 2PACz and 3-MPA 336 (where applicable) are shown in pink and blue, respectively; Sn and O atoms, shown in the 337 background, are depicted in yellow and red, respectively. f, Successive steps along an AIMD 338 trajectory showcasing the role of 3-MPA as a co-adsorbent. Large, encircled areas are 339 magnifications of small ones. 340
- Fig.2|Homogeneity of self-assembled monolayers formed on FTO substrates. a, XPS P 2p (left) 341 and S 2p (right) spectra for the control and mixed samples. The peaks were fitted using one S 2p 342 or P 2p doublet with a 2:1 peak area ratio. **b**, 2PACz (left) and 3-MPA (right) transmission FTIR 343 spectra, compared with ATR-FTIR spectra for the SAM (left) and 3-MPA (right) coated FTO 344 substrates. The FTIR peaks identified at 947 and 1021 cm<sup>-1</sup> for 2PACz powders correspond to the 345 symmetric and asymmetric stretching of P–OH, respectively. Cz = carbazole. c, KPFM images of 346 control (left) and mixed (right) SAM-coated FTO substrates. d, Cross-sectional HAADF-STEM 347 images of control (top) and mixed (bottom) SAMs sandwiched between MoO<sub>x</sub> and FTO. 348
- Fig.3|Materials properties of perovskite films on different FTO/SAM substrates. a, b, Top-349 view SEM images of perovskite films deposited on control (a) and mixed (b) SAMs. c, GIWAXS 350 images for perovskite films on control (left) and mixed (right) SAMs. The color bar shows the 351 diffraction intensity collected from the GIWAXS detector.  $q_{xy}$  and  $q_z$  represent in-plane and near 352 out-of-plane scattering vectors, respectively. d, e, TOF-SIMS profiles of perovskites on 353 FTO/control SAM (d) and FTO/mixed SAM (e) substrates measured in negative polarity. Traces 354 of SH<sup>-</sup> were identified at the perovskite/control SAM interface, possibly due to contamination of 355 precursor materials. f, Absolute intensity PL spectra of perovskite films on bare FTO (no SAM) 356 357 and control and mixed SAMs. g, PLOYs of perovskite films on control and mixed SAMs (5 samples for each condition). Statistical distribution represented in box plots (line within the box: 358 mean, box limit: standard deviation). h, UPS and IPES spectra of perovskite films on control and 359 360 mixed SAMs. Gaussian fit (grey line) was used to determine the conduction and valence bands. i, j, Schematic energy level diagrams of the perovskite/control SAM (i) and perovskite/mixed SAM 361 bilayer (j), respectively. 362
- 363 Fig.4|Photovoltaic performance of perovskites solar cells. a, Schematic illustration of the device architecture with textured FTO substrate. **b**, Solar cell parameters for the control (16 devices) 364 and mixed (32 devices) SAM devices. Statistical distribution represented in box-and-whisker plots 365 (line within box: median, box limit: standard deviation, whiskers: 1.5 outliers). c, EOE and 366 367 integrated  $J_{sc}$  (grey dashed line) curves of the mixed SAM device. d, QSS J-V curve of one representative mixed SAM device certified at Newport. Inset: PV parameters of the device. e, 368 369 Certified performance of inverted PSCs measured under the QSS condition. f, MPP tracking of the encapsulated mixed SAM device at heatsink temperature of 65°C with a relative humidity of ~50% 370 under 1-sun illumination. The device configuration is FTO/SAM/Perovskite/345FAn/C<sub>60</sub>/ALD-371 SnO<sub>2</sub>/Ag, having a PCE of 24.6% at room temperature. 372

