Direct Synthesis of Molybdenum Phosphide Nanorods on Silicon Using Graphene at the Heterointerface for Efficient Photoelectrochemical Water Reduction

Sang Eon Jun¹, Seokhoon Choi¹, Shinyoung Choi², Tae Hyung Lee¹, Changyeon Kim¹, Jin Wook Yang¹, Woon-Oh Choe¹, In-Hyuk Im¹, Cheol-Joo Kim² *, Ho Won Jang¹ *

HIGHLIGHTS

- MoP nanorod-array catalysts were directly synthesized on graphene passivated silicon photocathodes without secondary phase.
- Mo-O-C covalent bondings and energy band bending at heterointerfaces facilitate the electron transfer to the reaction sites.
- Numerous catalytic sites and drastically enhanced anti-reflectance of MoP nanorods contribute to the high solar energy conversion efficiency.

ABSTRACT Transition metal phosphides (TMPs) and transition metal dichalcogenides (TMDs) have been widely investigated as photoelectrochemical (PEC) catalysts for hydrogen evolution reaction (HER). Using high-temperature processes to get crystallized compounds with large-area uniformity, it is still challenging to directly synthesize these catalysts on silicon photocathodes due to chemical incompatibility at the heterointerface. Here, a graphene interlayer is applied between p-Si and MoP nanorods to enable fully engineered interfaces without forming a metallic secondary compound that absorbs a parasitic light and provides an inefficient electron path for hydrogen evolution. Furthermore, the graphene facilitates the photogenerated electrons to rapidly transfer by creating Mo-O-C covalent bondings and energetically favorable band bending. With a bridging role of graphene, numerous active sites and anti-reflectance of MoP nanorods lead to significantly improved PEC-HER performance with a high photocurrent density of 21.8 mA cm⁻² at 0 V versus RHE and high stability. Besides, low dependence on pH and temperature is observed with MoP nanorods incorporated photocathodes, which is desirable for practical use as a part of PEC cells. These results indicate that the direct synthesis of TMPs and TMDs enabled by graphene interlayer is a new promising way to fabricate Si-based photocathodes with high-quality interfaces and superior HER performance.

KEYWORDS Photoelectrochemical water splitting; Silicon; Molybdenum phosphide; Hydrogen evolution; Graphene
1 Introduction

Hydrogen is expected to become sustainable future energy owing to zero-emission, non-toxicity, and large energy density [1]. However, it usually exists in covalent compounds combined with other elements such as oxygen, nitrogen, and organic materials. One of the promising methods to obtain pure hydrogen from these compounds is the coupling of solar energy to electrochemical water splitting, such as photovoltaic-electrochemical (PV-EC), and photoelectrochemical (PEC) water splitting [2, 3]. Out of two systems, a single cost-effective device can be achieved by PEC water splitting which efficiently converts intermittent and unpredictable solar energy to chemical binding energy [4]. Silicon and III–V semiconductors have been extensively studied for photoabsorbers in PEC water splitting system [5–7]. Among them, p-type Si is one of the most promising photocathodes due to its earth abundance, advanced infrastructure, high carrier mobility, and suitable band gap, which closely matches the solar spectrum [8, 9]. However, it has the problem of insufficient output photovoltage arisen from its narrow band gap limited by the quasi-Fermi level difference at the interfaces under full illumination [10]. Also, the surface of p-type Si is catalytically inactive due to its high hydrogen adsorption Gibbs free energy ($\Delta G_{\text{H}}$). The other problem is that it suffers from photo-corrosion in aqueous solution because of the thermodynamical reduction potential of silicon adjacent to water reduction potential (H+/H2) [11].

By employing photoelectrochemical catalysts, both the interfacial reactions can be expedited via lowering the activation energy, and the silicon can be protected from photocorrosion [12]. Recently, transition metal phosphides (TMPs) and transition metal dichalcogenides (TMDs) have received tremendous attention because of strong interaction with solar irradiation, tunable optoelectronic properties, and abundant reaction sites for hydrogen evolution reaction (HER) [13–17]. In our previous studies, thickness-optimized 2D MoS2 [18], WS2 [19], and sulfur-doped MoP thin-film catalysts [20] synthesized by a simple in-vacuum thermolysis process exhibited remarkable catalytic activity and stability as PEC catalysts. Also, various transition metal-based catalysts synthesized by a vacuum process, such as NbS2 [21], MoS2/WS2/WSe2 [22], MoQ2Cl2 (Q=S, Se) [23], and CoP [24] have been widely investigated. However, with the high-temperature sulfurization or phosphorization process, which is well-known synthetic methods for highly uniform, scalable, and crystalline transition metal-based catalysts, there is a big issue in the direct synthesis of the catalysts just above the silicon photocathodes because they are chemically incompatible, leading to the formation of secondary compounds such as metal silicides. The inevitably formed secondary phase acts as a parasitic light absorption layer that causes the loss of photon reaching the silicon photocathode. Also, its low shunt resistance reduces catalytic activity by allowing light-generated electrons to move in an alternative path rather than to the surface of the catalyst. For example, Benck et al. [25] directly deposited Mo layers on p-Si using DC magnetron sputter, followed by sulfurization. However, inevitably formed interlayers such as Mo and MoSi induced the parasitic light absorption resulting in the low photocurrent density. Also, Hasani et al. [26] developed a hybrid sulfurization method of depositing MoO3 film on p-Si, spin-coating (NH4)2MoS2 on it, and converting them into MoS2 film. Even though the uniformly distributed and thickness-controlled thin films are synthesized, the photocathodes exhibited leakage current density arising from interfacial secondary phase between silicon and MoS2 [27, 28]. As a result, it is essential to apply a novel method that prevents the formation of metallic secondary phase acting as parasitic light absorption layer and providing inefficient electron path for hydrogen evolution reaction.

Here, we demonstrate the direct growth of highly uniform MoP catalysts on p-Si photocathodes by applying the graphene interlayer at the heterointerface. As graphene completely suppresses the chemical reaction between the silicon and Mo atoms, the Mo-Si layer does not exist. As a result, most of the incident light can be delivered to silicon photocathodes as graphene only absorbs 2.8–4.3% of visible light [29]. Also, no alternative current path between the silicon and catalyst is provided, leading to the absence of leakage current and enhanced catalytic activity. Furthermore, Mo-O-C covalent bondings, which connects graphene and MoP, is favorably formed to accelerate the electron transfer at heterointerface. Moreover, the large work function difference at the graphene/p-Si interface contributes to a strong band bending, facilitating the electron transfer. Based on these advantages of the graphene acting as a bridge between p-Si and MoP, vertically aligned MoP nanorods (NRs) are directly synthesized on silicon photocathode. To make good use of numerous catalytic sites ascribed to P atoms,
we designed a one-dimensional nanostructured MoP with a high surface-to-volume ratio [30, 31]. In addition, drastically enhanced anti-reflectance of nanorod-arrays contributes to the overall light absorption of photocathodes. With an optimized nanorod-arrays thickness of 100 nm, the photocathode exhibits a remarkable photocurrent density of 21.8 mA cm\(^{-2}\) at 0 V versus a reversible hydrogen electrode (RHE) and enhanced long-term stability. While MoS\(_2\) NRs/graphene/p-Si shows a dramatic degradation with increasing pH, MoP NRs/graphene/p-Si was considerably stable to changes in pH. Besides, it maintained high PEC performance with increasing electrolyte temperature up to 70 °C. Using graphene interlayer, not only MoP but also various catalysts such as nickel phosphide and cobalt phosphide can be directly synthesized on silicon photoelectrode. This study demonstrates the huge potential of the graphene to solve the interfacial incompatibility and will contribute to a large-scale direct synthesis of nanostructured transition metal-based materials exhibiting an efficient hydrogen production performance.

