Highly efficient and stable planar perovskite solar cells with reduced graphene oxide nanosheets as electrode interlayer

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Abstract

We demonstrate a simple solution and room-temperature processed reduced graphene oxide (RGO) as a novel hole-transporting material (HTM) to guarantee highly efficient and highly stable CH3NH3PbI3 perovskite solar cells (PeSCs). The effects of RGO HTM are systemically investigated in terms of PeSC efficiency, PeSC stability, morphology of perovskite film, recombination dynamics, and charge-transport through CH3NH3PbI3/HTM interface. The resultant PeSC with a planar configuration of glass/ITO/RGO/CH3NH3PbI3/PC61BM/bathocuproine (BCP)/Ag exhibits improved device efficiency (maximum PCE of 10.8%) with high reproducibility than those of the reference devices using conventional PEDOT:PSS and GO HTMs. Also, the RGO-based PeSCs show highly desirable device stability in comparison to the PEDOT:PSS PeSCs.

KEYWORDS

Perovskite solar cells; Reduced graphene oxides; Planar structures; Interlayers

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Introduction

Organic-inorganic perovskites employing methylammonium lead halide hold great promise for use as light harvesters in the field of next-generation photovoltaics, and this can be ascribed to their competitive efficiency and low-cost manufacturing [1-8]. The high levels of interests and research activities involving perovskite solar cells (PeSCs) have resulted in great progress in power conversion efficiency (PCE) - more than 17% - in a very short timescale [9-12]. However, the breakthroughs in PeSCs for PeSCs have been achieved mostly by the incorporation of dye-sensitized solar cell architectures that typically require high-temperature processing due to the use of metal oxides as charge-transporting materials, which is one of the big challenges for the market entry of PeSCs. To circumvent these issues, other device configurations based on organic photovoltaic (OPV) structures are being explored for various advantages such as the fabrication through simple and low-temperature processes with flexible formats [4,13-15]. General PeSCs using OPV structures are composed of an indium tin oxide (ITO) anode, poly(ethylenedioxythiophene):poly(styrene sulfonate) (PEDOT:PSS) as a hole-transporting layer, the perovskite layer as a light absorber, [6,6]-phenyl-C61-butyric acid methyl ester (PC61BM) as an electron-transporting material (ETM), and a low work function (WF) metal as a cathode. Among these layers, the hole-transporting and electron-transporting layers play a key role in determining the resultant device performance because the heterojunction between perovskite absorbers and adjacent layers could manipulate the p-i-n characteristics and charge extraction performances in the PeSCs [9,16-18].

The efforts to optimize the heterojunction of electron-transporting layer/perovskite have mostly focused on the fullerene derivatives since, as verified by several reports, the electron charge transfer from perovskites to fullerenes is both very fast (~0.4 ns) and efficient (92%) in the illumination state, thus, fullerene materials have been recognized as the perfect electron quenchers [4,19,20]. Jeng et al. applied fullerene derivatives as electron acceptors with diverse lowest unoccupied molecular orbital (LUMO) levels and demonstrated that the interface of CH3NH3PbI3/perovskite/fullerene could control photovoltaic performances with energy-level alignments [15]. Also, the effect of controlling PC61BM film thickness on device performances has recently been reported for PeSCs [21]. Unlike the intense interests in efficiency enhancements associated with the evolution of various fullerene derivatives and the optimization of process conditions that are noted above, very little interest has been paid to the efforts on hole-transporting materials (HTMs) in PeSCs based on OPV structures, although the efforts on searching other HTMs are as important as those of the electron-transporting materials in the enhancement of PeSC performance. Furthermore, almost exclusively used PEDOT:PSS HTM should be replaced due to the detrimental effects of its hygroscopicity and acidity on device durability as widespread acceptance.

