Metal-free carbon-based electrocatalysts for dye-sensitized solar cells (DSSCs) are sufficiently active in Co(II)/Co(III) electrolytes but are not satisfactory in the most commonly used iodide/triiodide ($I^-/I_3^-$) electrolytes. Thus, developing active and stable metal-free electrocatalysts in both electrolytes is one of the most important issues in DSSC research. We report the synthesis of edge-selenated graphene nanoplatelets (SeGnPs) prepared by a simple mechanochemical reaction between graphite and selenium (Se) powders, and their application to the counter electrode (CE) for DSSCs in both $I^-/I_3^-$ and Co(II)/Co(III) electrolytes. The edge-selective doping and the preservation of the pristine graphene basal plane in the SeGnPs were confirmed by various analytical techniques, including atomic-resolution transmission electron microscopy. Tested as the DSSC CE in both Co(bpy)$_3^{2+/3+}$ (bpy = 2,2'-bipyridine) and $I^-/I_3^-$ electrolytes, the SeGnP-CEs exhibited outstanding electrocatalytic performance with ultimately high stability. The SeGnP-CE–based DSSCs displayed a higher photovoltaic performance than did the Pt-CE–based DSSCs in both SM315 sensitizer with Co(bpy)$_3^{2+/3+}$ and N719 sensitizer with $I^-/I_3^-$ electrolytes. Furthermore, the $I^-_3$ reduction mechanism, which has not been fully understood in carbon-based CE materials to date, was clarified by an electrochemical kinetics study combined with density functional theory and nonequilibrium Green’s function calculations.

**RESULTS**

**Synthesis and characterization of SeGnPs**

SeGnPs were prepared by mechanochemical reaction between graphite and selenium (Se) powders (fig. S1). The mechanochemical reaction is driven by kinetic energy (20). For example, high-velocity stainless steel balls in a planetary ball mill reactor generate sufficient kinetic energy, which is delivered to graphite and Se powders and induces unzipping of graphitic C–C bonds and cracking of Se–Se bonds (fig. S1). Activated C and Se atoms have an opportunity to form covalent C=Se and C–Se–C bonds (as like C=O and C–O–C bonds) along the broken edges of SeGnPs. The resultant SeGnPs were characterized by various techniques including scanning electron microscopy (SEM; fig. S2) and transmission electron microscopy (TEM; Fig. 1 and fig. S3) with element mappings, energy-dispersive spectroscopy (EDS; fig. S4A and table S1), x-ray photoelectron spectroscopy (XPS; fig. S4), transmission electron microscopy (TEM; fig. S5A), Raman spectroscopy (fig. S5B), x-ray diffraction (XRD) patterns (fig. S5C), contact

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angle measurements (fig. S5D), thermogravimetric analyses (fig. S6), and solubility tests (fig. S7). SEM-EDS indicated that the content of Se is approximately 5.57 atomic % (table S1). More detailed characterization data with corresponding discussions are presented in the Supplementary Materials. In particular, the formation of C=Se and C-Se-C bonds at the edges of SeGnPs was demonstrated by a low-voltage spherical aberration (Cs)-corrected atomic-resolution TEM (AR-TEM; Fig. 1B and fig. S8A), high-resolution TEM (HR-TEM; fig. S8B), and scanning TEM (STEM; fig. S8, C and D). The AR-TEM images obtained at the edges of a thin platelet demonstrated in Fig. 1B and fig. S8A show that the Se atoms with the dark atomic contrast are present only along the edge lines of SeGnPs. In contrast, the inner part (basal area) of the sheets shows a perfect graphitic structure, indicating that the mechanochemical unzipping of graphitic C–C bonds does not destroy the crystallinity of the basal plane. A high-angle annular dark-field STEM image (fig. S8, C and D) with atomic Z contrast shows the existence of single Se atoms (arrow, bright spots) but not Se clusters. AR-TEM images (arrows, bright atomic contrast of Fig. 1, C and D) and corresponding IFFT images (bright atomic contrast of Fig. 1, E and F) confirm that the single Se atoms are attached at the edge of the armchair or zigzag configurations of graphene, which agrees with our model of edge-selective formations of C–Se bonds, proposed in Fig. 1A.