2 Experimental

2.1 Synthesis of Graphene

Monolayer graphene film was grown by chemical vapor deposition on Cu foil (Nilaco corporation, #CU-113213, 30 μm thick, 99.9% purity). First, the Cu foil was annealed at 1030 °C for 4 h under flow of H\(_2\) at 70 sccm with a total pressure of 5 Torr; then, monolayer graphene film was grown at 1040 °C for 2 h by additionally introducing 1% CH\(_4\) diluted in H\(_2\) at 2 sccm. During the cooling process after growth, the 1% CH\(_4\) diluted in H\(_2\) gas injection was kept until the film had cooled to 600 °C, then only H\(_2\) was introduced until the temperature reached under 150 °C to allow unloading of the sample.

2.2 Fabrication of MoP NRs/Graphene/p-Si Photocathodes

The boron-doped (100) p-type silicon wafers (1–10 Ω cm) for MoP nanorods(NRs)/graphene(Gr)/p-Si photocathodes and the conductive arsenic-doped (100) n-type silicon wafers (0.001–0.005 Ω cm) for MoP NRs/Gr/n++-Si were cut into pieces with a size of 1.5×1.5 cm\(^2\). The silicon pieces were cleaned using acetone, isopropyl alcohol, and DI water by ultrasonication for 15 min at each step. To remove the native oxide layer, cleaned Si wafers were soaked into HF for 30 s. The PMMA/graphene layer was transferred onto the silicon surface and PMMA was removed by acetone at 60 °C for 15 min. MoP NRs/Gr/p-Si photocathodes were fabricated by MoO\(_3\) precursor (Tae-won Co.) deposition and high-temperature phosphorization process. Vertically aligned molybdenum trioxide nanorods with 100 nm thickness were deposited on graphene/p-Si photocathodes by glancing angle deposition (GLAD) electron beam evaporator. The substrates were tilted at an angle of 80° and rotated with a speed of 80 rpm. The base pressure and growth rates were 1.0×10\(^{-6}\) mTorr and 1.5 Å s\(^{-1}\), respectively. By simple phosphorization process, MoO\(_3\) nanorods turned into MoP nanorods. High-purity H\(_2\) gases mixed with N\(_2\) gases were utilized for this process in a thermal chemical vapor deposition (CVD) system. At first, the temperature of the CVD chamber was increased to 500 °C and maintained for 15 min under the flow of H\(_2\) and N\(_2\) at 0.85 Torr. The flow rate of H\(_2\) and N\(_2\) was 100 and 500 cm\(^3\) min\(^{-1}\), respectively, measured by mass flow controllers. The temperature was increased to 900 °C maintaining for 15 min. The sublimation of phosphorus powder (Sigma-Aldrich, 99.9% purity) proceeded in the other heating zone. The temperature of the sublimation region was set at 400 °C and 0.7 g of powder was used.

2.3 Fabrication of MoP film/Graphene/p-Si and MoP NPs/p-Si Photocathodes

The MoO\(_3\) film with intended thickness of 5 nm was deposited on graphene/p-Si and HF-treated bare p-Si for the fabrication of MoP film/Gr/p-Si and MoP NPs/p-Si, respectively. During deposition, the substrates were kept horizontal without rotation. The base pressure and growth rates were 1.0×10\(^{-6}\) mTorr and 0.1 Å s\(^{-1}\), respectively. After film deposition, the phosphorization process with the temperature of 700 °C in H\(_2\) and N\(_2\) (100 and 500 cm\(^3\) min\(^{-1}\)) atmosphere proceeded for 15 min. The other heating zone where 0.7 g of phosphorus powder sublimates was maintained at 400 °C.
2.4 Fabrication of Various TMPs and TMDs/Graphene/p-Si Photocathodes

The metal oxide films of CuO, WO$_3$, Fe$_2$O$_3$, CoO$_3$, and NiO with intended thickness of 5 nm were deposited on graphene/p-Si for the synthesis of CuP$_x$, WP$_x$, FeP$_x$, CoP$_x$, and NiP$_x$ catalysts. Also, MoO$_3$ and WO$_3$ film with 5 nm thickness were deposited to synthesize MoS$_2$ and WS$_2$ on graphene/p-Si. During the phosphorization and sulfurization process, the temperature of the CVD furnace was maintained at 700 °C for CuP$_x$, CoP$_x$, WS$_2$, and 900 °C for WP$_x$, FeP$_x$, NiP$_x$, MoS$_2$ catalysts. The temperature of the sublimation region was set at 400 °C with 0.7 g of phosphorus powder and 300 °C with 0.7 g of sulfur powder.

2.5 Materials Characterization

The Raman spectra were obtained using Lab RAM HR (Horiba JobinYvon, Japan) with 532 nm excitation wavelength. The FT-IR spectra measurements were conducted with Nicolet iS50 (Thermo Fisher Scientific). The surface morphology and cross-sectional images were acquired by FE-SEM (MERLIN COMPACT, JEISS). Low magnification, high-resolution images, and EDS data were obtained with TEM (JEM-2100F, JEOL). The chemical bonding states and work function of MoP nanorods/graphene/p-Si were characterized by XPS (AXIS SUPRA, Kratos). The crystal structures of MoP film and MoP nanorods were analyzed by XRD (D8 discover, Bruker). The reflectance and absorption spectra were obtained with UV–visible spectroscopy (V-770, JASCO). The electrical properties of the deposited

![Fig. 1](https://example.com/fig1.png)  
**Fig. 1** Schematic illustrations of MoP synthesized on **a** bare p-Si with Mo$_x$Si and **b** graphene passivated p-Si. Linear sweep voltammograms of **c** MoP/p-Si photocathode with photon loss and low catalytic activity and **d** MoP/graphene/p-Si photocathode showing enhanced PEC performance.
catalysts were measured by an Agilent 4156C semiconductor analyzer.

2.6 Photoelectrochemical Measurements

The photoelectrochemical properties of as-synthesized photocathodes were measured with a potentiostat (Ivium Technologies, Nstat) using a three-electrode system consisting of a saturated calomel electrode (SCE) as the reference electrode and a graphite rod as the counter electrode. 0.5 M H₂SO₄ standard solution was used as an electrolyte. A quartz vessel showing high transmittance for UV wavelengths of 200 nm or less was utilized. A Xe arc lamp calibrated to 100 mW cm⁻² (AM 1.5 G condition) was used as a light source. In the process of the linear sweep voltammogram (LSV) measurements, the photocathodes were polarized at a scan rate of 10 mV s⁻¹ under AM 1.5 G. The IPCE data were obtained with a monochromator (MonoRa150) and the potential of 0 V vs. RHE was applied. Electrochemical impedance spectroscopy (EIS) was carried out with the frequency from 250 kHz to 1 Hz using 10 mV alternating current.