Recently, nickel oxides have been demonstrated as an alternative to PEDOT:PSS as a hole-transporting layer in PeSCs with better energy-level alignments between CH3NH3PbI3 perovskites and nickel oxides [22,23]. However, to achieve a reasonable PCE, the nickel oxide required either a high-temperature process or a vacuum-process and additional UV/O3 treatment, which can nullify the solution-processible manufacturing of PeSC based on OPV structures. Also, although very recent accomplishment of the solution-processible graphene oxide (GO) as an alternative HTM was demonstrated in PeSC with OPV architecture employing a CH3NH3PbI3.Clx absorber [24], the GOs could have their intrinsic disadvantages for HTMs such as an insulating property and a high degree of oxygen contents on their surfaces [25]. Furthermore, the important issues on PeSCs with alternative HTMs to PEDOT:PSS remain concerning the charge-transfer behaviors through a perovskite interlayer, and concerning the device stability of PeSCs. Thus, greater interest in hole-transporting materials should be devoted for development of higher efficiency, increased cost-effectiveness, and more stable perovskite solar cells.

The present study is the first to demonstrate how solution-based reduced graphene oxide (RGO) films processed at room temperature can dramatically raise the device performance of perovskite-based solar cells. As a novel electrode interlayer in PeSCs, we used RGO reduced through p-hydrazinobenzenesulfonic acid hemihydrate, because this RGO satisfies most of the requirements of an ideal HTM in terms of low cost, abundance, solution processibility, conductivity, and stability [25,26]. In particular, recently developed RGOs have shown an enhanced conductivity and a higher work function than conventional PEDOT:PSS and hydrazine-based RGOs [26]. Here, we combined CH3NH3PbI3 light absorbers with an ultra-thin RGO, and we systemically studied the effects of RGO HTM in terms of PeSC efficiency, PeSC stability, morphology of perovskite film, recombination dynamics, and charge transport through CH3NH3PbI3/HTM interface. As a result, a PeSC with a configuration of glass/ITO/RGO/CH3NH3PbI3/PC61BM/bathocuproine (BCP)/Ag exhibited better device efficiency (maximum PCE of 10.8%) with high reproducibility than those of the reference devices using PEDOT:PSS and GO HTMs. Furthermore, the RGO-based PeSCs showed highly desirable device stability in comparison to the conventional PEDOT:PSS PeSCs. This promising strategy will be a critical step toward the ideal HTM design for the advancement of practical perovskite solar cells.

Results and discussion

The PeSC configuration and the chemical structure of RGO used as a HTM are illustrated in Fig. 1(a). Aqueous suspensions of RGO with outstanding dispersibility presented simple fabrication technique with no thermal treatments, and resulted in a thin hole-transporting layer with a thickness of about 3 nm on ITO glass substrates. CH3NH3PbI3 perovskite film for use as a light harvester was grown by recently developed solvent additive process by using a small amount of N-cyclohexyl-2-pyrrolidone (CHP), which resulted in a highly uniform and flat surface of the perovskite film, thereby leading to highly reproducible PeSC performances [27]. Then, an optimized 100 nm-thick PC61BM layer and BCP/Ag cathodes...
covered CH3NH3PbI3 perovskite layers. Fig. 1(b) for the cross-section image of scanning electron microscopy (SEM) shows the well-fabricated and uniform structure of planar PeSC with RGO HTMs. The corresponding energy diagram of each layer is shown in Fig. 1(c). Recent reports have alluded that minimizing the mismatch between the energy levels of perovskite and transporting materials promotes the transfer of charge carriers and elevates the voltage output [9,16,22,28]. Accordingly, the 0.3 eV higher WF of RGO HTMs relative to PEDOT:PSS HTMs as provided in Fig. S1 can be anticipated to show desirable charge transfer across the heterojunction of RGO/CH3NH3PbI3 perovskite with minimal energy losses and minimal degradations in the photovoltage of PeSCs.

In order to explore the potential of the RGO as HTMs in PeSCs, we first evaluated the optical properties of CH3NH3PbI3 perovskite layers on ITO/RGO films. PEDOT:PSS and GO before reduction were chosen as HTMs for a comparative study. In the optical absorption shown in Fig. 2(a), the ability of light harvesting for each of the CH3NH3PbI3 perovskite film was similar, and no noticeable transition of the absorption pattern was obtained over the entire visible spectrum, regardless of a variety of HTM types. For further examination on the compatibility of RGO HTMs within the perovskite devices, we measured the steady-state photoluminescence (PL) spectra. Fig. 2(b) provides the PL characteristics of CH3NH3PbI3 perovskite layers on ITO/PEDOT:PSS, ITO/GO, and ITO/RGO substrates. The intensity of the PL response from the CH3NH3PbI3 perovskite film was reduced on the order of PEDOT:PSS, GO, and RGO. We noted that when the almost equal magnitude of absorbance among the three types of HTMs was accounted for, the RGO exhibited the best PL quenching ability, and the holes generated in perovskite absorbers could be more efficiently transferred into RGO HTMs compared to conventional PEDOT:PSS HTMs [4].

