- 1 Strong Optical Absorption in AgBiS<sub>2</sub> Nanocrystals enabled by Cation
- 2 Disorder Engineering for Highly Efficient Extremely Thin Absorber Solar
- 3 Cells
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- Strong optical absorption of a semiconductor is a highly desirable property for a material 16 17 to be considered in optoelectronic and photovoltaic applications. Strong light absorbers can enable ultrathin solar cells and photodetectors that in turn lead to significant 18 reductions in cost, weight and manufacturing throughput as well as improve quantum 19 20 efficiency and performance. The optimal thickness of a semiconductor absorber is primarily determined by its absorption coefficient. To date, this parameter has been 21 22 considered as a fundamental material property and efforts to realize thinner photovoltaics have relied on light-trapping structures that add complexity and cost. Here, 23 24 we demonstrate that by engineering cation disorder homogeneity in a ternary 25 chalcogenide semiconductor leads to significant absorption increase due to enhancement 26 of the optical transition matrix elements. We show that cation disorder engineered AgBiS<sub>2</sub> 27 colloidal nanocrystals offer an absorption coefficient that is higher than any other 28 photovoltaic material used to date, enabling highly efficient extremely thin absorber

(ETA) photovoltaic devices. Leveraging this high absorption and by further optimization

of the electron and hole blocking layers we report solution-processed, environmentally-

friendly, 30nm thick ETA solar cells with short circuit current density of 27 mA·cm<sup>-2</sup>, a

# 1 record power conversion efficiency of 9.17% (8.85% certified) and high stability under

2 ambient conditions.

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High-performance, lightweight solar cells featuring low cost and environmental friendliness have been a long-sought target in the photovoltaic field. Ultrathin solar cells can reduce material consumption and manufacturing demands, directly lowering the cost. Owing to an absorber thickness that is orders of magnitude lower than conventional solar cells, ultrathin solar cells further allow for flexible and light-weight form factors suited for building- or wearable- integrated photovoltaics. A thinner absorber layer also improves charge carrier collection and reduces bulk recombination, which are particularly prevalent in nanocrystalline solution-processed semiconductors<sup>1,2</sup>, thus improving performance provided strong light absorption. Furthermore, high absorption in small absorber volumes allows for operating the device at high photo-generated carrier densities, a physical regime which maximizes the opencircuit voltage  $(V_{oc})$  and permits advanced high-efficiency concepts such as hot-carrier solar cells<sup>3</sup>. Currently, high performance ultrathin solar cells are mainly realized via different light-trapping strategies, which are employed to compensate for the short circuit current drop due to incomplete absorption in the ultrathin absorbing layer<sup>4,5</sup>. The use of optical architectures, however, increases non-radiative recombination<sup>4,6</sup> and complicates the manufacturing process, introducing additional obstacles toward low-cost high-efficiency devices. Typically, the absorption coefficient of a material determines the required thickness for optimal light harvesting under photovoltaic operation. The absorption coefficient of a semiconductor has been considered as a fundamental parameter of the material, with efforts to enhance light absorption in ultrathin absorbers relying on optical means<sup>4,5</sup>. We instead took the view motivated by the fundamental relationship between atomic geometry, electronic structure and

- 1 optical absorption that the absorption coefficient of a semiconductor can be tuned by
- 2 engineering the atomistic material structure.
- 3 Cation disorder is a widely observed phenomenon in multinary materials, referring to the
- 4 deviation of atomic positions in the cationic sublattice from an ordered crystalline arrangement.
- 5 This phenomenon significantly impacts the optoelectronic properties of semiconductors<sup>7–9</sup> and
- 6 has traditionally been considered an undesirable (and often unavoidable) effect, due to its
- 7 entropically-driven nature<sup>7,8,10,11</sup>. Herein, we leverage the modulation of cation disorder
- 8 homogeneity in multinary semiconductors as a pathway to enhancing optical transition matrix
- 9 elements, in order to achieve improved absorption coefficient and photovoltaic device
- 10 efficiency.

# Cation disorder homogeneity and absorption behaviour

- 12 In this work, we focus on ternary AgBiS<sub>2</sub> nanocrystals (NCs), a solution-processed
- 13 nanomaterial comprising environmentally-friendly elements<sup>12,13</sup> with reported efficiencies
- exceeding 6% in thin-film solar cells<sup>1,14</sup>. The cation distribution around Ag sites in AgBiS<sub>2</sub>
- nanocrystals has shown evidence for non-random Ag-Ag correlation<sup>15</sup>, indicating the presence
- of inhomogeneous cation disorder (i.e. cation segregation with local Ag-rich and Bi-rich
- 17 regions), likely due to growth kinetics during synthesis and surface ligand interactions <sup>16–18</sup>.
- An illustration of inhomogeneous cation disorder within AgBiS<sub>2</sub> NCs is shown in Fig. 1a. The
- valence band maximum (VBM) of AgBiS<sub>2</sub> primarily derives from Ag d and S p states, while
- 20 the conduction band minimum (CBM) arises from Bi p and S p interactions <sup>19,20</sup> (Fig. 1b). A
- clear spatial separation of the Ag-derived VBM and Bi-derived CBM for the cation-segregated
- configuration can be observed in the local electronic density of states (LDOS), shown in Fig.
- 1c. In contrast, under homogeneous cation disorder, we predict a VBM and CBM delocalized