3 Results and Discussion

3.1 Graphene Interlayer for Enhanced Light Absorption and Catalytic Activity

In Fig. 1a, b, schematic illustrations of photocathodes with MoP catalysts synthesized on bare p-Si and graphene passivated p-Si are shown, respectively. As shown in Fig. 1a, the MoₓSi layer, a secondary compound of silicon and Mo atoms, is laterally formed between p-Si and MoP catalyst. It induces parasitic light absorption resulting in a reduction of photogenerated charges. In addition, due to its low shunt resistance, some of the light-generated electrons are trapped

![Image](image-url)
in the metallic layer and cannot reach the MoP catalyst. As unintended hydrogen evolution reaction occurs at the secondary phase with no suitable reaction sites, the photocathode shows low catalytic activity and PEC performance. One possibility for hydrogen production to be active in silicide layer may be a spillover effect in which hydrogen species adsorbed to reaction sites with negative $\Delta G_H$ diffuse to the sites with positive $\Delta G_H$ [32]. However, $\Delta G_H$ of $P$ sites, the most catalytic sites of MoP, changes from negative to positive when $H$ coverage increases, which means that as soon as the hydrogen is adsorbed, it is desorbed immediately before the spillover effect occurs [33]. As a result, active proton reduction from the spillover effect cannot occur in silicide layer. On the other hand, as shown in Fig. 1b, both the reduced photon loss and enhanced HER activity can be achieved by graphene interlayer at a heterointerface. Linear sweep voltammograms clearly identify the contribution of graphene to PEC performance of heterostructure photocathodes in Fig. 1c, d. With the direct synthesis of MoP on bare p-Si, the reduction of saturation current density can be observed in Fig. 1c, consistent with photon loss derived from the metallic layer. Also, a certain amount of leakage current with no light illumination clarifies the existence of a conducting layer having low shunt resistance. Because of decreased electrons involved in proton reduction at MoP catalyst, the onset potential of the LSV curve shows a shift toward a cathodic direction. On the contrary, large saturation current density and lowered onset potential with no leakage current can be achieved by the photocathode to which the graphene interlayer is applied in Fig. 1d.

### 3.2 Characterization of MoP Nanorods/Graphene/p-Si Photocathode

Figure 2a shows a schematic illustration of MoP nanorods(NRs)/graphene(Gr)/p-Si photoelectrode fabrication (see the Experimental Section and Fig. S1 for details). Graphene having tightly packed carbon atoms with an sp$^2$-hybridized lattice is grown on copper foil with large grain size (Fig. S2a) [34]. It is transferred to the p-Si substrate and completely covers the surface of silicon (Fig. S2b). Using electron beam evaporator, MoO$_3$ nanorods are deposited on graphene with a thickness of 100 nm. MoP nanorod-array catalysts are finally synthesized on graphene passivated silicon wafer by vapor phase phosphorization of MoO$_3$ precursor. As for the comparative group, MoP nanoparticles and MoP film are synthesized on p-Si and graphene/p-Si, respectively (see the Experimental Section for details). The photographic images of centimeter-scale Graphene/p-Si and MoP NRs/Gr/p-Si is represented in Fig. S3a, b. It indicates that the graphene monolayer is uniformly transferred to p-Si substrates, and MoP nanorod-arrays are successfully grown on graphene.

The vertically aligned nanorod structure is identified by SEM images. In Fig. 2b, the top-view SEM image of MoP nanorods is shown. It is confirmed that each MoP nanorod has a strictly uniform diameter (~20 nm) and is well-separated as glancing angle deposition (GLAD) induces a self-shadowing effect leading to homogeneous MoP nucleation [35]. The morphologies of MoP nanoparticles(NPs)/p-Si and MoP film/Gr/p-Si are shown in Fig. S4. As we can see in Fig. 2c, the thickness of the MoP NRs layer is 100 nm, which is quite thick compared to the thin-film catalysts. Even though a certain amount of incident light can be interrupted, it is compensated by the anti-reflectance of nanorod-arrays leading to no significant decrease in the overall amount of light absorption [36]. It will be identified by the reflectance vs. wavelength spectra. Meanwhile, what we aimed to achieve from nanorod structure is a high surface-to-volume ratio. In other words, by increasing the catalytically active area with a structural approach [37], the photoelectrochemical performance of MoP NRs/Gr/p-Si photocathode is enhanced compared to that of MoP film/Gr/p-Si photocathode. As MoP has numerous active sites ascribed to P atoms exhibiting low $\Delta G_H$ and acting as “$H$ delivery” [33, 38], increasing the surface area of MoP is a key strategy to obtain a high photocurrent density. The constituent elements and atomic structure of the synthesized MoP are investigated by transmission electron microscopy (TEM). In Fig. 2d, e, uniformly distributed Mo and P elements with Si are shown by scanning TEM (STEM) image and the EDS elemental mapping images. As shown in Fig. 2f, a high-resolution TEM image (HR-TEM) and fast Fourier transformation (FFT) pattern indicate the presence of highly-crystallized MoP nanorod. The clear and well-defined lattice fringes with a d-spacing of 0.278 nm and corresponding fast Fourier transform are consistent with the (100) plane for hexagonal MoP [39]. The high-resolution cross-sectional TEM image in Fig. 2g represents that MoP nanorod with high-crystallinity is formed just above the graphene-passivated silicon
substrate. In Fig. S5, the existence of graphene interlayer can be identified by zoom-in cross-sectional HR-TEM image. All of the MoO₃ precursors are successfully transformed into MoP catalysts without remaining MoO₃ layers. Moreover, the formation of MoₓSi layers is prevented by using graphene passivation.

The surface constituents and chemical nature of MoP nanorods and film are further investigated by X-ray photoelectron spectroscopy (XPS). The full XPS spectra show the existence of Mo, P, O, and C (Fig. S6a). The XPS profiles of Mo 3d and P 2p of both MoP film and MoP nanorods are presented in Fig. S6b and c. Two pairs of doublets at 231.5 eV/228.4 eV (Mo³⁺ 3d₅/₂/Mo³⁺ 3d₃/₂) and 130.5 eV/129.6 eV (P 2p½/P 2p¾) are ascribed to MoP [38], while the rest of the peaks corresponds to MoO₃, MoO₂, and PO₄³⁻ caused by surface oxidation [33].

In Fig. 3a, X-ray diffraction (XRD) patterns of crystalline MoP film and nanorod-arrays on graphene/p-Si are provided. Characteristic peaks of MoP film are observed at 28° and 58°, which correspond to the (001) and (110) facets of MoP, respectively [33]. In the case of MoP nanorods, an additional peak at 32° corresponding to the (100) facets is identified, matching the result of the HR-TEM image. In both samples, the peak corresponding to MoO₃ phase is not observed. The optical reflectance and absorbance of bare p-Si, MoP film/Gr/p-Si, and MoP NRs/Gr/p-Si photocathodes in the wavelength range from 300 to 900 nm are shown in Figs. 3b and S7a. MoP film grown on graphene/p-Si exhibits about 5% higher reflectance than the bare silicon in all wavelengths. However, even the same materials, the reflectance values of nanorod-structured MoP are much lower than that of film structure. It has a reflectance of 30% at a wavelength of 900 nm, and the value becomes smaller as the wavelength decreases, showing about 6% reflectance at a wavelength of 300 nm. Due to the porous nature of the material, MoP nanorod-arrays have a lower refractive index weakening broadband light reflectance [40, 41]. Such an outperforming anti-reflectance of MoP NRs/Gr/p-Si photocathode can contribute to the enhanced solar energy conversion efficiency. For MoP NPs/p-Si photocathode in Fig. S7b, though there is photon loss induced by metallic MoₓSi layer, it seems that MoP nanoparticles significantly affect light absorption and there is no big difference in the reflectance compared to MoP film/Gr/p-Si.

