A sustainable molybdenum oxysulphide-cobalt phosphate photocatalyst for effectual solar-driven water splitting

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HIGHLIGHTS
- Facile synthesis of Molybdenum oxysulphide-cobalt phosphate (MoOxSy-CoPi) novel material for solar water splitting.
- The MoOxSy-CoPi photocatalyst corresponded to the most stable oxidation states of Mo, Co, S, P, and O in the composite.
- Nanoflowers of MoOxSy-CoPi showed controlled thickness up to ~10 nm, and variable interspaces of 200-400 nm were produced.
- Band-gap energy of 2.44 eV and significantly higher (7-8 folds) photocurrent density of MoOxSy-CoPi is recorded compared to MoOxSy and CoPi.

GRAPHICAL ABSTRACT
A facile fabrication of MoOxSy-CoPi photocatalyst revealed a conformal growth of nanoflowers. The photoelectrochemical studies exhibited significantly enhanced photocurrents with a stable and repeatable photocurrent response under visible light (1SUN) simulated solar irradiation.

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ABSTRACT
Introduction: Hydrogen is considered as a clean alternative green energy future fuel. Since the Honda-Fujishima effect for photoelectrochemical water splitting is known, there has been a substantial boost in this field. Numerous photocatalysts based on metals, semiconductors, and organic-inorganic hybrid-systems have been proposed. Several factors limit their efficiency, e.g., a stable PEC-WS setup, absorbing visible light, well-aligned band energy for charge transfer, electrons and holes, and their separation to avoid recombination and limited water redox reactions. Metallic doping and impregnation of stable and efficient co-catalysts such as Pt, Ag, and Au showed enhanced PEC-WS. We used Cobalt-based co-catalyst with molybdenum oxysulfide photocatalyst for effectual solar-driven water splitting.

Objectives: To develop photocatalysts for efficient PEC processes capable of absorbing sufficient visible light, good band energy for effective charge transfer, inexpensive, significant solar-to-chemical energy

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Methods: We prepared Molybdenum oxide-sulphide-cobalt phosphate photocatalyst for PEC-WS through a facile hydrothermal route using ammonium heptamolybdate, thiourea, and metallic Cobalt precursors.

Results: An effectual photocatalyst is produced for solar-driven water splitting. The conformal morphophotocatalysts of MoO_{3x}CoPi nanoflowers is a significant feature, as observed under FE-SEM and HR-TEM. XRD confirmed the degree of purity and orthorhombic crystal structure of MoO_{3x}CoPi. EDX and XPS identify the elemental compositions and corresponding oxidation states of each atom. A 2.44 eV band-gap energy is calculated for MoO_{3x}CoPi from the diffused reflectance spectrum. Photo- Electrochemical Studies (PEC) under 1-SUN solar irradiation revealed 7-8 folds enhanced photocurrent (~3.5 mA/cm²) generated from MoO_{3x}CoPi/FTO in comparison to Co-Pi/FTO (~0.5 mA/cm²) and MoO_{3x}CoPi respectively, within 0.5 M Na_{2}SO_{4} electrolyte (@pH=7) and standard three electrodes electrochemical cell.

Conclusion: Our results showed MoO_{3x}CoPi as promising photocatalyst material for improved solar-driven photoelectrochemical water splitting system.

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# Introduction

Developing semiconducting nanomaterials for photoelectrochemical water splitting (PEC-WS) has gained substantial consideration [1–3] since the invention of the Honda-Fujishima effect [4,5]. Sunlight is a source of alternative energy, and its venture with photocatalytic nanomaterials can split water into molecular hydrogen and oxygen [6,7]. Lee et al. [8] and Wu et al. [9] illustrated photoelectrochemical solar water-splitting reaction consisting of an oxidation half-reaction, i.e., a multistep four-proton, four-electron process, which needs large overpotential to overawe the competing electron-hole recombination processes at the expense of solar–energy–conversion efficiency [8,10]. Significant research is carried out to investigate varieties of active photocatalytic materials [10–12]. S Kment et al. reported Fe-O based semiconductors for PEC-WS applications [13], and L Zhu et al. [14] prepared textured TiO_{2}/CuO composite nanofibers for enhanced solar H_{2} generation. It is quite evident that such photocatalysts are aiding to achieve the escalating global energy demand and to discover sustainable energy resources [13–16]. However, further investigation is vital for an economical photocatalyst for PEC-WS. Developing a prolific photocatalysts needs, a stable PEC-WS setup, an efficient PEC processes that must be capable of absorbing a large fraction of the visible light, retain well-aligned band energy for thermodynamically promising charge transfer, and last but not least should be inexpensive [10,17]. Consequently, all characteristics mentioned above are requisite for effective photocatalyst for PEC-WS with significant solar-to-chemical energy conversion efficiencies to make these systems commercially viable.