To directly access the performance of RGO as a contact material for CH3NH3PbI3 perovskites, we fabricated the planar PeSCs with an OPV-based configuration on ITO substrates using different HTMs: PEDOT:PSS (~30 nm), GO (~3 nm), and RGO (~3 nm). The detailed statistics and averaged photovoltaic parameters are summarized in Fig. S2 and Table 1. Also, to confirm the accurate characterization of PeSCs in this study, their corresponding hysteresis properties of the photocurrents were measured according to the previous reports as shown in Fig. S3 and S4 [11,29]. Encouragingly, the PeSCs employing RGO HTMs presented the PCE enhancement by ca. 20% relative to conventional devices using PEDOT:PSS as shown in Fig. 2(c) and (d). The representative photovoltaic parameters of the device using RGO HTM showed an open-circuit voltage (VOC) of 0.98 V, a short-circuit current density (JSC) of 15.4 mA cm$^{-2}$, a fill factor (FF) of 71.6%, and a PCE of 10.8%, whereas the control device incorporating PEDOT:PSS exhibited a VOC of 0.91 V, a JSC of 14.1 mA cm$^{-2}$, a FF of 70.8%, and a PCE of 9.1%. Interestingly, however, when the GO was employed as a HTM, where a main chemical structure of GO is also based on a graphene basal plane like RGO, a completely inferior performance to that of the RGO in all photovoltaic parameters was observed as follows: a VOC of 0.88 V; a JSC of 11.9 mA cm$^{-2}$; a FF of 38.5%; and, a PCE of 4.0%. This result could be originated from the electrically insulating properties of GO and the large potential of charge traps due to the oxygenated functionalities and vacancies on the GO sheets [25].

To improve the comparative investigation, we calculated the series resistance (Rs) and the shunt resistance (Rsh) from the photocurrent density-voltage (J-V) curves. We observed a reduced Rs of 0.92 Ω cm$^2$ in the RGO device compared to the PEDOT:PSS device (1.10 Ω cm$^2$), implying a superior charge-transport ability of RGO, which was probably due to a higher conductivity (~3 S cm$^{-1}$) and a higher WF than those of PEDOT:PSS HTMs. More importantly, an unexpected shunt
The external quantum efficiency (EQE) and internal quantum efficiency (IQE) confirmed the enhanced photocurrent of the RGO device, as shown in Fig. 2(e) and (f). Also, all the calculated J_{SC} values using EQE data were within a 3% error with the corresponding J_{SC} values detected in the J-V curves. The PeSC using RGO showed high and broad spectral responses in wavelength from 300 nm to 800 nm with a maximum peak approaching an 80% value of EQE. Furthermore, the IQE spectrum of RGO cells depicted higher values in the overall wavelength range compared to the reference PEDOT:PSS spectrum, indicating that the efficiency of photon-to-electron conversion was more advanced in the RGO device even with similar absorption ability using equal light harvesters and their similar optical transparency shown in Fig. S6 [35].

To further unveil the effects of RGO incorporation on the CH_{3}NH_{3}PbI_{3} perovskite devices and possible correlation with enhanced device performances, the morphological changes of the perovskite films on different substrates (ITO/PEDOT:PSS, PEDOT:PSS, GO, and RGO) were investigated. The absorbance and photoluminescence responses of CH_{3}NH_{3}PbI_{3} films on various substrates are shown in Fig. 2(a) and (b), respectively. The J-V curves for PeSCs with various HTMs are shown in Fig. 2(c), and the corresponding PCE statistics based on 20 devices are shown in Fig. 2(d).