**Electrocatalytic activity of SeGnPs**

Given the identification of structure and properties, the electrocatalytic activities of SeGnPs were evaluated for use as CEs in DSSCs. The formation of SeGnP thin films (fig. S9) and fabrication of DSSCs are described in the Supplementary Materials. Electrocatalytic activity was tested with a symmetrical dummy cell by using two identical electrodes through the electrochemical impedance spectroscopy (EIS) technique (fig. S10A). The EIS spectra obtained from Co(bpy)$_3^{2+/3+}$ and I$_3^−$/I$^−$ electrolytes are shown in Fig. 2 (A and B). Regardless of the electrodes, the EIS spectra for Co(bpy)$_3^{2+/3+}$ electrolyte exhibited two distinct semicircles in high- and low-frequency regions, and they were fitted to the Randles-type equivalent circuit (EC; fig. S10B) (21). The corresponding EIS parameters are summarized in Table 1. The SeGnPs exhibited profoundly better electrocatalytic activity than those of the reference Pt and edge-hydrogenated GnPs (HGnPs) (fig. S11). For example, the charge-transfer resistances ($R_{ct}$) of the Pt-CEs and SeGnP-CEs were 1.85 and 0.14 ohms-cm$^2$, respectively, with corresponding exchange current densities ($i_0$) of 13.9 and 234 mA cm$^{-2}$.

Unlike the Pt with two semicircles, for which the high- and low-frequency semicircles correspond to the charge-transfer and Nernst diffusion processes (Fig. 2B), respectively (21, 22), the EIS spectra of SeGnPs (Fig. 2B) and HGnPs (Fig. S12) in I$_3^−$/I$^−$ electrolytes exhibit three semicircles, which is a general feature of highly porous carbon materials (6, 13, 23, 24). Comparing the SeGnP and HGnP electrodes, there is only a slight difference in the magnitude of the first semicircles, but there is a considerable difference in the second semicircles. In this case, the effective charge-transfer resistance ($R_{ct}$) should be considered, and several ECs have been proposed to identify and quantify these effects (fig. S13) (14, 23, 24). For the first semicircle newly emerging in the carbon-based CEs, several proposals have been made as to its origin, including Nernst diffusion in the pores; however, some issues still remain. For example, several of our results conflict with the first semicircle arising from any type of Nernst diffusion in the pores: (i) there was no line of slope 1 (the Warburg line) at the beginning of the semicircle (Fig. 2B); (ii) the semicircles were not enlarged by increasing the electrode thickness (fig. S14, D and E), electrolyte viscosity (fig. S14F), or external bias potential (fig. S15); and (iii) the pore size of the SeGnPs was 26 Å (table S2), which is considerably larger than that of I$_3^-$ and Co(bpy)$_3^{2+}$ ions (table S3). Considering the size difference between I$_3^-$ and Co(bpy)$_3^{2+}$, apparently, the first semicircle that appeared at a high
frequency could not be explained as Nernst diffusion in the pores of the electrode, which can also be characterized by a potential step chronoamperometry (CA; fig. S16).

In addition to the EIS analysis, cyclic voltammetry (CV) with different scan rates was carried out in both electrolytes (figs. S17 to S19). The remarkably higher anodic peak current ($I_{pa}$) and cathodic peak current ($I_{pc}$) observed for SeGnPs compared to Pt suggest higher electrocatalytic activity for the redox reaction of $\text{Co(bpy)}_3^{2+/3+}$ and $\Gamma/\Gamma_3^-$, which could be attributed to the lower $R_{ct}$ at the SeGnP/electrolyte interface as observed in EIS measurements.

The electrochemical stability of CE materials is also one of the important factors that need to be considered for practical DSSC applications. On the basis of this consideration, the electrochemical stability of the SeGnPs was tested in both electrolytes (fig. S20). The reciprocal $R_{ct}$ values of SeGnPs were nearly invariable after 1000 CV cycles in both electrolytes, whereas those of Pt decreased by less than 10% in both electrolytes at the final cycle (Fig. 2, C and D). SeGnPs displayed similar electrochemical stability even with a 1-month rest after 1000 CV measurements under normal laboratory conditions (fig. S21).

### Theoretical calculations

We carried out DFT calculations for the CE/electrolyte interface to clarify the high electrocatalytic activity of SeGnPs for $\Gamma/\Gamma_3^-$ reduction. The $\Gamma_3^-$ reduction contains three elementary reaction steps

\[
\Gamma^- + e^- \rightarrow \Gamma (\text{sol}) \quad \text{(III)}
\]

where the asterisk indicates the free site on the CE and “sol” indicates the electrolyte solvent. The solution reaction step I typically proceeds rapidly and in equilibrium, or the molecular iodine reduction reaction (IRR) steps II and III occurring at the CE/electrolyte interface are the rate-determining steps (21, 22). Steps II and III in Pt occur rapidly and nearly simultaneously and induce the first of its two semicircles in the Nyquist plot (21, 22). It has been recently shown that the adsorption

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**Table 1.** EIS parameters of symmetrical dummy cells with Pt and SeGnP electrodes. $R_0$, serial resistance; $R_{ct}$, charge-transfer resistance; $C_{dl}$, double-layer capacitance; $R_{tm}$, transmission resistance; $C_{ad}$, capacitance due to the adsorption of $\Gamma/\Gamma_3^-$ on the graphitic basal plane; $J_0$, exchange current density.