#### 373 Methods

#### 374 Molecular dynamics simulations

Static DFT calculation and AIMD were performed with the CP2K<sup>40</sup> package utilizing a mixed 375 Gaussian/plane wave (GPW) basis. The PBE exchange-correlation functional<sup>41</sup>, DFT-D3 376 dispersion corrections<sup>42</sup>, and dipole corrections<sup>43</sup> necessary for periodic boundary conditions along 377 the perpendicular direction to the surface were included. Valence electron wave functions were 378 379 expanded in a double- $\zeta$  Gaussian basis set with polarization functions (DZVP)<sup>44</sup>. The energy cutoff for the electron density expansion in the GPW method was 400 Ry. Born-Oppenheimer AIMD 380 simulations were run with an integration time step of 0.5 fs and the system was kept at 300 K using 381 the thermostat of Bussi et al.<sup>45</sup> in a canonical isothermal-isochoric (NVT) ensemble, where the 382 total number of atoms N, the volume V and the temperature T of the system were held constant. 383 All AIMD simulations were performed at  $\Gamma$  point. Being the most stable surface, the SnO<sub>2</sub> (110) 384 was chosen as the substrate surface. The systems were equilibrated over 5 ps in the NVT ensemble 385 and the remaining 40 ps were used for the production run. To investigate the interactions between 386 387 the 2PACz and 3-MPA molecules, as well as molecules and the SnO<sub>2</sub> substrate, 5 scenarios were simulated: (1) a single molecule (2PCAz or 3-MPA) on SnO<sub>2</sub> (110), (2) one 2PACz and one 3-388 389 MPA concurrently on SnO<sub>2</sub>(110), (3) two 2PACz and one 3-MPA on SnO<sub>2</sub>(110), (4) four 2PACz and one 3-MPA on  $SnO_2(110)$ , and (5) six 2PACz and one 3-MPA on  $SnO_2(110)$ . In all the 390 scenarios, an in-plane SnO<sub>2</sub> (110) simulation cell of 22.690 Å  $\times$  27.756 Å with 45 Å of vacuum 391 392 between the slab repetitions was used. AIMD was also used to estimate the interaction energy of 2PACz and 3-MPA with the SnO<sub>2</sub>(110) surface. The final AIMD configurations were relaxed with 393 DFT to extract the interaction energies of the molecules. Binding energies were calculated as 394  $\Delta E_{\rm b} = E_{\rm tot} - E_{\rm S} - E_{\rm mol}$ , where  $E_{\rm tot}$  is the energy of the final relaxed configuration (molecule on 395  $SnO_2$  (110));  $E_S$  is the energy of the final configuration of the slab;  $E_{mol}$  is the energy of the final 396 configuration of the isolated molecule. The interaction energies of 2PACz clusters were computed 397 398 after DFT relaxations of isolated clusters. Dimers, trimers, and tetramers were considered. Interaction energies per molecule were calculated as  $\Delta E_i = (E_{tot} - nE_{mol})/n$ , where  $E_{tot}$  is the energy 399 of the final relaxed cluster; n is the number of molecules forming the cluster;  $E_{\rm mol}$  is the energy of 400 401 a single relaxed molecule assumed isolated. Vibrational power spectra were computed as the Fourier transform of the autocorrelation function of the atomic velocities. 402

Classical MD simulations with full atomic resolution were realized with the help of the LAMMPS 403 MD simulator<sup>46</sup>. The equations of motion were integrated using the Velocity-Verlet method<sup>47</sup>, with 404 a 1 fs time step, while bond stretching was not constrained for any bonds. To maintain isothermal 405 conditions, the deterministic Nosé – Hoover thermostat<sup>48,49</sup>, with a time constant equal to 0.1 ps, 406 was utilized. This ensured that thermostating was applied to both translational and rotational 407 degrees of freedom, crucial for the asymmetric 2PACz and 3-MPA molecules. All classical MD 408 simulations were realized in the canonical NVT ensemble at T = 300K comprising a V-shaped 409 SnO<sub>2</sub> (110) model system with dimensions 60 Å  $\times$  105 Å  $\times$  40 Å. Periodic boundary conditions 410 were used in x- and y-directions, whereas reflective boundary conditions were employed in the z-411 direction. The two considered systems, control and mixed, comprised a total of 18498 and 18714 412 atoms, respectively, whereas the experimental molar ratio (2PACz/3-MPA=9/1) was retained in 413 414 the simulated mixed system. The systems were equilibrated for 1 ns followed by a 10ns production simulation, where sampling took place every 500 fs for the calculation of ensemble averages. A 415 snapshot of the system was taken every 1000 fs for visualization purposes. The calculation of the 416 number distributions of formed 2PACz clusters was performed through an in-house code utilizing 417