- over the entire material (Fig. 1d), with the correlation between cation distribution and spatial
- 2 (de)localization of the band extrema further demonstrated by Fig. 1e.
- 3 The theoretically simulated absorption spectra of AgBiS<sub>2</sub> for both inhomogeneous and
- 4 homogenous cation disorder are provided in Fig. 2a. As expected, we find a substantially
- increased refractive index n and transition dipole matrix element (Supplementary Fig. 1), thus
- 6 enhanced optical absorption upon homogenizing disorder.
- 7 To experimentally tune cationic disorder, we first assessed the thermodynamics of atomic reordering, in particular the formation energy difference needed to induce disorder. Density 8 9 Functional Theory (DFT) was employed to calculate the inter-site cation exchange energy. Under the highest level of theory employed for phase energetics (hybrid DFT including spin-10 11 orbit coupling effects), the bulk order-disorder enthalpy difference is calculated as 17.4 12 meV/atom, indicating the accessibility of cation site-swapping in AgBiS<sub>2</sub> under mild annealing conditions. The mechanism behind cation inter-site exchange is likely defect-mediated ionic 13 migration. Indeed, cation mobility aided by Ag vacancies and bond anharmonicity has been 14 demonstrated in AgBiS<sub>2</sub> nanocrystals<sup>21–23</sup>. Considering the low inter-site cation exchange 15 energy in AgBiS2, we sought to anneal AgBiS2 NCs under low-temperature conditions to 16 facilitate cation inter-site exchange and thus an entropically-driven transition to homogenous 17 cation distribution. Fig. 2b plots the absorption coefficient of our NC films upon annealing 18 19 under different temperatures. Comparing Fig. 2a and 2b, the simulated absorption spectrum for cation segregation configurations (inhomogeneous disorder) matches well with the as-prepared 20 AgBiS<sub>2</sub> NC film, both exhibiting Urbach tailing at longer wavelengths, which is pernicious for 21 photovoltaic devices<sup>24,25</sup>. However, in the homogenous cation-disordered structure, the 22 simulated absorption coefficient is much higher than the cation-segregated case, with reduced 23 bandtailing, manifesting favourable properties for optoelectronic applications<sup>26</sup>. As shown in 24 Fig. 2b, the absorption coefficient of AgBiS<sub>2</sub> NCs films is enhanced by up to a factor of two 25

1 after annealing, alongside reduced Urbach energy from 173.2 meV to 25.7 meV

2 (Supplementary Fig. 2), as predicted by ab initio calculations. Furthermore, this enhanced

absorption is stable in ambient atmosphere, as spontaneous cation segregation is

thermodynamically and entropically unfavourable (Supplementary Fig. 3). With this low-

energy and scalable annealing process, we produce a semiconducting material that exhibits an

absorption coefficient 5-10 times greater than any other material currently used in photovoltaic

technology<sup>4,27–31</sup>, across a wide spectral range from 400 – 1000 nm (Fig. 2c).

With the high absorption coefficient of our films, an ultrathin layer of AgBiS<sub>2</sub> NCs would be sufficient to absorb most of the light in the visible range. In order to assess the potential of our strongly-absorbing NC films, we calculate the maximum achievable short-circuit current density  $J_{sc}$  using the transfer matrix method<sup>32,33</sup>, assuming 100% internal quantum efficiency (Supplementary Fig. 4). Fig. 2d plots the maximum  $J_{\rm sc}$  as a function of the active layer thickness for different annealing temperatures, corresponding to varying degrees of cation disorder homogeneity. Upon increased annealing temperature, we witness an increase in  $J_{\rm sc}$ reached at low active layer thicknesses (t < 200 nm). We predict a maximum  $J_{sc}$  of ~28 (32)  $mA \cdot cm^{-2}$  for AgBiS<sub>2</sub> NCs films annealed at 115°C (150°C), with thicknesses of only ~30 nm. Furthermore, the Spectroscopic Limited Maximum Efficiency (SLME) was calculated assuming only radiative recombination in devices<sup>34</sup>. As shown in Fig. 2e, a high photovoltaic efficiency up to 26% was predicted for a 30 nm absorber layer, indicating the performance potential of ultrathin solar cells based on AgBiS<sub>2</sub> NC films. Furthermore, with the thickness reduction of AgBiS<sub>2</sub> absorber from 500 nm (common photovoltaic thickness) to 30 nm, we estimate a ~50% cost reduction over the full PV device cost (Supplementary Fig.5 and Supplementary Note 1).

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# Cation configuration transition

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2 To further verify the proposed cation homogenization as the underlying mechanism responsible for the optical absorption enhancement in AgBiS2 NC films, film thicknesses were first 3 measured to exclude nanocrystal densification as the dominant factor. Negligible thickness 4 5 changes were observed between the as-prepared and 115 °C annealed samples (Supplementary Fig. 6), with only a slight decrease for the 150 °C annealed films, contributing to a very small 6 7 (~8%) absorption enhancement for this sample. These film thickness measurements demonstrate that densification effect is not the dominant factor in the absorption enhancement 8 9 of these materials. Furthermore, we used X-Ray Diffraction (XRD) and Transmission Electron 10 Microscopy (TEM) to probe the changes in crystal structure due to cation disorder homogenization upon annealing. Interestingly, while XRD analysis suggested increased 11 12 crystallite size (from 4.2 nm to 6.2 nm) upon annealing at 115°C, High Resolution Transmission Electron Microscopy (HRTEM) showed no measurable size differences in these 13 samples (Supplementary Fig. 7). The sharpening of XRD peaks despite negligible NC growth 14 indicates an improved crystallinity due to atomic rearrangements within the nanocrystals. 15 Further increasing the annealing temperature to 150°C and 200°C causes NCs to fuse to larger 16 17 crystals (28 nm and 47 nm, Supplementary Fig. 7). Point defects are observable from HRTEM after high temperature annealing, which could be harmful for optoelectronic applications<sup>35</sup> 18 19 (Supplementary Fig. 8). In addition to peak narrowing in XRD patterns, the peak positions 20 were also found to shift to higher angles upon annealing (Fig. 3a). To deconvolute the effect of crystal size and cation arrangement, and explain the apparent changes in crystallinity, we 21 22 calculated the expected XRD patterns for homogeneously disordered cubic AgBiS<sub>2</sub> (space group  $Fm\bar{3}m$ , Supplementary Fig. 9), with crystal size as the only variable parameter 23 (Supplementary Fig. 10). The full width half maximum (FWHM) of all peaks sharpens with 24 25 crystal growth, while the peak positions remain essentially the same, as expected. However,