<table>
<thead>
<tr>
<th>CE</th>
<th>Electrolyte</th>
<th>$R_s$ (ohm-cm²)</th>
<th>$R_{ct}$ (ohm-cm²)</th>
<th>$C_{dl}$ (μF cm²)</th>
<th>$R_{tm}$ (ohm-cm²)</th>
<th>$C_{ad}$ (μF cm²)</th>
<th>$J_0$ (mA cm⁻²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt</td>
<td>$\text{Co(bpy)}_3^{2+/3+}$</td>
<td>3.12</td>
<td>1.85</td>
<td>8.88</td>
<td>—</td>
<td>—</td>
<td>13.9</td>
</tr>
<tr>
<td>SeGnPs</td>
<td>$\text{Co(bpy)}_3^{2+/3+}$</td>
<td>3.20</td>
<td>0.13</td>
<td>17.5</td>
<td>—</td>
<td>234</td>
<td></td>
</tr>
<tr>
<td>Pt</td>
<td>$\Gamma/\Gamma_3^-$</td>
<td>3.42</td>
<td>0.61</td>
<td>6.02</td>
<td>—</td>
<td>21.1</td>
<td></td>
</tr>
<tr>
<td>SeGnPs</td>
<td>$\Gamma/\Gamma_3^-$</td>
<td>2.92</td>
<td>0.23</td>
<td>133</td>
<td>0.17</td>
<td>12.4</td>
<td>32.1</td>
</tr>
</tbody>
</table>

---

**Fig. 2.** Comparison between conventional Pt and SeGnP electrodes. (A and B) Nyquist plots of the Pt-CEs and SeGnP-CEs: $\text{Co(bpy)}_3^{2+/3+}$ (A) and $\Gamma/\Gamma_3^-$ (B). The inset in (A) is an enlargement of the EIS spectrum of the SeGnP-CEs in high-frequency range. (C and D) Normalized $R_{ct}$ and $\mathcal{R}_{ct}$ changes versus the EIS scan number: $\text{Co(bpy)}_3^{2+/3+}$ (C) and $\Gamma/\Gamma_3^-$ (D).
energy of I atom can serve as a good descriptor for the IRR. Specifically, it was argued that the adsorption energy should range between 0.33 and 1.20 eV, of which the former ensures the I₂ molecular dissociation and the latter desorption of I⁻ into I⁻(sol) (22, 25). Following the above-described experimental characterizations that have observed Se atoms only at the edges of GnP, we set up SeGnP models based on the representative HGNP armchair and zigzag edges (Fig. 3A). Carefully treating the ferromagnetically spin-polarized graphene zigzag edge states (26, 27), we performed DFT calculations to estimate the adsorption energies of atomic iodine under the vacuum and explicit acetonitrile solvation conditions (figs. S22 to S24) (28–31). For the pristine armchair and zigzag edges as well as the basal plane of GnP, the I atom adsorption energies were determined to be below the optimal range for the CE catalytic activity (−0.17, 0.17, and −0.49 eV, respectively). Edge models involving double-coordinated Se (C–Se–C) or oxidized Se atoms were also unable to support stable I atom adsorption (Fig. 1A).

Fig. 3. Theoretical calculation and proposed mechanism of IRR. (A) For the representative single-coordinated, double-coordinated, and hydrogenated Se [Se(c1), Se(c2), and SeH, respectively]–doped armchair (ac) and zigzag (zz) graphene edges (top panel), the adsorption energies of the I atom explicitly solvated by acetonitrile molecules were evaluated and compared with the undoped edge and basal plane cases (bottom panel). The Pt(111) value of 0.52 eV has been taken from the study of Li et al. (24). In the bottom panel, the shaded region indicates the IRR activity criterion. (B) For various I⁻ and I₃⁻-adsorbed graphene basal plane models (top panel and figs. S27 and S28), the current-voltage (I-V) curves were calculated and compared with those from pristine graphene (bottom panel). In the top panel, Mulliken charge populations were coded into atomic structures. (C) IRR mimetic diagram on the SeGnP surface. (D) Nyquist plots of SeGnP-CEs and their EC at room temperature.
and fig. S22). However, the I atom binding criterion was well satisfied for the single-coordinated Se-doping cases (C=Se), with adsorption energies of 0.77 and 0.48 eV for the armchair and zigzag edges, respectively.