- a threshold distance of 8 Å among the centre of masses of adjacent 2PACz molecules in accordance
- 419 with the nearest neighbour shell of the corresponding radial distribution function as clustering
- 420 criterion.
- 421 To accurately capture the interactions of SnO<sub>2</sub>, the thoroughly validated classical force field
- developed by Bandura et al.<sup>50</sup> was used. The all-atom OPLS force field was utilized for 3-MPA,
   wherein the corresponding interaction parameters were obtained with the help of the LigParGen
- wherein the corresponding interaction parameters were obtained with the help of the LigParGen
   server<sup>51</sup>. For 2PACz, containing the phosphonic functional group, the specially designed force field
- 425 developed by Meltzer et al.<sup>52</sup> based on the Generalized Amber Force Field<sup>53</sup> was utilized. The van
- 426 der Waals and electrostatic non-bonded interactions were calculated using a real-space cutoff
- 427 radius of 10 Å and 8 Å, respectively, whereas the particle-particle-particle-mesh (PPPM) scheme<sup>54</sup>
- with an accuracy of 0.0001 (kcal/mol)/Å was used for the consideration of long-range electrostatic
- interactions. Because the  $SnO_2$  force field uses a Buckingham potential for the van der Waals interactions, its combination with the Lennard-Jones potential used for the compounds was
- 430 realized by fitting the Buckingham parameters of SnO<sub>2</sub> to a suitable set of Lennard-Jones
- 432 parameters<sup>55</sup> and geometric mixing rules were applied for all dissimilar non-bonded interactions.

#### 433 Materials

All materials were used as received without further purification. Organic halide salts, including 434 methylammonium iodide (MAI), formamidinium iodide (FAI), methylammonium chloride 435 (MACl), were purchased from Great Cell Solar. Cesium iodide (CsI), 3-mercaptopropionic acid 436 (3-MPA), 3,4,5-trifluoroaniline (98%), and guanidinium thiocyanate (GuaSCN) were purchased 437 438 from Sigma-Aldrich. 3,4,5-trifluoroaniline was converted to its ammonium form following the published procedure. [2-(9H-Carbazol-9-yl)ethyl]phosphonic Acid (2PACz), lead iodide (PbI<sub>2</sub>, 439 99.99%), and bathocuproine (BCP) were purchased from TCI. Anhydrous solvents including N, 440 N-dimethylformamide (DMF, 99.8%), dimethyl sulfoxide (DMSO, 99.9%), 2-propanol (IPA, 441 99.5%), chloroform (CF, 99.8%), and anisole (99.7%) were purchased from Sigma-Aldrich. C<sub>60</sub> 442 was purchased from Xi'an Polymer Light Technology Corp. ITO and FTO (TEC 10) substrates 443 were purchased from Thin Film Devices Inc. and Ossila, respectively. 444

445 **Perovskite film fabrication** 

1.5 M of Cs0.05MA0.1FA0.85PbI3 perovskite precursor was fully dissolved in mixed solvents of DMF 446 and DMSO (4:1, v/v) with the molar ratio for FAI/MAI/CsI was 0.85:0.1:0.05. 10 mg mL<sup>-1</sup> MAC1 447 448 and 8 mg mL<sup>-1</sup> GuaSCN were added in the solution to improve the film morphology. The precursor solution was filtered through a 0.22 µm polytetrafluoroethylene (PTFE) membrane before use. 60 449 µL of perovskite solution was deposited on the substrate and spun cast at 1000 rpm for 10 s 450 followed by 6000 rpm for 30 s. 150 µL anisole was dropped onto the substrate during the last 5 s 451 of the spinning, resulting in the formation of dark brown films that were then annealed on a hot 452 plate at 100°C for ITO and 120°C for FTO for 20 min. 453