As the graphene is a pathway through which photogenerated electrons must pass until they reach the electrolyte, it is necessary to identify how the graphene chemically and electrically interacts with MoP and silicon. At first, to characterize the chemical interaction between the graphene and MoP, Raman and FT-IR spectra were investigated in each of the three steps where the MoP NRs/Gr/p-Si photocathode is fabricated. In Fig. 4a, Raman spectra show three distinct peaks of D peak at 1350 cm⁻¹, G peak at 1590 cm⁻¹, and 2D peak at 2680 cm⁻¹, which originated from the out of plane vibrations, the in-plane vibrations of sp²-bonded carbon atoms, and the second-order two phonon process, respectively [42]. As for Gr/p-Si and MoO₃ NRs/Gr/p-Si samples, typical I_D/I_2D peak ratio of monolayer graphene and the negligible D peaks are observed, implying a well-progressed graphene transfer and MoO₃ precursor deposition.
However, after phosphorization, the intensity of D peak is drastically increased while that of 2D peak is decreased. It is well known that defects are required to activate D peak and deactivate 2D peak of graphene [43]. Specifically, $sp^3$-type defects are likely introduced since D’ peak is not visible [44]. Among some kinds of $sp^3$-type defects, it is most likely that oxygen-containing functional groups are formed by receiving oxygen from the MoO$_3$ precursor as thermal energy was applied to the interface between graphene and MoO$_3$ during the heating process. To clearly verify that this Raman spectrum is derived from the bonding with oxygen, not affected by H$_2$ annealing or gas phase phosphorus, Raman spectra of Gr/p-Si and MoO$_3$ NRs/Gr/p-Si photo-cathodes after annealing at 900 °C in H$_2$/N$_2$ atmosphere are shown in Fig. S8a and b, respectively. For Gr/p-Si photo-cathode after annealing, $I_D/I_{2D}$ peak ratio increased, which attributes to the generation of defects caused by hydrogen etching [45]. However, $I_D/I_{2D}$ peak ratio is still small which is not appropriate to explain the Raman spectra of MoP NRs/Gr/p-Si. In Fig. S8b, although there is no flowing of phosphorus during the annealing, the same Raman spectrum with MoP NRs/Gr/p-Si was obtained. From this result, it is confirmed that interfacial bonding between graphene and MoO$_3$ are formed due to high thermal energy.

Fig. 4  a Raman spectra of graphene when transferred to a p-Si wafer, covered with MoO$_3$ nanorods precursor, and covered with MoP nanorods catalysts. b FT-IR spectra. c Ultraviolet photoemission spectroscopy (UPS) as a function of binding energy for the work function of p-Si, graphene/p-Si and MoP NRs/graphene/p-Si. d Schematic of the energy band diagram showing favorably formed band bending between p-Si and graphene.
To further investigate the presence of the functional group in graphene, the samples are analyzed by FT-IR spectroscopy in Fig. 4b. Unlike graphene oxide and reduced graphene oxide, monolayer graphene shows no C=C aromatic stretching [46]. As for MoO3 NRs/Gr/p-Si sample, the bands at 943 cm\(^{-1}\) and 910 cm\(^{-1}\) are apparent, attributed to Mo=O characteristic vibrations of the orthorhombic phase MoO\(_3\) [47, 48]. After phosphorization, with the MoP NRs/Gr/p-Si sample, MoO\(_3\) precursor has been completely converted to MoP as can be seen from the disappearance of Mo=O peaks and the most discernible band appears at 1242 cm\(^{-1}\) corresponding to C–O epoxy stretching [49–51]. The peaks at 1422 and 1625 cm\(^{-1}\) are related to the vibration of the Mo-OH bond and the bending of adsorbed water, respectively [52, 53]. Judging from the existence of C–O vibration, it appears that oxygen is bound to graphene interlayer and in the interface between graphene and MoP NRs. Considering that flowing hydrogen gas for phosphorization induces graphene etching and the broken C=C bonds appear with destroyed carbon lattice [45], it is likely that the broken bonds in lattice function as reactive sites and contribute to covalent bondings such as Mo-O-C under high-temperature condition [54]. Consequently, as strong chemical interaction can be achieved by covalent bondings, the photogenerated electrons transport easily at the graphene/MoP NRs interface with low interfacial charge resistance, enhancing the photoelectrochemical performance.

Secondly, the electrical interaction between graphene and silicon photocathode is analyzed using ultraviolet photoelectron spectroscopy (UPS) to demonstrate the band alignment of heterostructure in hydrogen production. In Fig. 4c, the secondary electron emission (SEE) cutoffs as a function of binding energy are shown for the work function of p-Si (4.65 eV), graphene/p-Si (4.2 eV), and MoP NRs/Gr/p-Si (4.8 eV). As the graphene and MoP show metallic properties, the energy band diagrams for the MoP NRs/Gr/p-Si heterostructure are described in Fig. 4d. The large work function difference of 0.45 eV at the graphene/p-Si interface induces a strong band bending, which allows electrons in the conduction band minimum of p-Si to move rapidly through the graphene interlayer. For the detailed explanation of band bending at graphene/p-Si contact, band diagrams for the graphene/p-Si contact in three cases (before equilibrium, in equilibrium, in steady-state illumination) are provided in Fig. S9.

### 3.3 Photoelectrochemical Hydrogen Evolution Reaction

The PEC characterization of MoP NRs/Gr/p-Si photocathode compared to bare p-Si, graphene/p-Si, MoS\(_2\) NRs/Gr/p-Si, MoP NPs/p-Si, and MoP film/Gr/p-Si is carried out under a simulated air mass 1.5 G condition with a standard three-electrode setup using 0.5 M H\(_2\)SO\(_4\) electrolyte. The graphite rod is used as a counter electrode. As shown in current density versus potential (J-V) curves in Fig. 5a, the bare p-Si photocathode shows a large negative onset potential because of its catalytic inactivity despite the high electron mobility and suitable bandgap. When MoP nanoparticles are applied to p-Si, catalytic activity is enhanced, resulting in the onset potential shift towards the anodic direction. The onset potential and the photocurrent density at 0 V versus RHE reach the values of 0.08 V versus RHE and 4.8 mA cm\(^{-2}\), respectively. However, from potential above −0.4 V versus RHE, a discernible leakage current is shown due to a low shunt resistance caused by MoSi layer between the crystalline silicon and MoP nanoparticles [55]. Though no leakage current was observed in MoS\(_2\)/Mo/MoSi/n+p-Si photocathode fabricated by Benck et al. [25], it is clearly due to the low synthesis temperature of 250 °C. As shown in Fig. S10, the LSV curves of MoP/p-Si obtained at 250 and 900 °C evidently show the temperature dependence of leakage current. In Fig. S10a, when synthesized at 250 °C, a negligible leakage current is observed, whereas it increases dramatically as the temperature rises to 900 °C in Fig. S10b.