1 transitioning from cation-segregated configurations to homogeneous cation disorder, while 2 fixing the crystallite size, the simulated XRD patterns of AgBiS<sub>2</sub> show distinct peak shifts to higher angles as well as peak narrowing (Fig. 3b), matching the experimental observations of 3 4 improved crystallinity upon annealing. The XRD peak shifts primarily originate from the shortening of Ag-S bond lengths, while peak narrowing occurs due to narrowing of the bond 5 6 length distributions and reduction of octahedral distortion, upon homogenization of the cation 7 distribution (Supplementary Fig. 11). This phenomenon was further confirmed with HRTEM measurements, as shown in Fig. 3c and d, with integrated line profiles of the {200}-plane 8 9 showing a slight shrinkage after annealing, further confirming the transition from cation segregation to homogeneous disorder. 10 Considering the difference in local bonding environments for different cation configurations, 11 changes in Madelung potential were expected<sup>36</sup>. We calculate the average Madelung potential 12 at Bi sites as 3.29 V and 4.66 V for segregated and homogeneous cation configurations, 13 respectively, using Bader atomic charges. A greater Madelung potential suggests a decrease in 14 XPS binding energy, upon transitioning from inhomogeneous to homogeneous cation 15 disorder $^{36,37}$ . As shown in Fig. 3e, the Bi 5d peaks in the simulated XPS spectra are noticeably 16 17 shifted to lower binding energies for homogeneous cation disorder, as compared to cationsegregated configurations. Likewise, the annealed AgBiS<sub>2</sub> NCs show a small but significant 18 19 chemical shift to lower energy, compared with as-prepared samples (Fig. 3f and g), in 20 agreement with our proposition of cation homogenization upon annealing.

# Ultrathin AgBiS<sub>2</sub> NC solar cells

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Having developed a material with such strong optical absorption, we next sought to produce highly efficient ultrathin solution-processed solar cells with these nanocrystals. Solar cells were fabricated with an architecture of Glass/ITO/SnO<sub>2</sub>/AgBiS<sub>2</sub>/HTL/MoO<sub>3</sub>/Ag, as shown in Fig.

1 4a. Cross-sectional Transmission Electron Microscopy (TEM) confirms the ultrathin nature of 2 the device layers (Fig. 4b). We first used PTB7 as an electron blocking layer, in accordance with previous studies<sup>1,14</sup>. The devices showed an average power conversion efficiency (PCE) 3 of 6.4  $\pm$  0.6%, with a champion device reaching PCE of 7.6% (Table 1) – higher than the 4 previously reported record performance of 6.3% 1,14,38-40. Atomic Force Microscopy (AFM) 5 6 revealed a surface roughness of 0.6 nm for a 4 nm PTB7 film (Supplementary Fig. 12), which undermines the performance of the cells by introducing interface recombination<sup>41</sup>. We sought 7 to replace PTB7 with an alternative electron-blocking layer with improved morphological 8 9 characteristics. We found that PTAA yielded improved uniformity (RMS roughness of 0.4 nm), a prerequisite for suppressing current leakage and interface recombination. In order to further 10 assess the superiority of PTAA, transient photocurrent and photo-voltage were measured under 11 12 one-Sun light bias (Supplementary Fig. 13). Devices with PTAA showed a faster photocurrent decay and longer carrier lifetime than that with PTB7 as the HTL, which indicates improved 13 charge extraction and reduced interface recombination. As a result, replacing PTB7 with PTAA 14 15 leads to significant improvement in  $V_{oc}$  and fill factor (FF) (Supplementary Fig. 12), and results in a ~20% increase in power conversion efficiency to 8.7  $\pm$  0.3%, with a best device reaching 16 9.17% (Fig. 4c and d). One of our champion devices was sent to an accredited PV calibration 17 laboratory (Newport, USA), which certified a PCE of 8.85% under AM1.5G full sun 18 illumination, with negligible hysteresis (Fig. 4e, Supplementary Fig. 14). The measured  $J_{\rm sc}$  of 19 27 mA·cm<sup>-2</sup> was further confirmed by the external quantum efficiency (EQE) spectrum, which 20 gives a value of 26.5 mA·cm<sup>-2</sup> (Fig. 4f). In addition, the measured EQE spectrum matched well 21 with the TMM predicted EQE, indicating a near-unity internal quantum efficiency (IQE). 22 23 However, unannealed devices showed not only lower absorption (i.e. predicted EQE) in near infrared region, but also a lower IQE (Fig. 4f) and larger  $V_{oc}$  deficit, which is ascribed to slower 24 charge extraction, Urbach band tailing and higher trap density (Supplementary Fig. 15). Higher 25

temperature annealing leads to significantly lower performance, which is consistent with the defects forming from uncontrolled nanocrystal fusing (Supplementary Fig. 8 and Fig. 16).