Transition metal dichalcogenide, i.e., Molybdenum disulphide (MoS_{2}) and oxy-sulphides (MoOxSy) semiconducting nanomaterials [18,19] have been recently explored for PEC-WS. Owe to their entralling electrical, optical, and catalytic properties, these nanomaterials are used to produce hydrogen and oxygen as alternative and sustainable green energy resources [20]. Different strategies, for instance, mechanical exfoliation [21], solution-based chemical exfoliation [22], chemical vapor deposition (CVD) [23], hydro and solvothermal synthesis [24], etc have been adopted to fabricate Molybdenum-oxide-sulphides (MoOxSy) and disulphide (MoS_{2}) photocatalysts. MoS_{2} based nanomaterials are well-investigated for PEC-WS in comparison to MoOxSy photocatalysts. MoS_{2} nanomaterials showed good electrical conductivity, i.e., charge carrier mobility ≥ 200 cm²/V·s in monolayer and up to 517 cm²/V·s in multilayers. Besides, their optical studies showed that MoS_{2} exhibits a band-gap of ~ 1.2 eV- 2.2 eV in its monolayer and multilayers/bulk forms [25]. On the contrary, limited studies on Mo oxides or oxy sulphides are contained in the literature, likely because of solubility issues, mostly in acidic electrolytes and inadequate PEC characteristics. Nevertheless, Mayhall et al. [26], Sinaim et al. [27], and Xie et al. [28] reported that incorporation of oxygen into Mo/MoS_{2} greatly enhance the stability of the photocatalyst, limits its solubility, and improves the PEC properties of Mo based oxides and oxy sulphides. Therefore, our focus is to enhance those properties mentioned earlier for Mo, MoS_{2}, and MoOxSy and their hybrid nanocomposites to be used as potential photocatalytic nanomaterials in solar cells and solar-driven PEC-WS applications [29].

In PEC-WS, hydrogen is generated following a hydrogen evolution reaction (HER). In such electrochemical reactions, the catalyst plays a decisive role in reducing the electrodes’ overpotential that subsequently promote reaction efficiency. Likewise, the edges in 2-D Mo photocatalysts, i.e., MoS_{2} and MoOxSy, the layered nanostructures and active sites located along with them, are coordinately unsaturated and thermodynamically unfavorable in addition to their basal surfaces that are chemically inert. These features correspond to their significant electrochemical properties [30]. Conversely, the utility of Mo based materials becomes limited in energy storage and conversion processes due to its high surface energy and strong interlayer π- π interactions that result in the restacking of layered nanostructures, hence showed poor stability, loss of active sites, and infrequent properties [31]. However, due to inadequate HER catalysts, there is an ongoing demand for substituting such nanomaterials and strategies. Recent research efforts have identified several attractive photocatalytic nanomaterials and molecular catalysts in photoelectrochemical (PEC) water splitting with numerous advantages. Despite its success, some factors in PEC hydrogen production need more improvements such as energy conversion efficiencies, limited charge recombination, and inefficient catalytic reduction and oxidation reactions. The incorporation of oxygen and a co-catalyst with Mo based compounds for both oxidation and reduction reactions have been used as an alternative approach to improve the photocatalytic activity of the nanomaterials in PEC-WS processes [32]. Among the co-catalysts, noble metal co-catalysts are well recognized, but the use of precious metals (Au, Ag, Pt, Pd) is not cost-effective. Hence this prompts the quest for alternative co-catalysts such as cobalt phosphate (CoPi). The literature revealed that cobalt phosphate (CoPi) significantly enhanced the PEC-WS properties in 1D-WO_{3}/BiVO_{4}. That is attributed to a reduction in charge recombination, facilitating electron-hole transfer, and reducing the overpotential for OER in the water-splitting process [33].