In this respect, by applying the graphene interlayer, we can completely suppress the formation of the metallic secondary phase as seen that no leakage current is visible with MoP film/Gr/p-Si and MoP NRs/Gr/p-Si photocathodes. In Fig. S11, resistance analysis clearly shows that the leakage current flowing laterally through the silicide layers can be effectively suppressed by applying graphene interlayer. Though the graphene shows no remarkable PEC performance as catalysts, MoP film/Gr/p-Si and MoP NRs/Gr/p-Si photocathodes show the dramatic improvement in PEC performance. Mainly, MoP NRs/Gr/p-Si photocathode exhibits a remarkable catalytic activity for the HER with an onset potential of 0.24 V vs. RHE to reach -1 mA cm\(^{-2}\) and a photocurrent density of 21.8 mA cm\(^{-2}\) at 0 V versus RHE. The 100 nm thickness of MoP nanorod-arrays is an optimized value to get maximum photocurrent density at 0 V versus RHE. Figure S12 indicates that the photocathode with 100 nm MoP NRs shows the highest photocurrent density at 0 V versus RHE compared to the photocathodes with 70 and 130 nm MoP NRs catalysts.
Electrochemical impedance spectroscopy (EIS) measurements were conducted to understand the PEC performance concerning charge transfer resistance, as shown in Fig. 5b. The obtained EIS data were fitted to a simplified equivalent circuit. The equivalent circuit is composed of charge transfer resistance ($R_{ct}$) and constant phase elements (CPEs). The smaller arcs in Nyquist plots indicate the lower charge transfer resistance at each interface. Compared to each semicircle of MoP NPs/p-Si photocathode, the photocathodes to which the graphene interlayer is applied show smaller arcs. Notably, the resistances of MoP NRs/Gr/p-Si photocathode at each interface are remarkably decreased.

The exact values of fitted charge transfer resistance corresponding to each interface are tabulated in Table 1. The
The summarized charge transport resistance values of bare p-Si, MoP NPs/p-Si, MoP film/Gr/p-Si, and MoP NRs/Gr/p-Si photocathodes.

The capability of photocathodes to convert the incident photons to electrical current is investigated by the incident photon-to-current conversion efficiency (IPCE) at an applied bias of 0 V versus RHE, as shown in Fig. 5c. The MoP NRs/Gr/p-Si photocathode shows the efficiency of ~68% over the entire visible light wavelength, while the MoP film/Gr/p-Si photocathode exhibits the IPCE value of ~33% which elucidates that a high quantum efficiency can be attained by nano-structured MoP catalysts.

The stability of bare p-Si, MoP NPs/p-Si, MoP film/Gr/p-Si, and MoP NRs/Gr/p-Si photocathodes are evaluated by chronoaaperometric measurements in Fig. 6. These measurements are conducted to determine whether MoP catalysts constructed on graphene can act as a passivation layer and maintain their activity without deformation. A critical degradation is shown with a bare p-Si photocathode. Also, in the case of MoP nanoparticles, the photocurrent density of ~4.8 mA cm\(^{-2}\) at 0 V versus RHE is not endured 80% of photocurrent density until 1 h. As the catalysts do not fully cover the silicon surface, the formation of an insulating layer is accelerated, hindering charge transfer and finally shutting down the overall activity. On the other hand, it is found that the long-term stability of over 10 h is maintained with the MoP film and MoP nanorod-arrays directly synthesized on graphene. For the MoP NRs/Gr/p-Si photocathode, significantly, the photocurrent degradation is considerably suppressed with retaining over 88% of the performance for 10 h.

Table 1

| Photocathode       | Resistance | \(R_{c,t,1}\) (\(\Omega \text{ cm}^2\)) | \(R_{c,t,2}\) (\(\Omega \text{ cm}^2\)) | \(R_{c,t,3}\) (\(\Omega \text{ cm}^2\)) |
|--------------------|------------|-------------------------------------|-------------------------------------|-------------------------------------|
| Bare p-Si          | 1.60       | 3131.13 (p-Si→electrolyte)          | N/A                                 | N/A                                 |
| MoP NPs            | 1.46       | 16.48 (p-Si→MoP NPs)               | 44.34 (MoP NPs→electrolyte)         | N/A                                 |
| MoP film/Gr        | 1.47       | 5.27 (p-Si→Graphene)               | 5.47 (Graphene→MoP film)            | 4.75 (MoP film→electrolyte)         |
| MoP NRs/Gr         | 1.12       | 5.34 (p-Si→Graphene)               | 5.45 (Graphene→MoP NRs)             | 3.68 (MoP NRs→electrolyte)          |

Fig. 6 Long-term stability tests for the bare p-Si, MoP NPs/p-Si, MoP film/Gr/p-Si, and MoP NRs/Gr/p-Si photocathodes at 0 V versus RHE and Faradaic efficiency for MoP NRs/Gr/p-Si photocathode

\(R_{c,t,1}\) of bare p-Si shows two orders of magnitude higher than the total resistance of other MoP deposited photocathodes. It reflects that molybdenum phosphate enables the electrons to move faster at the interfaces and provides a large number of active sites for energetic kinetics. For the resistances at MoP/electrolyte interface, \(R_{c,t,1}\) of MoP NRs/Gr/p-Si photocathode exhibits the lowest value compared to \(R_{c,t,2}\) of MoP film/Gr/p-Si and \(R_{c,t,3}\) of MoP NPs/p-Si photocathodes. It confirms that for the nanorod-structured MoP, electron transfer to the redox couples is enhanced, which is ascribed to the highly-exposed 1D structure showing the shortest distance towards MoP/electrolyte interface and the largest amount of active sites. Moreover, to identify the synergistic effects of graphene, we compared the resistances at heterointerfaces with and without graphene. The sum of \(R_{c,t,1}\) and \(R_{c,t,2}\) of both the graphene-passivated photocathodes is lower than \(R_{c,t,1}\) of MoP NPs/p-Si photocathode, which provides compelling evidence that the graphene can act as a highway to help the electrons move from p-Si to MoP catalysts. This result is in good agreement with the formation of Mo-O-C covalent bondings and energetically favorable band bending identified by Raman, FT-IR spectroscopy, and UPS.
As shown in Fig. S13, the SEM image reveals that MoP nanorod-arrays show no structural change after a 10-h stability test. Interestingly, it is estimated that the total amount of hydrogen generated at the wafer scale for 10 h is apparently larger than that required for the synthesis of MoP nanorod catalysts, but much lower than that required for the synthesis of graphene in Fig. S13.

The Faradaic efficiency is obtained by conducting a gas chromatography (GC) measurement that collects the evolved hydrogen gas. In Fig. 6, the MoP NRs/Gr/p-Si photocathode shows almost 100% Faradaic yield during the photoelectrochemical process.

To compare the PEC performance of MoP NRs/Gr/p-Si photocathode with that of state-of-the-art photocathodes synthesized by the in-vacuum high-temperature process, photocurrent density at 0 V versus RHE and long-term stability are plotted in Fig. 5d. Most of the photocathodes [22, 24–26, 56] provide a lower photocurrent density at 0 V versus RHE than our photocathode though some of them maintain their catalytic activity for a long time. In contrast, NbS₂/p-Si NWs [21] and MoSₓClₓ/n⁺pp⁺-Si MPs [23] photocathodes show significantly enhanced photocurrent density while it does not last more than 2~3 h. However, our photocathode shows a superior PEC performance in balance with photocurrent density and stability. The details about the synthesis method and numerical values are tabulated in Table S1.