The photovoltaic parameters with average values for PEDOT:PSS, GO, and RGO-based PeSCs are summarized in Table 1. The series resistance and the shunt resistance were calculated from illuminated J-V plots converging to VOC and J_{SC}, respectively. The data in Table 1 show that the RGO device exhibited a higher VOC (9.5 V) and J_{SC} (14.8 mA cm^{-2}) compared to the PEDOT:PSS device (8.8 V and 13.6 mA cm^{-2}), respectively. This suggests that carrier recombination could be more efficiently blocked through the RGO/perovskites heterojunction than through the PEDOT:PSS/perovskites. The PeSC incorporating RGO also exhibited a weaker dependence of VOC on the light intensity as shown in Fig. S5. The PeSC using RGO exhibited a weaker dependence of VOC on the light intensity compared to the device using PEDOT:PSS with a slope of 1.24 kT/e, which indicates that the strength of recombination was significantly reduced in the RGO device compared to the PEDOT:PSS device [13,30,31].

The high electron mobility of RGO nanosheets allows for effective electron transport within the RGO layer, reducing recombination at the RGO/perovskites interface. This, in turn, leads to a higher Voc and J_{SC} in the RGO-based PeSC. Additionally, the lower series resistance (2.95 Ω cm^{2}) of the RGO device that was higher than that of the PEDOT:PSS device (1.65 Ω cm^{2}) was also obtained, suggesting that carrier recombination could be more efficiently blocked through the RGO/perovskites heterojunction than through the PEDOT:PSS/perovskites. This observation is in good agreement with the dependence of Voc on the light intensity as shown in Fig. S5.

The EQE and IQE spectra of RGO cells depicted higher values in the overall wavelength range compared to the reference PEDOT:PSS spectrum, indicating that the efficiency of photon-to-electron conversion was more advanced in the RGO device even with similar absorption ability using equal light harvesters and their similar optical transparency shown in Fig. S6 [35].

To further unveil the effects of RGO incorporation on the CH_{3}NH_{3}PbI_{3} perovskite devices and possible correlation with enhanced device performances, the morphological changes of the perovskite films on different substrates (ITO/PEDOT:PSS...
and ITO/RGO) were investigated using a series of SEM and X-ray diffraction (XRD) analyses, and the surface morphology for each substrate obtained by atomic force microscopy (AFM) was also shown in Fig. S7. All samples were fabricated using procedures identical to those of working devices. As can be seen in the top view SEM images of Fig. 3(a) and (b), both films grown on ITO/PEDOT:PSS and ITO/RGO exhibited densely covered and well-crystallized cubic structures of CH$_3$NH$_3$PbI$_3$ perovskites, which ensure the uniform coverage of upper PC$_{61}$BM layer. The cross-sectional images in Fig. 3(c) and (d) confirm again a compact morphology and an uniform film thickness of 160 nm without any pin-holes across the entire region in both films, which prevents direct contact between the PC$_{61}$BM layer and the HTMs to induce an electrical shunting [14,21]. More notably, the grain size was the most distinguishing morphological characteristics of the two types of samples. The enlarged perovskite grains in the range of 100-200 nm were observed in the perovskite film grown on RGO compared to the perovskite film grown on PEDOT:PSS with grain size below 100 nm. The larger size of the perovskite grains resulted in a reduction in the total number of grain boundaries (GBs) in a given volume as can be seen in the SEM images. Very recently, it was found that the GBs in perovskite film could activate a charge recombination owing to their higher surface potential and defect states [10,29,36]. In this respect, the reduced number of GBs shown in RGO HTMs are believed to facilitate the flow of charge carrier with a minimal recombination loss through the GBs. The improved morphology of the CH$_3$NH$_3$PbI$_3$ perovskite film on an RGO layer is also evident in the corresponding XRD patterns shown in Fig. 3(e). Stronger reflection peaks of the perovskite structure were clearly obtained in the case of RGO HTMs where peaks of 14.1°, 28.4°, and 31.7° are assigned to (110), (220), and (310) crystal planes of the CH$_3$NH$_3$PbI$_3$ perovskites [11]. In particular, the smaller full-width-at-half-maximum (FWHM) of (110) XRD peak (0.49) shown in Fig. S8 represents the better textured domains with a preferred in-plane orientation to substrates, suggesting that the RGO/perovskites interface
could encourage a high degree of crystallinity and texturing [10].