#### 454 **Device fabrication**

FTO glasses were sequentially sonicated in aqueous detergent, deionized water, acetone, and IPA each for 10 min. After drying with nitrogen, the substrates were exposed to UV-ozone treatment for 15 min to remove organic contaminants and then transferred into a nitrogen-filled glovebox maintaining less than 0.1 ppm of  $O_2$  and  $H_2O$ . 100 µL of 2PACz or a mixture of 2PACz:3-MPA (in

a molar ratio of either 9:1 in characterization and device fabrication, or 8:2 in XPS measurements)

460 in anhydrous ethanol (1 mM) solution was uniformly spread on the substrates and allowed to rest for 15 s, followed by spinning the films at 3000 rpm for 30s. The films were then annealed at 461 100°C for 10 min and washed by dropping 500 µL ethanol during the film spinning at 3000 rpm. 462 463 Perovskites solution was deposited on the SAM-modified substrate as detailed above. 200 µL of 345FAn solution (1 mg/mL) in CF with an additional 3% of IPA was then drop cast within 2-3 s 464 on the perovskite film spinning at 4000 rpm (i.e., dynamic spinning) and annealed at 100°C for 5 465 min. Both control and mixed SAM films were then transferred to the thermal evaporator 466 (Angstrom engineering). C<sub>60</sub> (30 nm) and BCP (7 nm) were deposited sequentially with a rate of 467 0.3 Å s<sup>-1</sup> and 0.5 Å s<sup>-1</sup>, respectively, at a pressure of ca. 2 x 10<sup>-6</sup> mbar. Finally, Ag contact (140 468 nm) was deposited on top of BCP through a shadow mask with the desired aperture area. For the 469 stability testing, ALD-SnO<sub>2</sub> were used as the barrier layer. The deposition of ALD-SnO<sub>2</sub> was 470 carried out in the PICOSUN R-200 Advanced ALD system. H<sub>2</sub>O and TDMASn were used as 471 oxygen and tin precursors. Precursor and substrate temperature were set to 75°C and 85°C, 472 respectively. 90 SCCM N<sub>2</sub> was used as carrier gas. Pulse and purge times for H<sub>2</sub>O were 1 s and 5 473 s, and 1.6 s and 5 s for TDMASn. The total deposition cycle is 120, corresponding to 20 nm of 474 475 SnO<sub>2</sub>.

#### 476 Solar cell characterization

The current-voltage (I-V) characteristics of solar cells were measured using a Keithley 2400 477 sourcemeter under the illumination of solar simulator (Newport, Class AAA) at the light intensity 478 479 of 100 mW cm<sup>-2</sup> as checked with a calibrated reference solar cell (Newport). The stabilized PCE was measured by setting the bias voltage to the  $V_{MPP}$  and then tracing the current density. The  $V_{MPP}$ 480 was determined from the reverse I-V curve. The active area was determined by the aperture shade 481 mask (0.049 cm<sup>2</sup>) placed in front of the solar cell to avoid overestimation of the photocurrent 482 density. EQE spectra were recorded with a commercial system (Arkeo-Ariadne, Cicci Research 483 s.r.l.) based on a 300 W xenon light source and a holographic grating monochromator (Cornerstone, 484 Newport). 485

#### 486 Stability tests of solar cells

Devices were placed in a homemade stability tracking station. The illumination source is a white 487 light LED with intensity calibrated to match the 1-sun condition. For the ISOS-L-3 ageing protocol 488 (65 or 85°C; 50% relative humidity; MPP)<sup>56</sup>, the device chamber was left open in a room with 489  $50\pm10\%$  humidity and solar cell was mounted on a metal plate kept at  $65^{\circ}$ C by a heating element. 490 A thermal couple attached to the metal plate was used to monitor and provide feedback control to 491 the heating element to ensure temperature consistency. MPP was tracked using a home-build 492 493 MATLAB-based MPP tracking system using a 'perturb and observe' method. The MPP was updated every 1000 minutes. Encapsulation was done by capping the device with a glass slide, 494 using UV-adhesive (Lumtec LT-U001) as a sealant. 495