For further practical use of MoP NRs/Gr/p-Si photocathode as part of PEC cell, it needs to be operated with high reliability against variations in the pH and temperature of electrolyte. In Fig. 5e, 2D plots show how the current density of MoP NRs/Gr/p-Si and MoS₂ NRs/Gr/p-Si photocathodes changes at each voltage as the pH changes from 0.26 to 2. The pH of the electrolyte is controlled by mixing 0.5 M H₂SO₄ with neutral DI water. In the case of MoS₂ nanorod-array catalysts (inset), considering the contours with the current density of 1, 10, and 20 mA cm⁻², the potential required to obtain each current density value dramatically increases as the acidity of electrolyte becomes weaker. On the other hand, the MoP nanorod-arrays catalysts show no definite change up to pH 1.2, and after that, the PEC performance gradually weakened. Similarly, the current density variation depending on electrolyte temperature is plotted in Fig. 5f. As for the temperature effects on the photoelectrodes, it is well known that a net decrease of efficiency is observed with increasing temperature as the reduction in photovoltage induced by radiation losses surpasses an enhancement of interfacial kinetics [57]. However, as shown in Fig. 5f, both samples show high stability against the increase of electrolyte temperature from 30 to 70 °C. It appears that both the incorporated photoelectrochemical catalysts boost the surface reaction enough to alleviate the reduction in the absorber efficiency.

We also investigated how the nanorod structure, which caused a significant increase in PEC performance of MoP having a 3D atomic structure, affects that of MoS₂ having a 2D atomic structure. MoS₂ nanorod-arrays and film are synthesized in the same process with MoP nanorod-arrays and MoP film, respectively, except for using sulfur powder. The Raman and XRD spectrums of MoS₂ nanorods are provided.

![Fig. 7](https://doi.org/10.1007/s40820-021-00605-7)
in Fig. S14a, b. As shown in Fig. S14c, there is a noticeable degradation in PEC performance when the nanorod structure is applied to MoS$_2$. Besides, it does not maintain its activity for a long time (Fig. S14d). Considering that MoS$_2$ has a layered crystal structure held by weak van der Waals force [13], the photogenerated electrons should move through continuous hopping transport at each layer, which causes a large overpotential. Though the nanorod structure contributes to a numerical increment of active sites, large overpotential hinders an efficient hydrogen evolution reaction.

The electrochemical (EC) HER performances of MoP-based catalysts on graphene/p-Si are also investigated in Fig. S15 and Table S2. To achieve graphene-based direct growth on silicon, highly doped n$^{++}$-Si wafers are used as metallic cathodes. In Fig. S15a, MoP nanorod-array catalysts exhibit a smaller onset potential and overpotential at a current density of 10 mA cm$^{-2}$ compared to MoP film. Also, the Tafel slope for MoP NRs/Gr/n$^{++}$-Si is 97.19 mV per decade, which is lower than the values of 108.99 mV per decade for MoP film/Gr/n$^{++}$-Si, as shown in Fig. S15b. From these results, a larger amount of surface active sites is exposed and faster catalytic activity occurs for MoP nanorods than film electrocatalyst. Since the thickness of our MoP nanorods catalysts is optimized for the PEC system, an enhanced electrocatalytic performance can be obtained by increasing the thickness in the EC system.

To further demonstrate that the synergistic direct growth on graphene interlayer can be applied to various TMPs and TMDs, we have synthesized Cu$_x$P$_y$, WP$_x$, FeP$_x$, CoP$_x$, NiP$_x$, MoS$_2$, and WS$_2$ on graphene/p-Si substrates. They were prepared by phosphorization of CuO, WO$_3$, Fe$_2$O$_3$, CoO$_3$, NiO, and sulfurization of MoO$_3$, WO$_3$. The synthesis temperature of CuP$_x$, CoP$_x$, and WS$_2$ is 700 °C, while that of WP$_x$, FeP$_x$, NiP$_x$, and MoS$_2$ is 900 °C. In Fig. 7a, the current density vs. potential (J-V) curves of TMPs/Gr/p-Si photocathodes under a simulated air mass 1.5 G condition are shown. For all the TMPs, the onset potentials are shifted toward the anodic direction compared to bare p-Si, indicating that the various TMPs work as efficient PEC catalysts on graphene/p-Si photocathodes. Among them, NiP$_x$ and CoP$_x$ exhibit a remarkable current density of 12.9 and 6.1 mA cm$^{-2}$ at 0 V vs. RHE, respectively. Also, Fig. 7b presents the current density vs. potential (J-V) curves of TMDs/Gr/p-Si photocathodes. Similarly, the onset potentials of MoS$_2$/Gr/p-Si and WS$_2$/Gr/p-Si photocathodes are shifted toward the anodic direction. If the thickness control or nanostructuring is achieved with these catalysts, the enhanced photoelectrochemical activity will be obtained.

4 Conclusions

We successfully demonstrated that MoP nanorod-array catalysts grown on graphene-passivated silicon photocathodes remarkably improve PEC-HER performance in balance with the high photocurrent density of 21.8 mA cm$^{-2}$ at 0 V versus RHE and long-term stability of over 10 h. Through the graphene passivation process, not only the secondary compound which absorbs a parasitic light and provides an inefficient electron path are prevented, but also the synergistic effects of Mo-O-C bondings and strong band bending enhanced the photogenerated electron transport. Benefiting from a large surface area exposing numerous active sites and anti-reflectance of MoP nanorods, the obtained photocathodes exhibited a high quantum efficiency. Furthermore, its high performance was maintained with the increase of pH and temperature. This study provides the framework to rationally design nanostructured PEC catalysts directly on graphene/p-Si photocathodes with high-quality interfaces and superior HER performance.

Acknowledgements This research was financially supported by the Basic Science Research Program (2017R1A2B3009135), the Korean government MSIT (2019M3E6A1103818), the Basic Research Laboratory Program (2018R1A4A1022647), and the Future Material Discovery Program (2018M3D1A1058793) through the National Research Foundation of Korea. H.W.J. acknowledges KOREA HYDRO & NUCLEAR POWER CO., LTD. (No. 2018-Tech-21).

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain
Supplementary Information

The online version contains supplementary material available at (https://doi.org/10.1007/s40820-021-00605-7).