To further examine the influence of RGO on the charge dissociation and recombination processes, we performed time-resolved photoluminescence (TRPL) as shown in Fig. 3(f). We analyzed the PL curves with a bi-exponential decay function including a fast decay (\( \tau_1 \)) component and a slow decay (\( \tau_2 \)) component, and the detailed parameters are summarized in Table 2. The fast decay would be originated from the quenching of charge carriers, meaning that the free carriers generated from the perovskite region were transported to the respective hole or electron contact, and the slow decay component could be attributed to the radiative recombination of free charge carriers before the charge collection [14,36]. In the case of the PEDOT:PSS/CH3NH3PbI3 perovskite, the fast decay lifetime of 5.16 ns with a 0.26 fraction and the slow decay lifetime of 47.95 ns with a 0.48 fraction were detected, which suggests that the depopulation of photogenerated charges was dominated by radiative recombination rather than by an appropriate charge collection through the PEDOT:PSS/perovskite interface. In contrast, for the RGO/CH3NH3PbI3 perovskite film, the fast decay lifetime and the slow decay lifetime were both shortened (4.44 ns and 26.92 ns, respectively) and the fraction of the fast decay process was much more enhanced than the reference of PEDOT:PSS HTMs, which can be interpreted that the RGO/CH3NH3PbI3 perovskite junction could induce a faster charge transfer across its interface with a reduced recombination strength. Furthermore, when the perovskite film forms the junction with PC61BM in addition to RGO, closely resembling the working devices, both types of decay lifetimes were further decreased with negligible fraction of the slow decay component. Therefore, in the case of RGO HTMs, it was apparent that the majority of charge carriers generated under illumination were swept to the respective contacts with minimal recombination, while the recombination process still takes some degree of fraction in PEDOT:PSS HTMs even with a PC61BM electron quencher. The fast charge transfer and collection shown in RGO-based devices may be closely related to the high conductivity, the better-aligned energy levels, the improved morphology such as the reduced GBs, and the high quality of the crystal phase as discussed above, thus contributing to an enhancement of the photovoltaic parameters under working conditions.

To further demonstrate the potential of RGO HTMs, we recorded the device performance as a function of exposure time to ambient conditions with a humidity of approximately 50% without encapsulation according to the ISOS-D-1 protocol [37]. As evident in Fig. 4, the device efficiency of PEDOT:PSS-based cells was drastically deteriorated and showed complete failure in photovoltaic characteristics with 0% just after 120 h of exposure. It is generally acknowledged that a device incorporating PEDOT:PSS is degraded mostly by the erosion of ITO electrodes [26]. In addition to damaged ITO electrodes, for these CH3NH3PbI3 perovskite devices, the hygroscopic and highly acidic properties of PEDOT:PSS can activate the decomposition of CH3NH3PbI3 perovskite components because the lightly fixed CH3NH3 cation in perovskite crystals could be actively converted to CH3NH2 with the aid of moisture [38]. As a consequence, the CH3NH3PbI3 perovskite was changed to hexagonal PbI2 with the emission of CH3NH2 components owing to its very low boiling point (17 °C), which could be accelerated by a PEDOT:PSS/CH3NH3PbI3 perovskite junction. In contrast, the RGO-based devices showed a promising device-stability retaining about 6% (62% of the initial value) of PCE even after 140 h of exposure instead of device failure. The stability of

Table 2 Values for time-resolved PL characteristics by fitting the decay curves of various films.

<table>
<thead>
<tr>
<th>HTM</th>
<th>ETM</th>
<th>( \tau_1 ) (ns)</th>
<th>( \tau_2 ) (ns)</th>
<th>Fraction 1</th>
<th>Fraction 2</th>
<th>Average ( \tau ) (ns)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PEDOT:PSS</td>
<td>-</td>
<td>5.16</td>
<td>47.95</td>
<td>0.26</td>
<td>0.48</td>
<td>45.55</td>
</tr>
<tr>
<td>RGO</td>
<td>-</td>
<td>4.44</td>
<td>26.92</td>
<td>0.42</td>
<td>0.32</td>
<td>22.93</td>
</tr>
<tr>
<td>PEDOT:PSS</td>
<td>PC61BM</td>
<td>2.16</td>
<td>11.11</td>
<td>0.48</td>
<td>0.28</td>
<td>8.89</td>
</tr>
<tr>
<td>RGO</td>
<td>PC61BM</td>
<td>2.04</td>
<td>8.21</td>
<td>0.67</td>
<td>0.09</td>
<td>4.22</td>
</tr>
</tbody>
</table>
devices with RGO stems from the nearly neutral properties of RGO with few surface oxygen functionalities and the inherent passivation ability of RGO against oxygen and moisture, which reduces the decomposition probability of CH$_3$NH$_3$PbI$_3$ perovskite films [26,39]. From this point of view, the incorporation of RGO HTMs is more suitable than the conventional PEDOT:PSS HTMs in terms of excellent lifetime of PeSCs as well as the advances made in device efficiency.