Herein, based on the characteristics above for Mo, the significant role of oxygen incorporation into MoS_{2} and support of Co-Pi as co-catalyst, we designed a novel Molybdenum oxide-sulphide-cobalt phosphate (MoO_{3x}CoPi) photocatalyst for PEC-WS. Our research strategy has some intriguing features. Firstly, the novelty of the MoO_{3x}CoPi nanocomposite material in terms of its elemental composition and conformal growth of MoO_{3x}CoPi...
nanoflowers. Secondly, the ease and straightforward synthetic protocol, i.e., the in-situ growth of MoOxSy nanoflowers and cobalt phosphate (CoPi) co-catalysts in between MoOxSy layers. Last but not least, the cost-effectiveness in the fabrication of MoOxSy-CoPi, consolidating simple materials, and experimentation. In addition, our experiments showed that MoOxSy-CoPi/FTO enhanced the PEC signal by a factor of seven and eight in comparison to MoOxSy/FTO and Co-Pi/FTO photocatalysts, respectively. Therefore, considering the above traits, it is foreseen that this novel photocatalyst material will open new research directions to augment the challenges in solar-driven PEC-WS.

Experimental strategies

Materials and chemicals

The chemicals, reagents, and materials used in this study are of analytical grade and used as received unless otherwise stated. Ammonium heptamolybdate tetrahydrate, Thio-urea, cobalt powder, phosphoric acid (85%), ammonia solution, sodium sulfate (Na2SO4), ethanol and acetone are purchased from Sigma Aldrich. We used deionized water in all experiments.

Synthesis of nanomaterials

Synthesis of cobalt phosphate (CoPi)

CoPi synthesis is carried out from metallic cobalt powder and phosphoric acid (85%). The equimolar reaction mixture is sonicated at 50 °C continuously with the dropwise addition of 10 ml deionized water at the rate of 2 ml/10 min. After the complete addition of deionized water, the reaction mixture is sonicated for a further half an hour at 60 °C and then cooled to room temperature. In the next step, 25% ammonia solution (5 ml or as required) is added to initiate precipitation, while maintaining pH at 13. Later on, the reaction mixture is transferred into a steeliness steel autoclave containing Teflon vessel. The solvothermal reaction has proceeded at 60 °C for 2 h at 200 °C. Afterward, the reaction mixture is centrifuged at 4000 RPM for 5 min. The violet color product collected and washed with deionized water several times to remove undesired and unreacted moieties before drying in a vacuum oven for 2 h at 150 °C.

Synthesis of molybdenum oxysulfide-Cobalt phosphate nanocomposite (MoOxSy-CoPi)

The equimolar amounts of ammonium heptamolybdate and thiourea mixed in deionized water are stirred to obtain a clear mixture. In this mixture, an equimolar amount of as prepared CoPi is added slowly with continuous stirring. The reaction mixture is placed in an ultrasonic bath for one hour at 60 °C to acquire homogeneous mixing. The reaction mixture is transferred into a steeliness steel autoclave containing a Teflon vessel to proceed with a hydrothermal reaction for 24 h at 180 °C. The white precipitate of MoOxSy-CoPi is centrifuged at 4000 RPM for 5 min, washed with ethanol and acetone, respectively, before drying in a vacuum oven for 200 °C for two hours. Finally, the MoOxSy-CoPi photocatalyst nanomaterial is annealed at 400 °C for 4 h in the air.

Synthesis of molybdenum oxysulfide (MoOxSy)

MoOxSy is prepared by the same strategy as mentioned above in section-b but without Co-Pi. The purpose of MoOxSy synthesis is only to study and compare its morphology and PEC properties with MoOxSy-CoPi and CoPi.