We next theoretically investigate the origin of the first semicircle appearing in the Nyquist plot of the SeGnP-CEs in $\Gamma/\Gamma'$ electrolytes (Fig. 2B). As mentioned above, it is not fully understood why the EIS spectra of carbon-based materials, including HGNPs and SeGnPs, have three semicircles. Having identified the Se-doped graphitic edges as the primary sites where the $I_3^-$ reduction takes place and as being responsible for the charge-transfer process that results in the second semicircle, we can assume that the first semicircle originates from the graphitic basal plane. To confirm this scenario, we carried out combined DFT (29, 32–34) and NEGF calculations (35, 36) and analyzed the structural, electronic, and charge transport properties of graphene adsorbed by $\Gamma/\Gamma'$ (figs. S25 and S26). The $I_3^-$ and $\Gamma'$ ions were calculated to be adsorbed on the graphitic basal plane at the distances of 3.18 to 3.37 Å. This induces a n-type shift of the graphene band and significantly enhances the transmission or increases currents at finite bias voltages (Fig. 3B and fig. S28). The current increase has also been experimentally observed in CV measurements (figs. S18B and S19B), and we propose that although the graphitic basal plane cannot drive IRR by itself, its presence of the first semicircle with Co(bpy)$_3$$^{2+}$ electrocatalysts shows significant Nernst diffusion impedance ($Z_{\text{ct}}$) directly related to IRR activity at the Se-doped edge of the graphene and electrolyte interface. The third semicircle is the Nernst diffusion impedance ($Z_{\text{ct}}$) of the redox couple between two electrodes. As a result, we can propose a new EC to elucidate the EIS spectrum of SeGnPs obtained from the IRR (Fig. 3D), with their EIS parameters summarized in Table 1. From the second semicircle of SeGnPs, the $R_{\text{ct}}$ of SeGnPs (0.23 ohm-cm$^2$) was considerably lower than that of Pt (0.61 ohm-cm$^2$). From the first semicircles, the $R_{\text{ct}}$ was 0.17 ohm-cm$^2$, and thus, the effective charge-transfer resistance ($R_{\text{ct}} = R_{\text{ct}} + R_{\text{dl}}$) of SeGnPs is 0.40 ohm-cm$^2$, which is also lower than that of Pt. Thus, SeGnP-CEs are expected to exhibit a higher short-circuit current density ($J_{\text{sc}}$) and fill factor (FF) in DSSCs.

**DISCUSSION**

This resulted in the net enhancement of the PCE, which was also confirmed with photocurrent transient dynamics and EIS measurements (fig. S30). The mass transport effect of the Co(bpy)$_3$$^{2+}$ and $\Gamma'/\Gamma$ redox couples in actual DSSCs with different CEs was investigated by monitoring photocurrent transients using a multistep on/off modulation of the mass transfer limitation (fig. S30, A and B) (37, 38). As can be seen, the ratio of the initial peak current to the steady-state current in the photocurrent transients is almost consistent in Pt-DSSCs and SeGnP-DSSCs with Co(bpy)$_3$$^{2+}$, whereas Pt-DSSCs and SeGnP-DSSCs with $\Gamma/\Gamma'$ redox couple significantly different. Considering that the DSSCs were fabricated under similar conditions, the mass transport of Co(bpy)$_3$$^{2+}$ and $I_3^-$ ions in the pores of SeGnPs does not affect the overall cell efficiency.

To further understand the improved performance of the DSSCs with the SeGnP-CEs, EIS measurements were carried out on the DSSCs (fig. S30, C and D). The first semicircles of the DSSCs with the SeGnP-CEs are smaller than those of the Pt-CEs. From the first semicircles, the $R_{\text{ct}}$ values calculated by using the EC (inset in fig. S30E) on the DSSC-SeGnP-CEs were 0.34 and 1.54 ohms-cm$^2$ for the Co(bpy)$_3$$^{2+}$ and $\Gamma'/\Gamma'$ redox couples, respectively, whereas the corresponding values of the DSSC-Pt-CEs were 1.27 and 1.95 ohms-cm$^2$. The lower $R_{\text{ct}}$ values for the DSSCs with the SeGnP-CEs suggest higher electrocatalytic activities for the reduction of Co(bpy)$_3$$^{2+}$ and $I_3^-$ ions than...
Table 2. Photovoltaic performance of Pt-DSSCs and SeGnP-DSSCs with different electrolytes. $J_{sc}$, short-circuit current density; $V_{oc}$, open-circuit voltage.