#### 496 **XPS measurements**

497 XPS measurements were performed with a Thermo Scientific K-Alpha system with  $180^{\circ}$  double-498 focusing, hemispherical analyser. The system is equipped with a 128-channel detector and 499 monochromatic small spot XPS. An Al K $\alpha$  source (1486.6 eV) was used for excitation and a pass 500 energy of 50 eV was used for XPS acquisition. Samples mounted on a metal specimen holder. All 501 data were analysed with CasaXPS and Thermo Avantage software.

#### 502 KPFM measurements

503 KPFM measurements were performed using an Asylum Research Cypher S atomic force 504 microscope (Oxford Instruments, Santa Barbara, USA) with an ASYELEC.01-R2 Ti-Ir coated 505 cantilever (Asylum Research). Scans were performed over 2 μm at 512 pixels and 0.5 Hz using a 506 two-pass method where the first pass is a tapping mode topography scan and the second in KPFM 507 mode with a tip potential of 3 V and a surface clearance of 5 nm. Cantilever calibration was 508 performed using the Thermal method from the Asylum Research GetReal database.

#### 509 HAADF-TEM measurements

HAADF-TEM images were acquired at an aberration corrected FEI (Titan Cubed Themis G2)
operated at 300 kV equipped with an XFEG gun and Bruker Super-X EDS detectors. The SAM
samples had a structure of glass/FTO/SAM/MoO<sub>x</sub> (50 nm), where the MoO<sub>x</sub> layer was deposited

through thermal evaporation with a low rate of  $0.1 \text{ Å s}^{-1}$  at a pressure below 2 x  $10^{-4}$  Pa. The cross-

- sectional samples were prepared by using a focused ion beam system (FIB, Helios G4 UX).
- 515 Another protective layer of carbon was thermally evaporated before ion-beam cutting and etching.

#### 516 CV measurements

CV measurements were conducted using a three-electrode configuration with a potentiostat 517 (PGSTAT204, Autolab). The working electrodes were prepared using a spin coating method on 518 an FTO electrode. The exposed area of the working electrode to the electrolyte measures 8 mm  $\times$ 519 520 21.4 mm. A platinum plate and an Ag/AgCl electrode (in a 3.0 M KCl solution) were employed as the counter and reference electrodes, respectively. The measurements were performed in an Ar-521 saturated solution of 1,2-dichlorobenzene (o-DCB) with 0.1 M tetrabutylammonium 522 hexafluorophosphate (TBA<sup>+</sup>PF $_{6}$ ) serving as the supporting electrolyte. All potentials are 523 referenced against the ferrocene redox couple, serving as an internal standard. The effective 524 coverage of the self-assembled monolayers on the FTO surface is measured by the slope of a linear 525 526 dependency of the oxidative peak intensity against the scan rate as follows:

527 
$$i_{p,o} = \frac{n^2 \mathrm{F}^2}{4 \mathrm{R} T \mathrm{N}_{\mathrm{A}}} A \Gamma^* v$$

528  $i_{p,o}$  is the oxidative peak current, v is the voltage scan rate, n is the number of electrons transferred, 529 F is the Faraday constant (96,485 C mol<sup>-1</sup>), R is the universal gas constant (8.314 J K<sup>-1</sup> mol<sup>-1</sup>), T

is the temperature, N<sub>A</sub> is the Avogadro constant, A is the electrode area, and  $\Gamma^*$  is the areal density.