References

1. W. Lubitz, W. Tumas, Hydrogen: an overview. Chem. Rev. 107, 3900 (2007). https://doi.org/10.1021/cr050200z
2. N. Armaroli, V. Balzani, Solar electricity and solar fuels: status and perspectives in the context of the energy transition. Chem. Eur. J. 22, 32–57 (2016). https://doi.org/10.1002/chem.201503580
3. X. Li, L. Zhao, J. Yu, X. Liu, X. Zhang et al., Water splitting: from electrode to green energy system. Nano-Micro Lett. 12, 131 (2020). https://doi.org/10.1007/s40820-020-00469-3
4. I. Roger, M.A. Shipman, M.D. Symes, Earth-abundant catalysts for electrochemical and photoelectrochemical water splitting. Nat. Rev. Chem. 1, 1–13 (2017). https://doi.org/10.1038/s41570-016-0003
5. S.A. Lee, T.H. Lee, C. Kim, M.G. Lee, M.-J. Choi et al., Tailored NiOx/Ni cocatalysts on silicon for highly efficient water splitting photoanodes via pulsed electrodeposition. ACS Catal. 8, 7261–7269 (2018). https://doi.org/10.1021/acscatal.8b01999
6. R.J. Britto, J.D. Benck, J.L. Young, C. Hahn, T.G. Deutsch et al., Molybdenum disulfide as a protection layer and catalyst for gallium indium phosphide solar water splitting photocathodes. J. Phys. Chem. Lett. 7, 2044–2049 (2016). https://doi.org/10.1021/acs.jpclett.6b00563
7. D. Kang, J.L. Young, H. Lim, W.E. Klein, H. Chen et al., Printed assemblies of GaAs photoelectrodes with decoupled optical and reactive interfaces for unassisted solar water splitting. Nat. Energy 2, 17043 (2017). https://doi.org/10.1038/nenergy.2017.43
8. O. Gunawan, L. Sekaric, A. Majumdar, M. Rooks, J. Appenzeller et al., Measurement of carrier mobility in silicon nanowires. Nano Lett. 8, 1566–1571 (2008). https://doi.org/10.1021/nl072646w
9. C. Strümpfel, M. McCann, G. Beaucaurne, V. Arkhipov, A. Slaoui et al., Modifying the solar spectrum to enhance silicon solar cell efficiency—An overview of available materials. Sol. Energy Mater. Sol. Cells 91, 238–249 (2007). https://doi.org/10.1016/j.solmat.2006.09.003
10. M.G. Walter, E.L. Warren, J.R. McKone, S.W. Boettcher, Q. Mi et al., Solar water splitting cells. Chem. Rev. 110, 6446–6473 (2010). https://doi.org/10.1021/cr1002326
11. S. Chen, L.-W. Wang, Thermodynamic oxidation and reduction potentials of photocatalytic semiconductors in aqueous solution. Chem. Mater. 24, 3659–3666 (2012). https://doi.org/10.1021/cm302533s
12. C.-J. Chen, K.-C. Yang, C.-W. Liu, Y.-R. Lu, C.-L. Dong et al., Silicon microwire arrays decorated with amorphous hetero-metal-doped molybdenum sulfide for water photoelectrolysis. Nano Energy 32, 422–432 (2017). https://doi.org/10.1016/j.nanoen.2016.12.045
13. G. Li, D. Zhang, Q. Qiao, Y. Yu, D. Peterson et al., All the catalytic active sites of MoS2 for hydrogen evolution. J. Am. Chem. Soc. 138, 16632–16638 (2016). https://doi.org/10.1021/jacs.6b05940
14. S. Zheng, L. Zheng, Z. Zhu, J. Chen, J. Kang et al., MoS2 nanosheet arrays rooted on hollow rGO spheres as bifunctional hydrogen evolution catalyst and supercapacitor electrode. Nano-Micro Lett. 10, 62 (2018). https://doi.org/10.1007/s40820-018-0215-3
15. M.-L. Tsai, S.-H. Su, J.-K. Chang, D.-S. Tsai, C.-H. Chen et al., Monolayer MoS2 heterojunction solar cells. ACS Nano 8, 8317–8322 (2014). https://doi.org/10.1021/nn502776h
16. T. Chu, H. Ilitikhameneh, G. Klimeck, R. Rahman, Z. Chen, Electrically tunable bandgaps in bilayer MoS2, Nano Lett. 15, 8000–8007 (2015). https://doi.org/10.1021/acs.nanolett.5b03218
17. J. Lin, Y. Yan, C. Li, X. Si, H. Wang et al., Bifunctional electrocatalysts based on Mo-doped NiCoP nanosheet arrays for overall water splitting. Nano-Micro Lett. 11, 55 (2019). https://doi.org/10.1007/s40820-019-0289-6
18. K.C. Kwon, S. Choi, K. Hong, C.W. Moon, Y.-S. Shim et al., Wafer-scale transferable molybdenum disulfide thin-film catalysts for photoelectrochemical hydrogen production. Energy Environ. Sci. 9, 2240–2248 (2016). https://doi.org/10.1039/C6EE00144K
19. K.C. Kwon, S. Choi, K. Hong, D.M. Andoshe, J.M. Suh et al., Tungsten disulfide thin film/p-type Si heterojunction photocathode for efficient photochemical hydrogen production. MRS Commun. 7, 272 (2017). https://doi.org/10.1557/mrc.2017.37
20. K.C. Kwon, S. Choi, J. Lee, K. Hong, W. Sohn et al., Drastically enhanced hydrogen evolution activity by 2D to 3D structural transition in anion-engineered molybdenum disulfide thin films for efficient Si-based water splitting photocathodes. J. Mater. Chem. A 5, 15534–15542 (2017). https://doi.org/10.1039/C7TA03845C
21. P. Gnanasekar, D. Periyangounder, P. Varadhan, J.-H. He, J. Kulandaivel, Highly efficient and stable photoelectrochemical hydrogen evolution with 2D-NbS2/Si nanowire heterojunction. ACS Appl. Mater. Interfaces 11, 44179–44185 (2019). https://doi.org/10.1021/acsami.9b14713
22. S. Seo, S. Kim, H. Choi, J. Lee, H. Yoon et al., Direct in situ growth of centimeter-scale multi-heterojunction MoS2/WSe2 thin-film catalyst for photo-electrochemical hydrogen evolution. Adv. Sci. 6, 1900301 (2019). https://doi.org/10.1002/advs.201900301
23. Q. Ding, J. Zhai, M. Cabán-Acevedo, M.J. Shearer, L. Li et al., Designing efficient solar-driven hydrogen evolution photocathodes using semitransparent MoQxCl (Q= S, Se) catalysts on Si Micropyramids. Adv. Mater. 27, 6511–6518 (2015). https://doi.org/10.1002/adma.201501884
24. T.R. Hellstern, J.D. Benck, J. Kibsgaard, C. Hahn, T.F. Jaraszillo, Engineering cobalt phosphate (CoP) thin film catalysts for enhanced hydrogen evolution activity on silicon photocathodes. Adv. Energy Mater. 6, 1501758 (2016). https://doi.org/10.1002/aenm.201501758

25. J.D. Benck, S.C. Lee, K.D. Fong, J. Kibsgaard, R. Sinclair et al., Designing active and stable silicon photocathodes for solar hydrogen production using molybdenum sulfide nanomaterials. Adv. Energy Mater. 4, 1400739 (2014). https://doi.org/10.1002/aenm.201400739

26. A. Hasani, Q. Van Le, M. Tekalgon, M.-J. Choi, T.H. Lee et al., Direct synthesis of two-dimensional MoS2 on p-type Si and application to solar hydrogen production. NPG Asia Mater. 11, 1–9 (2019). https://doi.org/10.1038/s41427-019-0145-7

27. R. Liu, D. Williams, W. Lynch, A study of the leakage mechanisms of silicided n+/p junctions. J. Appl. Phys. 63, 1990–1999 (1988). https://doi.org/10.1063/1.341099

28. P. Sharma, T. Kaur, O.P. Pandey, In situ single-step reduction and silicidation of MoO3 to form MoSi2. J. Am. Ceram. Soc. 102, 1522–1534 (2019). https://doi.org/10.1111/jace.15994

29. K.F. Mak, J. Shan, T.F. Heinz. Seeing many-body effects in single-and few-layer graphene: observation of two-dimensional saddle-point excitons. Phys. Rev. Lett. 106, 046401 (2011). https://doi.org/10.1103/PhysRevLett.106.046401

30. E. Samuel, B. Joshi, M.-W. Kim, M.T. Swihart, S.S. Yoon, Morphology engineering of photoelectrodes for efficient photoelectrochemical water splitting. Nano Energy (2020). https://doi.org/10.1016/j.nanoen.2020.104648

31. J. Li, W. Xu, J. Luo, D. Zhou, D. Zhang et al., Synthesis of 3D hexagram-like cobalt-manganese sulfides nanosheets grown on nickel foam: a bifunctional electrocatalyst for overall water splitting. Nano-Micro Lett. 10, 6 (2018). https://doi.org/10.1007/s40820-017-0160-6