Analysing the absorption loss of our record performance ultrathin devices, a large part of the incident light (~10 mA·cm<sup>-2</sup>) was reflected (Supplementary Fig. 17) and further improvement

of device performance is expected by introducing an antireflection layer.

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- On the other hand, thicker (200 nm) devices are also fabricated for direct comparison with 6 ultrathin devices and, as expected, a low  $J_{sc}$  (~15 mA·cm<sup>-2</sup>) was obtained (Supplementary Fig. 7 18). In order to further understand the devices, time-of-flight (ToF) was employed to 8 9 investigate the charge carrier mobilities in our AgBiS<sub>2</sub> NC films (Supplementary Fig. 19). The hole and electron mobility are calculated to be  $1.2 \times 10^{-4}$  and  $5.7 \times 10^{-5}$  cm<sup>2</sup>·V<sup>-1</sup>·s<sup>-1</sup>, respectively. 10 Combining with the carrier lifetime extracted from TPV measurements, the diffusion length is 11 12 estimated to be ~25.4 nm, which rationalizes the near-unity IQE in our 30 nm devices and the low performance in 200 nm thick devices. Mid-gap trap density was estimated from TPC/TPV 13 measurements (Supplementary Fig. 15). In our optimal AgBiS<sub>2</sub> NC films, the trap density is ~ 14  $9\times10^{16}$  cm<sup>-3</sup>. Although this trap density is four times lower than before annealing (~  $4\times10^{17}$  cm<sup>-3</sup>) 15  $^{3}$ ), efforts are needed to further passivate the trap states for higher  $V_{\rm oc}$  and thus, higher 16 17 efficiency.
  - In addition to power conversion efficiency, stability is another important figure-of-merit of photovoltaic devices. Therefore, in order to assess the stability of our AgBiS<sub>2</sub> NC solar cells, we first measured their shelf lifetime by storing the un-encapsulated devices in ambient conditions with a temperature of ~25°C and relative humidity of ~30%. The device showed a gradual improvement in the first 20 days (Supplementary Fig. 20), which is likely a result of oxidation of MoO<sub>x</sub> layer and a better band alignment<sup>44</sup> (Supplementary Fig. 21). Furthermore, the device retained its original performance after four months' aging. The operational stability was further investigated by subjecting the un-encapsulated device to AM1.5G one sun

1 illumination in ambient atmosphere with relative humidity ~ 60%. The device performance 2 was measured by applying a forward bias fixed at the maximum power point (MPP). As shown in Fig. 4g, the PCE of the PTB7 device dropped below 2% after 20 minutes illumination, while 3 4 the PTAA device demonstrated much better operational stability under continuous operation. After 10 hours MPP testing on un-encapsulated cell in ambient conditions, the device retained 5 85% of its original efficiency. Longer stability at MPP was also recorded with a class AA solar 6 light source, showing ~80% of original performance after 40h test in ambient air 7 (Supplementary Fig. 22). To our knowledge the devices reported herein set a record among 8 9 low-temperature and solution-processed, non-toxic inorganic solar cells in terms of stability and performance<sup>45–48</sup>. These results support that AgBiS<sub>2</sub> NCs is extremely promising material 10 for low-cost, efficient, stable and environmentally friendly solar cells. 11 12 In conclusion, we have demonstrated that the absorption coefficients of ternary AgBiS<sub>2</sub> NCs can be enhanced via cation disorder homogenization at mild annealing conditions. Ultra-high 13 absorption coefficients were obtained in annealed AgBiS<sub>2</sub> NC films, with a calculated SLME 14 of over 26% for a 30 nm NC film. The transition in cation configuration was further confirmed 15 by the combination of ab initio calculations with XRD, HRTEM and XPS measurements. 16 17 Ultrathin solar cells are fabricated based on ultra-absorbing AgBiS<sub>2</sub> NCs. A high  $J_{\rm sc}$  of 27 mA·cm<sup>-2</sup> and a record efficiency up to 9.17% were obtained with an independent certification 18 19 of 8.85% from Newport. The air stability and photostability was also recorded in high 20 performance devices. Our work not only establishes the potential of ultrathin AgBiS<sub>2</sub> NC solar cells, which are solution-processable and RoHS-compliant, but also demonstrates the 21 importance of atomic configuration engineering in multinary systems. 22

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#### Methods

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Density Functional Theory (DFT) simulations. The AiiDA infrastructure was used to 2 maintain data provenance for all calculations performed in this study<sup>49</sup>. Calculations were 3 performed using Density Functional Theory (DFT) within periodic boundary conditions 4 through the Vienna *Ab Initio* Simulation Package (VASP)<sup>50–52</sup>. Using the projector-augmented 5 wave method, scalar-relativistic potentials were employed to describe the interaction between 6 core and valence electrons.<sup>53</sup> Calculations were carried out using  $\Gamma$ -centred k-point meshes 7 with a reciprocal space sampling of 0.11 Å<sup>-1</sup> and a plane-wave kinetic energy cutoff of 300 eV 8 9 (increased to 450 eV for structural relaxations, to avoid the possibility of Pulay stress). With these parameter choices, the ground-state energies of all known structures in the Ag-Bi-S 10 system were converged to within 1 meV/atom. 11 12 Cation Disorder. To model the effects of varying degrees of cation order/disorder, two approaches were employed. Firstly, to simulate total random cation disorder, a 64-atom 13 supercell for AgBiS<sub>2</sub> in the  $Fm\bar{3}m$  rocksalt structure was generated using the special quasi-14 random structure (SQS) method<sup>54</sup>, in which the cation-cation cluster correlations are optimised 15 to obtain the best approximation to an ideal infinite random distribution for a given supercell. 16 Here, the Alloy Theoretic Automated Toolkit (ATAT)<sup>55</sup> was used to generate the SQS supercell 17 via Monte-Carlo simulated annealing loops<sup>56</sup>. Eight Monte-Carlo simulations were performed, 18 19 with the structure giving the best match to the true disordered solid solution chosen for further calculations. This corresponded to an 'objective function' of -81.18; considering 2-atom 20 clusters up to 12 Å separation, 3-atom clusters up to maximum 10 Å pair separation and 4-21 atom clusters up to 8 Å pair separation (using the experimental rocksalt crystal structure)<sup>37</sup>. 22 The screened hybrid DFT exchange-correlation functional of Heyd, Scuseria and Ernzerhof 23  $(HSE06)^{38}$  was used to calculate the structural and electronic properties of  $Fm\overline{3}m$  (using this 24 SQS structure) and  $R\bar{3}m$  AgBiS<sub>2</sub>, unless otherwise stated, having been demonstrated to 25