Characterization of nanomaterials

The purity and crystalline phases of MoOxSy-CoPi nanoflowers and CoPi are characterized by X-ray diffraction (XRD) technique using a Benchtop MiniFlex -X-ray Diffraction instrument (Mini-XRD) from Rigaku (40 kV and 15 mA) in the range of 10–70° (2θ) at a scanning rate of 3° min⁻¹ with CuKα radiation. The structural composition and crystalline phases prevailing in MoOxSy-CoPi nanoflowers and CoPi are interpreted with library data. Also, the elemental composition and oxidation states of metals involved during MoOxSy-CoPi formation are verified by X-ray photoelectron spectroscopy (XPS) using PHI 5000 Versa Probe II spectrometer (UV/Vac-PhII) employing Al Kα as the incident radiation source. The C1s (E = 284.5 eV) level served as the internal standard. The morphology of the MoOxSy-CoPi nanoflowers, MoOxSy, and CoPi is observed under TESCAN Lyra 3 Field Emission Dual Beam (Electron / Focused Ion Beam) system combined with high-end high-resolution electron microscope (FE-SEM). This machine is also facilitated with EDX for elemental determination/mapping. HR-TEM study is carried out via JEM2100F, JEOL at 200 kV imaging to comprehend the MoOxSy and CoPi interfaces formed in the composite nanoflowers. Optical properties, i.e., diffused reflectance spectroscopy of MoOxSy-CoPi nanoflowers and CoPi, are carried out using Agilent Cary 5000 high-performance UV-Vis-NIR spectrophotometer containing praying mantis accessory with alignment tools and powder cell sample cups. In situ Raman spectroscopy of MoOxSy-CoPi nanoflowers and CoPi was performed with a laser (300 mW Green 532 nm) by iHR-320 Horiba imaging spectrometer. The vibrational modes in the MoOxSy-CoPi nanoflowers and CoPi are recorded by FT-IR 6700 Nicolet™ FT-IR spectrometer. A fluorolog-3 imaging spectrophotometer tests the photoluminescence activity of the nanomaterials at an excitation wavelength of 350 nm and a slit width of 2 nm.

Fabrication of MoOxSy-CoPi/FTO, MoOxSy /FTO, and CoPi/FTO photocatalysts for PEC

Studies

Fluorinated Tin Oxide (FTO) conducting glass substrates are used for photoelectrochemical studies of MoOxSy-CoPi and CoPi. Initially, FTO glass substrates are washed by acetone (10 min) and water (10 min), respectively, with continuous ultrasonication. In the next step, slurries of MoOxSy-CoPi nanoflowers, MoOxSy, and CoPi from 0.25 g of the materials are prepared in ethanol/water (70%/30%) mixture and coated over pre-treated FTO glass substrates by drop-casting, respectively. The optimized film is highly uniform and completely covered the desired area of the FTO substrate. The MoOxSy-CoPi/FTO, MoOxSy/FTO, and CoPi/FTO electrode were heated at 110 °C for 2 h to evaporate the solvents and harden the MoOxSy-CoPi nanoflowers and CoPi layers so that they could withstand during the PEC measurements. The PEC measurements are carried out by a three-electrode system in 0.5 M Na2SO4 (pH = 7) solution, where a Pt wire served as a counter electrode. The FTO coated MoOxSy-CoPi nanoflowers, and CoPi substrates served as photocatalysts or working electrodes, and the standard calomel (SCE) as a reference electrode. The PEC measurements are performed by Metrohm Autolab Potentiostat (PGSTAT302N). Moreover, for simulated solar light, we utilized Oriel sol 3A class AAA Solar Simulator-Newport (100 mW/cm²) IEC/IEC/ASTM certified containing 450Watt Xenon lamp, Air Mass 1.5G Filter, and 2x2 inch aperture for output beam.