32. J. Li, H.-X. Liu, W. Gou, M. Zhang, Z. Xia et al., Ethylene glycol ligand environment facilitates highly efficient hydrogen evolution of Pt/CdP through proton concentration and hydrogen spillover. Energy Environ. Sci. 12, 2298–2304 (2019). https://doi.org/10.1039/C9EE00752K

33. P. Xiao, M.A. Sk, L. Thia, X. Ge, R.J. Lim et al., Molybdenum phosphate as an efficient electrocatalyst for the hydrogen evolution reaction. Energy Environ. Sci. 7, 2624–2629 (2014). https://doi.org/10.1039/C4EE00957F

34. D. Zhan, J. Yan, L. Lai, Z. Ni, L. Liu et al., Engineering the electronic structure of graphene. Adv. Mater. 24, 4055–4069 (2012). https://doi.org/10.1002/adma.201200011

35. M.G. Lee, W. Sohn, C.W. Moon, H. Park, S. Lee et al., Conformally coated BiVO4 nanodots on porosity-controlled WO3 nanorods as highly efficient type II heterojunction photoanodes for water oxidation. Nano Energy 28, 250–260 (2016). https://doi.org/10.1016/j.nanoen.2016.08.046

36. D.M. Ando, S. Choi, Y.-S. Shim, S.H. Lee, Y. Kim et al., A wafer-scale antireflection protection layer of solution-processed TiO2 nanorods for high performance silicon-based water splitting photocathodes. J. Mater. Chem. A 4, 9477–9485 (2016). https://doi.org/10.1039/C6TA02987F

37. S. Deng, K. Zhang, D. Xie, Y. Zhang, Y. Zhang et al., High-index-faceted Ni3S2 branch arrays as bifunctional electrocatalysts for efficient water splitting. Nano-Micro Lett. 11, 12 (2019). https://doi.org/10.1007/s40820-019-01242-8

38. R. Ye, P. del Angel-Vicente, Y. Liu, M.J. Arellano-Jimenez, Z. Peng et al., High-performance hydrogen evolution from MoS21−xPx solid solution. Adv. Mater. 28, 1427–1432 (2016). https://doi.org/10.1002/adma.201504866

39. Z. Xing, Q. Liu, A.M. Asiri, X. Sun, Closely interconnected network of molybdenum phosphide nanoparticles: a highly efficient electrocatalyst for generating hydrogen from water. Adv. Mater. 26, 5702–5707 (2014). https://doi.org/10.1002/adma.201401692

40. M.L. Brongersma, Y. Cui, S. Fan, Light management for photovoltaics using high-index nanostructures. Nat. Mater. 13, 451–460 (2014). https://doi.org/10.1038/nmat3921

41. J.-Q. Xi, M.F. Schubert, J.K. Kim, E.F. Schubert, M. Chen et al., Optical thin-film materials with low refractive index for broadband elimination of Fresnel reflection. Nat. Photon. 1, 176–179 (2007). https://doi.org/10.1038/nphoton.2007.26

42. U. Sim, T.-Y. Yang, J. Moon, J. An, J. Hwang et al., N-doped monolayer graphene catalyst on silicon photocathode for hydrogen production. Energy Environ. Sci. 6, 3658–3664 (2013). https://doi.org/10.1039/c3ee42106f

43. L.G. Cançado, A. Jorio, E.M. Ferreira, F. Stavale, C.A. Achete et al., Quantifying defects in graphene via Raman spectroscopy at different excitation energies. Nano Lett. 11, 3190–3196 (2011). https://doi.org/10.1021/nl104132g

44. A. Eckmann, A. Felten, I. Verzbistkly, R. Davey, C. Casiragli, Raman study on defective graphene: Effect of the excitation energy, type, and amount of defects. Phys. Rev. B 88, 035426 (2013). https://doi.org/10.1103/PhysRevB.88.035426

45. Y. Zhang, Z. Li, P. Kim, L. Zhang, C. Zhou, Anisotropic hydrogen etching of chemical vapor deposited graphene. ACS Nano 6, 126–132 (2012). https://doi.org/10.1021/nn202996r

46. V.K. Singh, S. Kumar, S.K. Pandey, S. Srivastava, M. Mishra et al., Fabrication of sensitive bioelectrode based on atomically thin CVD grown graphene for cancer biomarker detection. Biosens. Bioelectron. 105, 173–181 (2018). https://doi.org/10.1016/j.bios.2018.01.014

47. S. Bai, C. Chen, R. Luo, A. Chen, D. Li, Synthesis of MoO3/reduced graphene oxide hybrids and mechanism of enhancing H2S sensing performances. Sensors Actuat. B Chem. 216, 113–120 (2015). https://doi.org/10.1016/j.snb.2015.04.036

48. A. Chithambararaj, N. Sanjini, A.C. Bose, S. Velmathi, Flower-like hierarchical h-MoO3: new findings of efficient visible light driven nano photocathode for molybene. Catal. Sci. Technol. 3, 1405–1414 (2013). https://doi.org/10.1039/C3CY20764A

49. D. He, Z. Peng, W. Gong, Y. Luo, P. Zhao et al., Mechanism of a green graphene oxide reduction with reusable potassium carbonate. RSC Adv. 5, 11966–11972 (2015). https://doi.org/10.1039/C4RA14511A

50. S. Verma, R.K. Dutta, A facile method of synthesizing ammonia modified graphene oxide for efficient removal of...
uranyl ions from aqueous medium. RSC Adv. 5, 77192–77203 (2015). https://doi.org/10.1039/C5RA10555B

51. H. Zhang, D. Hines, D.L. Akins, Synthesis of a nanocomposite composed of reduced graphene oxide and gold nanoparticles. Dalton Trans. 43, 2670–2675 (2014). https://doi.org/10.1039/C3DT52573B

52. M. Dhanasankar, K. Purushothaman, G. Muralidharan, Effect of temperature of annealing on optical, structural and electrochromic properties of sol-gel dip coated molybdenum oxide films. Appl. Surf. Sci. 257, 2074–2079 (2011). https://doi.org/10.1016/j.apsusc.2010.09.052

53. A. Klinbumrung, T. Thongtem, S. Thongtem, Characterization of orthorhombic α-MoO3 microplates produced by a microwave plasma process. J. Nanomater. 2012, 5 (2012). https://doi.org/10.1155/2012/930763

54. J.E. Johns, M.C. Hersam, Atomic covalent functionalization of graphene. Acc. Chem. Res. 46, 77–86 (2013). https://doi.org/10.1021/ar300143e

55. C.M. Proctor, T.-Q. Nguyen, Effect of leakage current and shunt resistance on the light intensity dependence of organic solar cells. Appl. Phys. Lett. 106, 083301 (2015). https://doi.org/10.1063/1.4913589

56. Q. Ding, F. Meng, C.R. English, M. Cabán-Acevedo, M.J. Shearer et al., Efficient photoelectrochemical hydrogen generation using heterostructures of Si and chemically exfoliated metallic MoS2. J. Am. Chem. Soc. 136, 8504–8507 (2014). https://doi.org/10.1021/ja5025673

57. S. Haussener, S. Hu, C. Xiang, A.Z. Weber, N.S. Lewis, Simulations of the irradiation and temperature dependence of the efficiency of tandem photoelectrochemical water-splitting systems. Energy Environ. Sci. 6, 3605–3618 (2013). https://doi.org/10.1039/C3EE41302K