accurately predict the electronic structures of semiconductor materials<sup>59,60</sup>. To account for 1 2 relativistic effects, due to the presence of heavy-atom elements Ag and Bi, spin-orbit coupling effects were included in all electronic and optical calculations (HSE06+SOC). 3 4 Secondly, to investigate the effects of both partial order and inhomogeneous disorder (i.e. cation segregation with Ag-rich and Bi-rich regions) on the cation sublattice, all possible 5 6 configurations for AgBiS<sub>2</sub> in a 32-atom rocksalt structure were enumerated using the Transformer package<sup>61</sup>, yielding 440 symmetry-inequivalent arrangements. Here, the PBEsol<sup>42</sup> 7 semi-local exchange-correlation functional was used for geometry optimisations and energetic 8 analysis, motivated by its well-established accuracy for the structural relaxation of bulk solids<sup>43</sup> 9 and moderate computational cost. Moreover, this functional was tested against both experiment 10 and the computationally-intensive HSE06 hybrid DFT functional for bulk geometry 11 12 optimisation and energetics respectively, using all known structures in the Ag-Bi-S system. Here, PBEsol was found to accurately reproduce experimentally-reported crystal structures, 13 with a mean absolute error <1.5% in the lattice parameters, as well as reproducing the energetic 14 15 ordering of phases on the convex hull according to both HSE06 and experiment. To ensure consistency when comparing the results of these calculations to that of  $Fm\bar{3}m$  (SQS) 16 17 AgBiS<sub>2</sub>, a new 32-atom SQS supercell was generated using the same procedure as outlined above, and relaxed with PBEsol. In this case, a perfect 'objective function' match to the true 18 disordered solution was obtained. While appropriate for structural and energetic analysis of 19 cation configurations in AgBiS<sub>2</sub>, semi-local DFT is known to severely underestimate electronic 20 bandgaps<sup>64,65</sup>, and so the HSE06+SOC hybrid DFT functional was used to calculate the optical 21 22 and electronic properties of these configurations. In this case, downsampling of the Fock exchange matrix by a factor of 2 (NKRED = 2) was employed to reduce the computational cost 23 to a manageable level. This choice was confirmed to affect the electronic bandgap by <0.05 eV 24 25 for the 32-atom SQS supercell.

- 1 To investigate the impact of supercell size on the calculated properties, the 32-atom SQS
- 2 supercell was also relaxed with the HSE06 functional, to then compare with the 64-atom
- 3 structure. The total energy was found to match that of the 64-atom SQS supercell to <1
- 4 meV/atom, while the bandgap was found to increase slightly from 0.69 eV to 0.83 eV for the
- 5 32-atom supercell.
- 6 Post-processing. Primitive and unfolded electronic band structure diagrams were generated
- vsing sumo and PyVaspWfc<sup>46,47</sup> respectively. Effmass was used to calculate carrier effective
- 8 masses<sup>68</sup>, and photoemission spectra were generated using Galore<sup>69</sup>. COHP analysis and
- 9 charge-density partitioning was performed using LOBSTER,<sup>50</sup> and the vasppy package<sup>51</sup> was
- used to calculate radial distribution functions. The pymatgen package was used throughout for
- manipulation and analysis of calculation inputs and outputs<sup>72</sup>.
- 12 Chemicals and materials. Reagents were purchased from Sigma Aldrich, except SnO<sub>2</sub> colloid
- precursor (tin (IV) oxide, 15% in H<sub>2</sub>O colloidal dispersion), which was obtained from Alfa
- Aesar. Poly (triaryl amine) (PTAA) was purchased from EM index and Poly [[4,8-bis](2-
- ethylhexyl)oxy]benzo[1,2-b:4,5-b']dithiophene-2,6-diyl][3-fluoro-2-[(2-ethylhexyl) carbonyl]
- thieno [3,4-b]thiophenediyl]] (PTB7) was purchased from 1-materials.
- 17 Synthesis of AgBiS<sub>2</sub> nanocrystals. The Schlenk technique was used to synthesize AgBiS<sub>2</sub>
- nanocrystals, following previous report with modifications<sup>1,14</sup>. 4 mmol Bi(OAc)<sub>3</sub>, 3.2 mmol
- 19 Ag(OAc), 24 mL oleic acid (OA) and 15 mL 1-octadecene (ODE) was pumped at 100°C for 2
- 20 hours (~0.2 mbar) to remove oxygen and moisture. 4 mmol hexamethyldisilathiane (HMS)
- 21 dissolved in 5 mL ODE was quickly injected into the flask. The heating mantel was removed
- 22 and the reaction was cooled down with water bath for ~5 min. After that, the crude solution
- 23 was left stirring for 1 hour at room temperature. The nanocrystals were isolated by adding
- acetone, followed by centrifugation. The extracted nanocrystals were re-dispersed in toluene
- and acetone was added to precipitate nanocrystals. The re-dispersion/precipitation was