<table>
<thead>
<tr>
<th>CE</th>
<th>Dye</th>
<th>Electrolyte</th>
<th>$J_{sc}$ (mA mA$^{-2}$)</th>
<th>$V_{oc}$ (mV)</th>
<th>FF (%)</th>
<th>PCE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt</td>
<td>SM315</td>
<td>Co(bpy)$_3^{2+/3+}$</td>
<td>15.30 ± 0.36</td>
<td>863 ± 2.0</td>
<td>76.6 ± 0.4</td>
<td>10.11 ± 0.25</td>
</tr>
<tr>
<td>SeGnP</td>
<td></td>
<td></td>
<td>16.27 ± 0.03</td>
<td>876 ± 6.5</td>
<td>77.0 ± 1.4</td>
<td>10.98 ± 0.17</td>
</tr>
<tr>
<td>Pt</td>
<td>N719</td>
<td>$I^-/I_3^-$</td>
<td>17.26 ± 0.32</td>
<td>729 ± 6.8</td>
<td>72.2 ± 1.1</td>
<td>9.07 ± 0.20</td>
</tr>
<tr>
<td>SeGnP</td>
<td></td>
<td></td>
<td>18.16 ± 0.44</td>
<td>692 ± 2.9</td>
<td>73.1 ± 1.3</td>
<td>9.17 ± 0.04</td>
</tr>
</tbody>
</table>

the Pt counterparts, which could enhance the DSSC performance, in particular FF and $J_{sc}$ (Table 2). The result is consistent with the feasibility of the heterogeneous electron-transfer process at the SeGnP/electrolyte interface, as indicated by the smaller $R_{ct}$ (Fig. 4B).

In summary, we have proven SeGnP to be Pt-free alternative CE for the DSSCs in both Co(bpy)$_3^{2+/3+}$ and $I^-/I_3^-$ electrolytes. Whereas carbon-based materials do not work properly for IRR, SeGnP displayed unusual dual electrocatalytic activities with very high electrochemical stability in both electrolytes. Furthermore, the IRR mechanism, which has not been fully understood on carbon-based materials to date, was clarified by an electrochemical kinetics study combined with DFT and NEGF calculations. Se doping at the edges of GnP, especially single-coordinated armchair and zigzag edges, was revealed as the source of the remarkably enhanced electrocatalytic activity. The well-preserved graphitic basal plane then rapidly transfers electrons from the external circuit to $I_3^-$ ions at the edges, thus reducing the $R_{ct}$ and hence improving the photovoltaic performance. On the basis of this new understanding, we proposed a new EC that should help in interpreting the IRR mechanism for other porous carbon-based materials such as heteroatom-doped graphene.

**MATERIALS AND METHODS**

**Synthesis of SeGnPs**

The SeGnPs were prepared simply by ball milling in a planetary ball mill capsule. Graphite (5.0 g) and Se (20.0 g) were placed into a stainless steel capsule containing stainless steel balls (500 g; diameter, 5 mm). After five charging/discharging cycles of argon under reduced pressure (0.05 mmHg), the capsule was sealed. Then, it was fixed in the planetary ball mill machine and agitated at 500 rpm for 48 hours. The resultant product was Soxhlet-extracted with CS$_2$ and further washed with 2.0 M aqueous KOH solution to remove unreacted Se and then repetitively extracted with 1 M aqueous HCl solution to get rid of remaining metallic impurities, if any. The final product was then freeze-dried at −120°C under a reduced pressure (0.05 mmHg) for 48 hours to yield 6.52 g (SeGnP contained at least 1.52 g of Se) of dark black SeGnP powder.