#### 531 UPS and IPES measurements

UPS measurements were taken with an Excitech H Lyman- $\alpha$  photon source (10.2 eV) with a 532 nitrogen-filled beam path coupled with a PHI 5600 UHV and analyser system. A sample bias of -533 5 V was applied and a pass energy of 5.85 eV was used for UPS acquisition. IPES measurements 534 were performed in the Bremsstrahlung isochromatic mode with electron kinetic energies below 5 535 eV and an electron gun emission current of 2 µA was used to minimize sample damage. A Kimball 536 537 Physics ELG-2 electron gun with a BaO cathode was used to generate the electron beam. Emitted photons were collected with a bandpass photon detector consisting of an optical bandpass filter 538 (254 nm, Semrock) and a photomultiplier tube (R585, Hamamatsu Photonics). Samples were held 539 at a -20 V bias during all IPES measurements and the UHV chamber was completely dark. 540

#### 541 Other characterizations

542 Particle size distributions in solutions were determined by the DLS technique using a Malvern Zetasizer Nano ZS. Contact angles were measured using VCA-Optima XE. The image was taken 543 with CCD camera within 1 s after water droplet. GIWAXS measurements were performed at CMS 544 beamline, NSLS II. The monochromatic X-ray with the energy of 13.5 keV shone upon the 545 samples at different grazing incident angles of 1°, 0.5°, and 0.08° with an exposure time of 10 s. 546 547 A Pilatus800K detector was placed 259 mm away from the sample to capture the 2D diffraction 548 pattern. Absolute intensity photoluminescence spectra were measured using an integrating sphere, and Andor Kymera 193i spectrograph, and a 660 nm continuous-wave laser set at 1-sun equivalent 549 photon flux (1.1 µm beam full-width half-maximum, 632 µW); photoluminescence was collected 550 551 at normal incidence using a 0.1 NA, 110 µm-diameter optical fiber. TOF-SIMS was conducted on the IONTOF M6 instrument with a Bi<sup>3+</sup> (30 keV) primary ion beam for analysis and an Cs-cluster 552 gun (2 keV) for sputtering. Data was acquired in positive mode with an analysis area of  $49 \times 49$ 553  $\mu m^2$  centred and a raster area of 200 × 200  $\mu m^2$ . IR spectra were obtained in the attenuated total 554 reflectance mode using a Fourier-transform IR spectrometer (Thermo Scientific iS50). Samples 555 were prepared on the FTO substrate and scanned in the spectral range of 4000 to 550 cm<sup>-1</sup> with a 556 minimum number of 500 scans and a resolution of 4  $cm^{-1}$ . The triangular apodization function was 557 used to improve the signal-to-noise ratio. High-resolution SEM images were obtained using the 558 Hitachi S5200 microscope with an accelerating voltage of 1.5 kV. A low accelerating voltage and 559 a low beam current were deployed to reduce surface damage of perovskite films under electron 560 beam bombardment. The diffuse and specular light reflected from the substrate surface was 561 measured using a Cary 5000 UV-Vis-NIR double-beam spectrophotometer in diffuse reflectance 562 mode. Pure BaSO<sub>4</sub> was used for the baseline collection. 563

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#### 623 Author contributions

- 624 S.M.P., M.W., M.G., and E.H.S. conceived the idea and proposed the experimental and modelling
- design. N.L., L.A., V.C., and U.R. carried out the MD simulation. S.M.P. fabricated all the devices
- and conducted the characterization. T.H., H.R.A and K.R.G performed XPS, UPS, and IPES
- 627 characterization and data analysis. W.Y. and L.X. carried out the HAADF-STEM measurements.
- 628 F.T.E., M.W., S.M.Z., and M.G. conducted the PL and EQE characterization and data analysis.
- 629 M.W. measured DLS. H.S. conducted CV measurements and data analysis. D.C. performed UV-
- 630 Vis spectroscopy characterization. Y.Y. and M.G.K. measured TOF-SIMS. K.D. and A.A.
- 631 performed the GIWAXS measurements. M.V., E.D.J. and D.B.K. helped with the device
- fabrication and material characterization. P.S. and T.F. performed the KPFM measurements. M.W.,
- 633 S.M.P., N.L., M.G., and E.H.S. co-wrote the manuscript. All authors contributed to data analysis,
- read, and commented on the manuscript.