- 1 repeated once more. Finally, the obtained AgBiS<sub>2</sub> nanocrystals powder was dispersed in
- 2 anhydrous toluene (20 mg mL<sup>-1</sup>) and stored in ambient atmosphere for device fabrication.
- 3 Characterization of AgBiS2 nanocrystals and films. Films exchanged with 3-
- 4 mercaptopropionic acid (3-MPA) were grown on glass substrates, followed by annealing at
- 5 different temperatures in glovebox. The thicknesses were measured using a profilometer. The
- 6 complex refractive index was measured at various angles using broadband Sopra Ellipsometer
- 7 GES5E. The software SEMILAB Spectroscopic Ellipsometry Analyzer was utilized to fit a
- 8 model of stacked layers of appropriate optical constants and the thickness from profilometer
- 9 was used as an input. The absorption coefficients are calculated from extinction coefficients.
- 10 XRD data were collected using a Rigaku Smartlab powder diffractometer in Bragg-Brentano
- 11 geometry with Cu Kα radiation. XPS measurements was performed with a SPECS PHOIBOS
- 12 150 hemispherical analyser (SPECS GmbH, Berlin, Germany) in ultra-high vacuum conditions
- 13 (10<sup>-10</sup> mbar), with a monochromatic  $K\alpha$  x-ray source (1486.74 eV) at the Institut Catala de
- Nanociencia i Nanotecnologia (ICN2). TEM was performed at the Scientific and Technological
- 15 Centres of the University of Barcelona (CCiT-UB). TEM images were obtained using a JEOL
- 2100 microscope operating at an accelerating voltage of 200 kV.
- 17 **Optical modelling and Spectroscopic limited maximum efficiency.** A homemade MATLAB
- code was used to implement the transfer matrix formalism. We assumed that each layer was
- 19 flat and considered no scattering effects. Short circuit current density was calculated with the
- 20 assumption of 100% internal quantum efficiency. Spectroscopic Limited Maximum Efficiency
- 21 (SLME) was calculated according to:

$$SLME = P_{max}$$

$$P = -V * \left\{ J_0 \left[ \exp\left(\frac{eV}{kT}\right) - 1 \right] - J_{SC,TMM} \right\}$$

- where e the elementary charge, k the Boltzmann constant, T the temperature, and  $J_0$  the dark
- 2 saturation current density, calculated with the assumption of only radiative recombination:

$$J_0 = e * \int_0^{+\infty} EQE_{TMM} * \emptyset_{BB}(E) dE$$

- 4 where EQE<sub>TMM</sub> is the calculated total absorption in AgBiS<sub>2</sub> NCs layer, and  $\emptyset_{BB}$  is the black
- 5 body radiation spectra at 300 K.
- 6 Solar cell fabrication. All solar cell fabrication steps were performed in ambient air, unless
- 7 with specific descriptions. ITO covered glass substrates (Universität Stuttgart, Institut für
- 8 Großflächige Mikroelektronik) were cleaned by ultra sonication in soapy water, acetone and
- 9 isopropanol for 20 min each and dried with nitrogen, followed with 0.5 h UV/Ozone treatment.
- SnO<sub>2</sub> electron transport layer was then spin cast from diluted Alfa SnO<sub>2</sub> colloid solution (1:5.6
- 11 v/v with H<sub>2</sub>O) with spin speed of 2000 rpm and annealed at 270°C for 15 min. Afterwards,
- three layers of AgBiS<sub>2</sub> nanocrystals were deposited from 20 mg mL<sup>-1</sup> toluene solution via layer-
- by-layer (LbL) method. For each AgBiS<sub>2</sub> nanocrystals layer, one drop of AgBiS<sub>2</sub> NCs solution
- was spin-coated onto SnO<sub>2</sub>/ITO substrates during spinning (2000 rpm). 3-Mercaptopropionic
- acid (MPA)/methanol (1% v/v) solution was then applied to the nanocrystals film for 45s,
- 16 followed by two rinse-spin steps with methanol and once with toluene. The films were
- transferred into glovebox for 10 min annealing at 115°C and then stored in dry air before spin-
- coating PTB7 (5 mg mL<sup>-1</sup> in dichlorobenzene) or PTAA solution (2 mg mL<sup>-1</sup> in toluene) at
- 19 3000 rpm. Finally, a Kurt J. Lesker Nano36 system was used to deposit 3 nm of MoO<sub>3</sub> and 120
- 20 nm of Ag through a shadow mask to produce solar cells with a diameter of 2 mm (area of 3.14
- 21  $\text{mm}^2$ ).
- 22 Solar cell characterization. All device characterization was performed in air under ambient
- conditions. Current-voltage measurements were performed with a Keithyley 2400 Sourcemeter