**Fabrication and characterization of DSSCs**

FTO plates were cleaned in detergent solution, water, and ethanol using an ultrasonic bath. The FTO substrates were immersed in 40 mM aqueous TiCl$_4$ solution at 70°C for 30 min and washed with water and ethanol. A TiO$_2$ colloidal paste (18NR-T, Dyessol) was screen-printed onto FTO/glass and sintered at 500°C for 30 min in air. The thickness of the transparent layer was measured by an Alpha-step 250 surface profilometer (Tencor Instruments), and a paste for the scattering layer containing 500-nm-sized anatase particles (STP-500N, ENB Korea) was screen-printed and then dried for 2 hours at 25°C. The TiO$_2$ electrodes were sintered at 500°C for 30 min. The resulting TiO$_2$ photoanodes were immersed in a THF/ethanol (v/v, 2:1) solution containing 0.3 mM SM315 sensitizer (39) and 0.3 mM HC-A4 coadsorbent (or 0.3 mM N719 sensitizer only) (40) and kept at room temperature for 12 hours (see Supplementary Materials and Methods). The dye-adsorbed TiO$_2$ photoanodes were assembled with Pt-CEs and SeGnP-CEs using a thermal adhesive film (25-μm-thick Surlyn, DuPont) as a spacer to produce a sandwich-type cell. Electrolyte solution was introduced through a drilled hole on the CEs via vacuum backfilling. The hole was sealed with cover glass using Surlyn. The light intensity was adjusted with a Si solar cell that was double-checked with a National Renewable Energy Laboratory–calibrated Si solar cell (PV Measurement Inc.). The applied potential and cell current were measured using a Keithley digital source meter (model 2400). Photoelectrochemical data were measured using a 1000-W xenon light source (Oriel, 91193) that was focused to give 100 mW cm$^{-2}$ (1 sun at AM 1.5 G). The $J$-$V$ characteristics of the cell under these conditions were determined by externally biasing the cell and measuring the generated photocurrent. This process was fully automated using the WaveMetrics software. The measurement settling time between applying a voltage and measuring a current for the $J$-$V$ characterization of DSSC was fixed to 80 ms.

**DFT calculations**

To determine the Se-doping geometries and iodine atom adsorption energies, we carried out DFT calculations using the VASP package (28) within the Perdew-Burke-Ernzerhof (PBE) parametrization of generalized gradient approximation (GGA) (29) augmented by DFT-D3 corrections (30). Valence electronic states were expanded in plane-wave basis sets with a 400-eV energy cutoff, and project-augmented wave method was used to represent core-valence electron interactions (31). All the structures were relaxed until the Hellmann-Feynman forces on each ion were less than 0.01 eV Å$^{-1}$. The vacuum distance between periodic images was modeled to have more than 16 Å. The single Γ-point in the Brillouin zone was sampled.

**NEGF calculations**

Electron transport properties were studied by conducting DFT-based NEGF calculations using the SIESTA-TranSIESTA package (32, 35). For the initial DFT calculations, we used the PBE GGA (29) corrected by the DFT-D2 scheme (33). Core electrons were replaced by Troullier-Martins–type norm-conserving pseudopotentials (34), and the Kohn-Sham wave functions were expanded in terms of double-$\zeta$ plus polarization-level numerical atomic basis sets defined by a confinement energy of...
100 meV. Real-space mesh corresponding to a cutoff energy of 200 rydbergs was used for grid operations, and a Monkhorst-Pack $2 \times 2 \times 1$ $k$-point mesh was used to sample the Brillouin zone. For the subsequent NEGF calculations, we chose the charge transport direction along the armchair dimer lines, and starting from an orthogonal graphene cell composed of four C atoms, constructed a $5 \times 12$ supercell (5 dimer lines and 24 zigzag chains along the transport-normal and transport directions, respectively). The supercell was then divided into three equal-area regions, each of which was assigned as electrode 1, scattering region, or electrode 2. A fine $20 \times 1 \times 1$ Monkhorst-Pack grid was used for the $k_z$-point sampling along the transport-normal direction. To generate surface Green’s functions, 15 $k_z$ points along the transport direction were sampled for each electrode. In obtaining transmission functions, an energy grid from 2.0 eV below to 2.0 eV above the Fermi level was sampled at the 0.01-eV resolution. Using the transmission functions, the $I-V$ characteristics were calculated according to the Landauer-Büttiker formula.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/2/6/ea1501459/DC1

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Edge-selenated graphene nanoplatelets as durable metal-free catalysts for iodine reduction reaction in dye-sensitized solar cells

Myung Jong Ju, In-Yup Jeon, Hong Mo Kim, Ji Il Choi, Sun-Min Jung, Jeong-Min Seo, In Taek Choi, Sung Ho Kang, Han Seul Kim, Min Jong Noh, Jae-Joon Lee, Hu Young Jeong, Hwan Kyu Kim, Yong-Hoon Kim and Jong-Beom Baek

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