Conclusions

In conclusion, we have introduced simple solution and room-temperature processed reduced graphene oxide, for the first time, as a novel HTM to guarantee highly efficient and highly stable CH$_3$NH$_3$PbI$_3$ PeSCs. The resultant PeSCs with device-architectures comprised of glass/ITO/RGO/CH$_3$NH$_3$PbI$_3$/PC$_{61}$BM/BCP/Ag provided excellent and reproducible device efficiency of up to 10.8%, which was superior to the reference PEDOT:PSS and GO-based devices. Systematic investigation on the mechanisms of enhancement in photovoltaic characteristics revealed that the improved PeSC performances of RGO devices could be due to the facilitated charge collection with a retarded recombination, high RGO conductivity, better-passivation ability greatly extended the cell-operation time compared to PeSCs employing conventional PEDOT:PSS. These successful demonstrations showed RGO to be an advanced interfacial material to surpass conventional PEDOT:PSS HTMs in terms of photovoltaic performance coupled to ambient stability test, we found that the use of RGO HTMs is more suitable than the conventional PEDOT:PSS HTMs in

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Appendix A. Supporting information

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References

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Supporting Information

Highly Efficient and Stable Planar Perovskite Solar Cells with Reduced Graphene Oxide Nanosheets as Electrode Interlayer

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Experimental Section

Preparation of RGO: The GO was synthesized via previously reported approaches and the prepared GO was dispersed in DI water with a concentration of 3 mg ml\textsuperscript{-1} giving brown solution for the synthesis of RGOs [25,26]. For RGO synthesis, \(\mathrm{p}\)-hydrazinobenzenesulfonic acid hemihydrate (5 g) was added into the prepared GO dispersion (3 mg ml\textsuperscript{-1}, 30 ml) to reduce GO without any insulating surfactants [26], and the product solution was stirred at 60 °C for 24 h. After the completion of the reduction process, sodium chloride was added into the solution to precipitate RGO flakes, and subsequently the mixture solution was centrifuged at 10000 rpm for
Finally, for the device fabrication, GO and RGO products were re-dispersed in DI water at a concentration of ~0.6 mg ml$^{-1}$.

**Sample characterizations:** The measurements of ultraviolet photoemission spectroscopy (UPS) (AXIS NOVA, Kratos) were performed using the He I (hν = 21.2 eV) excitation source with energy resolution of 0.1 eV under a pressure of 10$^{-8}$ torr. The reference of Fermi energy was determined with a cleaned gold surface. UV-vis/NIR spectra were recorded using a Jasco V-570 UV-vis/NIR spectrophotometer and the steady-state photoluminescence were measured using a Hitachi F-7000 UV with an excitation at 400 nm. The images of scanning electron microscope (SEM) were collected using a HITACHI S-4700, and the data of X-ray diffraction (XRD) were taken using a Rigaku Micromax-002 diffractometer with CuKα (λ=0.1542 nm) radiation. Time-resolved photoluminescence (TRPL) measurements were conducted by the 2nd harmonic generation (at 400 nm) of a mode-locked Ti:Sapphire laser (Chameleon Ultra II, Coherent) with a pulse duration of 150 fs equipped with an external pulse-picker (9200 series, Coherent Inc.) to reduce the repetition rate at 3.6 MHz. The fluorescence radiation was spectrally filtered to a narrow bandwidth and focused into the entrance slit of a 300 mm spectrograph (Acton SpectraPro 2300i, Princeton Instruments) with a spectral resolution of about 1nm. Then, PL lifetime was monitored with a picosecond streak camera (C11200, Hamamatsu Photonics).