1	and a Newport Oriel Sol3A solar simulator with an AM1.5G filter. The intensity of the solar
2	light was calibrated using a Hamamatsu S1336 silicon photodiode that had been calibrated at
3	the Fraunhofer Institute of Solar Energy Systems, Freiburg, Germany. The solar cells were
4	measured with and without masks, and slightly lower $V_{ m oc}$ was observed when measuring with
5	masks, due to masking effect. For certified cells, appropriate masks have always been used.
6	The EQE was measured using a Newport Cornerstone 260 monochrometer, a Thorlabs
7	MC2000 chopper, a Stanford Research SR570 trans-impedance amplifier and a Stanford
8	Research SR830 lock-in amplifier. A calibrated Newport 818-UV photodetector was used as a
9	reference. Shelf stability was obtained from devices stored in air without encapsulation. For
10	the maximum power point (MPP) measurement, the MPP voltage ( $t = 0$ ) was measured before
11	MPP testing. The device was then held at the MPP voltage $(t = 0)$ for operating stability test.
12	The device was unencapsulated under AM1.5G illumination. All devices were characterized
13	under ambient condition with relative humidity > 60% and ambient temperature ~25°C. Longer
14	time photo-stability was measured with a low-cost solar simulator from ABET technologies.
15	Transient photocurrent and photovoltage of the devices were measured with an in-house-built
16	set-up. The set-up comprises a LED lamp to provide steady state white light bias, a 637nm
17	laser and an Agilent 4000X oscilloscope. The Voc of devices are controlled by the light
18	intensity of the LED lamp and the transient voltage induced by laser pulse are controlled within
19	5% of the Voc. Time-of-flight method is used for the mobility measurements. Electron only
20	and hole only devices are fabricated with the structure of ITO/SnO $_2$ /AgBiS $_2$ /PCBM/Ag and
21	$ITO/NiO_x/AgBiS_2/PTAA/MoO_x/Ag. \ The \ thickness \ of \ AgBiS_2 \ layer \ was \ controlled \ to \ \sim 200 \ nm. \ controlled \ control$
22	A 520 nm nanosecond laser was used as the excitation light. Photocurrent decay was recorded
23	at various bias with an oscilloscope. The mobility was calculated by fitting the $d^2/t$ - $V_{bias}$ plots.

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#### **Author contributions**

G.K. supervised and directed the study. Y.W. and G.K. conceived the idea, designed this study and co-wrote the manuscript with feedback from co-authors. Y.W. synthesized the AgBiS<sub>2</sub>

- 1 NCs, performed material characterization, fabricated, characterized the solar cells, and
- analysed the data, with the help of I.B. Y.W. did the optical modelling. S.K. designed and
- 3 conducted the theoretical modelling, analysed the DFT simulations, interpreted the data,
- 4 provided insights and contributed to manuscript writing. D.O.S and A.W. supervised the
- 5 theoretical modelling.

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#### Additional information

- 8 Supplementary information is available in the online version of the paper. Correspondence and
- 9 request materials should be addressed to G.K.

# 10 Data and code availability

- 11 The data and code that supported the current study are available from the corresponding
- author on request.

#### 13 Competing financial interests

14 The authors declare no competing financial interests.

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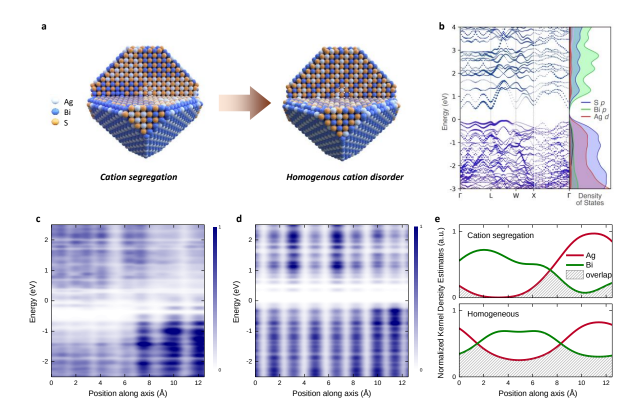
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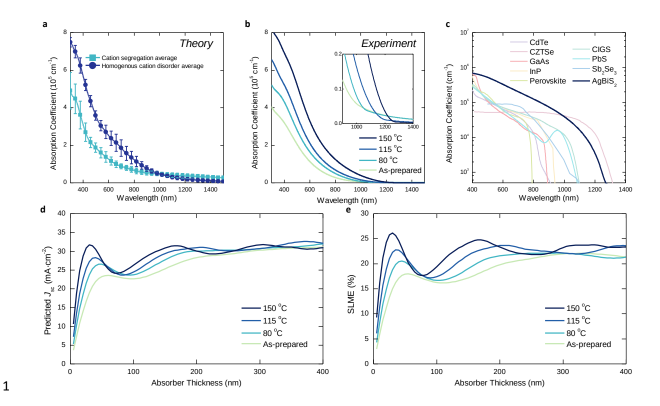
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#### 1 FIGURES