#### 635 Competing interests

636 The authors declare no competing interests.

#### 637 Data availability

- All data are available in the main text or the supplementary materials. Further data are available
- 639 from the corresponding author on reasonable request.

#### 640 Code availability

- 641 The codes and post-analysis tools for MD simulations are available in the following repository:
   642 <u>https://doi.org/10.5281/zenodo.8393081</u>.
- 643 **Correspondence and requests for materials** should be addressed to M.G. or E.H.S.





60 mV

30

0

-30

-60

Wavenumber (cm<sup>-1</sup>)

а





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Corresponding author(s): Michael Graetzel; Edward H. Sargent

### Solar Cells Reporting Summary

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#### Experimental design

#### Please check: are the following details reported in the manuscript?

1.	Dimensions	

	Area of the tested solar cells	Yes	0.049 cm2
	Area of the tested solar cens	No	
	Method used to determine the device area	Yes	The active area was determined by the aperture shade mask (0.049 cm2) placed in front of the solar cell
2.	Current-voltage characterization		
	Current density-voltage (J-V) plots in both forward	X Yes	Figure 4d (reverse and quasi-steady-state scans)
	and backward direction	No	
	Voltage scan conditions	X Yes	Supplementary Figure 21
	For instance: scan direction, speed, dwell times	No	
	Test environment	X Yes	Supplementary Figure 21
	For instance: characterization temperature, in air or in glove box	No	
	Protocol for preconditioning of the device before its	Yes	No preconditioning is needed
	characterization	🔀 No	
	Stability of the J-V characteristic	🔀 Yes	Supplementary Fig. 20b
	Verified with time evolution of the maximum power point or with the photocurrent at maximum power point; see ref. 7 for details.	No	
3.	Hysteresis or any other unusual behaviour		
	Description of the unusual behaviour observed during	Yes	PCE from reverse scan matches well with those from the quasi-steady-state scan and
	the characterization	🔀 No	steady-state measurements.
	Related experimental data	X Yes	Figure 4d and Supplementary Fig. 20
		No	
4.	Efficiency		
	External quantum efficiency (EQE) or incident	X Yes	Figure 4c
	photons to current efficiency (IPCE)	No	
	A comparison between the integrated response under	X Yes	The difference was found to be less than 1.5% (Main text - Solar cell efficiency and
	the standard reference spectrum and the response	No	stability)
			Net valavant
	For tandem solar cells, the bias illumination and bias voltage used for each subcell		
E	Calibration		
э.			Mathada Salar call characterization
	Light source and reference cell or sensor used for the characterization	No	
	Confirmation that the reference cell was calibrated	Yes	Methods - Solar cell characterization
	and certified	No	

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Calculation of spectral mismatch between the	X Yes
reference cell and the devices under test	No

6. Mask/aperture

Size of the mask/aperture used during testing

Variation of the measured short-circuit current density with the mask/aperture area

7. Performance certification

Identity of the independent certification laboratory that confirmed the photovoltaic performance

A copy of any certificate(s) Provide in Supplementary Information

8. Statistics

Number of solar cells tested

Statistical analysis of the device performance

9. Long-term stability analysis

Type of analysis, bias conditions and environmental conditions

For instance: illumination type, temperature, atmosphere humidity, encapsulation method, preconditioning temperature

No	
X Yes	0.049 cm2
No	
Yes	The accuracy of short-circuit current density was confirmed from EQE measurements.
📉 No	
X Yes	Newport
No	
X Yes	Supplementary Figure 21
No	
Yes No.	Figure 4b and Supplementary Figure 22
	Figure 4b and Supplementary Figure 22
X Yes	Figure 4f, Supplementary Figure 26 and Supplementary Figure 28
No	

M = 1.019 for the certified device