**PeSC fabrications and measurements:** The patterned glass/ITO substrates (12 Ω sq$^{-1}$) were treated under UV/O$_3$ for 20 min before the interfacial layer coating. The PEDOT:PSS (Clevios$^\text{TM}$ P VP AI 4083, Heraeus), GO (0.6 mg ml$^{-1}$), and RGO (0.6 mg ml$^{-1}$) solutions were spin-coated at 5000 rpm for 40 s onto prepared substrates, and then the only PEDOT:PSS film was annealed at 150 °C for 10 min. For the deposition of CH$_3$NH$_3$PbI$_3$ perovskites films, the substrates were transferred to a N$_2$-filled glovebox (<1 ppm O$_2$, and H$_2$O). Methylammonium iodide (MAI) was
synthesized via a previously reported method [8], and all reagents including solvents were purchased from Sigma Aldrich and used without any further purification. The precursor solution (35 wt%) was prepared by dissolving MAI and PbI$_2$ with a 1:1 molar ratio (0.262 g: 0.759g) in anhydrous N,N-dimethylformamide (DMF) solvent (2 ml), stirring for 12 h inside a N$_2$-filled glovebox at 60 °C. After that, 5 vol% N-cyclohexyl-2-pyrrolidone (CHP) was added into the precursor solution as a morphology controller before the fabrication of the perovskite film [27]. The mixture solution was spin-coated on top of each interfacial layer at 5000 rpm for 90 s. The films became darker and glassy after subsequent annealing at 100 °C for 2 min. After that, a chlorobenzene solution consisting of PC$_{61}$BM (Nano-C) with the concentration of 20 mg ml$^{-1}$ was spin-coated at 1000 rpm for 40 s onto the perovskite layer. Finally, cathodes composed of bathocuproine (BCP) (10 nm)/Ag (80 nm) were deposited using a thermal evaporator under a pressure of 10$^{-6}$ torr and an active area of 4.64 mm$^2$ was defined by a shadow mask. The thickness of deposition was checked with the quartz crystal sensor. The photocurrent density-voltage curves were measured with a Keithley 2400 instrument under 100 mW cm$^{-2}$ illumination at AM 1.5 G condition. For accurate measurement, a standard Si solar cell certified by the International System of Units (SI) (SRC 1000 TC KG5 N, VLSI Standards, Inc) was used for the calibration of light intensity. The EQE and IQE spectra were simultaneously obtained via an Oriel® IQE-200 QE Measurement System. The device-operation times of PeSC without encapsulations were recorded as a function of exposure time to ambient environment with a humidity of approximately 50% using equal instrumental conditions.
**Figure S1** UPS spectra in the secondary electron cutoff region of the glass/ITO/PEDOT:PSS and glass/ITO/RGO substrates.
**Figure S2** The statistics of photovoltaic parameter for (a) open-circuit voltages ($V_{OC}$), (b) short-circuit currents ($J_{SC}$), and (c) fill factors (FF) obtained from 20 devices.
Figure S3 The J-V characteristics for PeSCs based on various HTMs: (a) PEDOT:PSS, (b) GO, and (c) RGO. The different sweep rates (from about 0.5 V/s to 0.21 V/s) and different sweep directions were compared. No noticeable hysteresis was observed except for devices with GO HTMs.
Figure S4 Comparison of the photocurrent-output for PeSCs using different HTMs measured at the respective maximum-voltages for the maximum-currents, and the 0 voltage for the $J_{SC}$ as a function of time. The 301 scan points were measured during ~ 50 s scan times. There were no obvious current transients in all types of devices.
Figure S5 The measured $V_{OC}$ of PeSCs with PEDOT:PSS, GO, and RGO HTMs as a function of the light intensity with linear fitting to the data in a semi-logarithmic scale. The slopes of linear fitting are 1.89, 2.95, and 1.24 kT/e for the devices with PEDOT:PSS, GO, and RGO, respectively.
Figure S6 Optical transmittance spectra with various HTMs on glass/ITO substrates.
Figure S7 AFM topographic images for the surfaces of (a) PEDOT:PSS film, and (b) RGO film on ITO substrates. The scan window is 2×2 μm².
Figure S8 The calculated values via Gaussian fitting of full-width-at-half-maximum (FWHM) for the (110) XRD peak.