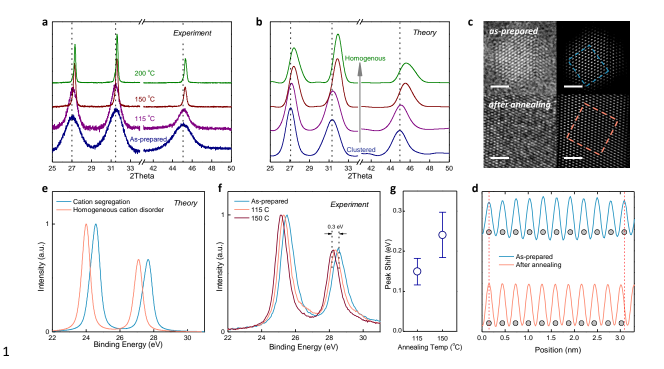




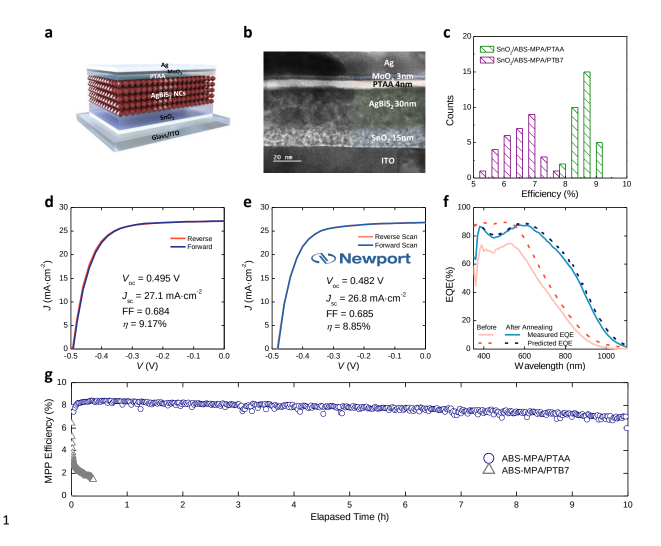
**Figure 1. Absorption enhancement via cation disorder homogenisation. a,** Schematic of AgBiS<sub>2</sub> NCs with cation segregation and homogeneous cation disorder. **b,** Effective electronic band structure of homogeneous disordered AgBiS<sub>2</sub>, alongside the electronic density of states. Atomic orbital contributions have been projected with the colour map (S p: blue, Bi p: green, Ag d: red). VBM set to 0 eV. **c,** Planar-averaged local electronic density of states (LDOS) for cation segregated AgBiS<sub>2</sub>. **d,** Planar-averaged local electronic density of states for homogeneous cation disordered AgBiS<sub>2</sub>. **e,** Normalized Kernel density estimates (KDEs) of Ag and Bi for cation segregation and homogeneous cation disordered AgBiS<sub>2</sub>. Shadow area shows the overlap of Ag and Bi KDEs.



**Figure 2. Absorption coefficients and optical modelling. a,** Simulated absorption coefficients of AgBiS<sub>2</sub> with different cation distributions. Error bars indicate the standard deviation. **b,** Absorption coefficients of AgBiS<sub>2</sub> NCs annealed at different temperatures. Inset: Zoom-in of long wavelength region. **c,** Absorption coefficient of AgBiS<sub>2</sub> NCs films compared with other photovoltaic materials (CdTe<sup>27</sup>, CZTSe<sup>28</sup>, GaAs<sup>4</sup>, InP<sup>29</sup>, Perovskite<sup>30</sup>, CIGS<sup>4</sup>, PbS, Sb<sub>2</sub>Se<sub>3</sub><sup>31</sup>). **d,** Predicted short circuit current density ( $J_{sc}$ ) of AgBiS<sub>2</sub> NCs using the transfer matrix method (TMM). **e,** Spectroscopic Limited Maximum Efficiency (SLME) of AgBiS<sub>2</sub> NCs as a function of film thickness.



**Figure 3.** Characterization of cation configuration transition. (Clockwise) a, Experimental XRD patterns of AgBiS<sub>2</sub> NCs annealed at different temperatures. b, Simulated XRD of 10nm AgBiS<sub>2</sub> NCs with varying cation distribution homogeneity, from cation segregation to homogeneous cation disorder. c, High resolution transmission electron microscope (HRTEM) images of AgBiS<sub>2</sub> NCs before and after 200 °C annealing. Scale bar: 2 nm. d, Integrated line profiles of the {200} crystal plane indicated by dashed lines in c. e, Simulated valence band x-ray photoelectron (XPS) spectra of AgBiS<sub>2</sub> with different cation configurations. f, Experimental XPS spectra of AgBiS<sub>2</sub> NCs annealed at different temperatures. Dashed lines indicates the peak positions before and after annealing. g, Statistics of XPS peak shifts from three different batches of samples. Error bars indicate the standard deviation.



**Figure 4. Ultrathin AgBiS**<sub>2</sub> NCs solar cells. **a,** Schematic of the AgBiS<sub>2</sub> NCs solar cells. **b,** Cross-sectional TEM image of the device. **c,** Statistical histogram of AgBiS<sub>2</sub> NCs solar cells with PTB7 and PTAA as hole transport layer (HTL). **d,** *J-V* curves of the champion device with PTAA as HTL. **e,** *J-V* curves of AgBiS<sub>2</sub> NCs solar cells certified at Newport, USA. **f,** Measured and predicted EQE curves of devices before and after annealing. **g,** Maximum power point (MPP) test of AgBiS<sub>2</sub> devices with PTB7 and PTAA as HTLs.

# 1 TABLES

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# 3 Table 1 Photovoltaic parameters of devices based on different hole transport layers

	$V_{\rm oc}\left(\mathbf{V}\right)$	<b>J</b> <sub>sc</sub> (m <b>A</b> ⋅cm <sup>-2</sup> )	FF	PCE (%)
AgBiS <sub>2</sub> NCs/PTB7	$0.437 \pm 0.011$	$25.59 \pm 0.81$	$0.57 \pm 0.03$	$6.42 \pm 0.55$
Champion device	0.450	26.75	0.63	7.63
AgBiS <sub>2</sub> NCs/PTAA	$0.489 \pm 0.005$	$26.99 \pm 0.76$	$0.66 \pm 0.02$	$8.70 \pm 0.31$
Champion device	0.495	27.11	0.68	9.17
Certified	0.482	26.8	0.68	8.85

Note: statistics are based on 31 different devices.