Results and discussions

FE-SEM & HR-TEM studies of MoOxSy-CoPi and CoPi nanomaterials

The surface studies of MoOxSy, CoPi, and MoOxSy-CoPi nanocomposite are carried out via the FE-SEM technique, as shown in Figs. S1-A, S1-B (In supplementary information), and Fig. 1 respectively. The solid MoOxSy shows distorted nanorods or flakes like growth, supported in literature through a different synthesis
The nanomaterial (MoOxSy) is synthesized by the same procedure as adopted for the MoOxSy-CoPi, i.e., the hydrothermal synthesis. However, the morphology of MoOxSy differs from what we observed for MoOxSy-CoPi at the same reaction conditions. Similarly, the FE-SEM of bare CoPi revealed particulate morphology, with EDS, indicated the constituent elemental ratios (Figs. S2 & S3) in a good agreement. Fig. 1 (a,b) and (c,d) show low and high-resolution micrographs of MoOxSy-CoPi nanoflowers. The MoOxSy-CoPi exhibit a uniform nanoflowers like morphology, with controlled nanosheets thickness, variable internal spaces, and good crystallinity (as supported by XRD studies). Under high magnification FE-SEM, the nanoflower growth of MoOxSy is observed with CoPi incorporation within the flakes of MoOxSy as a result of the hydrothermal reaction by taking equimolar amounts of both precursors. While looking at Fig. 1d, the thickness of the curved nanopetals in MoOxSy-CoPi nanoflowers can be estimated, which ranges beyond 10 nm. However, the internal distances between curved channels show variation between 200 and 400 nm. To further confirm, we performed energy dispersive spectroscopy (EDS) and transmission electron microscopy (TEM) analysis. Fig. 2 shows the EDS spectrum performed at different sites of the MoOxSy-CoPi composite nanoflowers with their corresponding atomic ratios that substantiated significant atomic ratios of Mo, Co, S, O, and P, respectively. Thus it is confirmed the equimolar amounts of both the precursors resulted not only in optimal material growth but also showed enhanced PEC-WS properties.

The TEM was performed to confirm the dispersion of CoPi within the channels of MoOxSy nanoflowers supported by high angle annular dark-field (HAADF) scanning transmission electron microscopy (STEM). Two different morphologies corresponded to MoOxSy and CoPi can be seen in the higher magnification TEM image in Fig. 3a. Likewise, Fig. 3b presents an evaluation of the blue cross-section via HRTEM elucidating two kinds of d-spacing patterns that are usually characteristics of nanocomposite materials.
with good crystallinity. **Fig. 3 (c & e)** shows high resolution (HR)-
TEM micrograph, the Fast-Fourier transform (FFT) micrographs
further demonstrate the MoO$_x$S$_y$-CoPi nanoflower composite for-
mation. It is evident from these micrographs that the MoO$_x$S$_y$ exhi-
bbit larger d-spacing (0.56 nm) than CoPi (0.28 nm). The linear EDX
spectra agree with EDX obtained from FESEM (**Fig. S4 in
supplementary information**). Thus, the HR-TEM studies provided
ample evidence for the successful formation of MoO$_x$S$_y$-CoPi
nanocomposite.

**X-Ray diffraction (XRD) analysis**

X-ray diffraction analysis is performed to explore the crystal
structure and the degree of purity of CoPi and MoO$_x$S$_y$-CoPi nano-
structures. It revealed that MoO$_x$S$_y$-CoPi materials have a polycrys-
talline structure with orthorhombic crystalline phases. XRD
patterns of MoO$_x$S$_y$-CoPi and CoPi nanostructures are shown in

**Fig. 2.** EDS studies of MoO$_x$S$_y$-CoPi nanoflowers a single-phase composite material fabricated $\Theta$ 2 h at 180°C via a hydrothermal route (a), (b) & (c) present EDS spectrums with corresponding elemental peaks for Mo, Co, S, P, and O respectively. The onset SEM micrographs present element distribution images of MoO$_x$S$_y$-CoPi as well as atomic ratios of Mo, Co, S, P, and O.

**Fig. 4.** Both MoO$_x$S$_y$-CoPi and CoPi display diffraction peaks of MoO$_x$S$_y$ and CoPi at their corresponding indexes through matching with the library data under JCPDS card No. 65-2421 and 01-078-6161. The XRD pattern of MoO$_x$S$_y$-CoPi matches with MoO$_3$ pattern, suggesting that most of the sulfur is replaced by oxygen during annealing. The CoPi showed peak intensities at 2$\theta$ angle are $\approx$18, 21, 26, 31, 36, 39, 42, 45, 52, 55 and 57 with corresponding indexes observed at (200), (101), (211/020), (200), (220), (102), (202), (212), (131), (231) and (412), respectively. The sharp and high intensities diffraction peaks exposed the formation of high crystalline orthorhombic CoPi that is also supported by various literature studies [36,37]. The XRD spectra of MoO$_x$S$_y$-CoPi composite presented in **Fig. 4** have main diffraction peaks at $\approx$13, 24, 26, 28, 39, 46, 49 with corresponding indexes at (002/200), (110/101), (040), (021/201), (060/600), (002) respectively, which are in correspondence to MoO$_3$. Some CoPi diffraction peaks at $\approx$18, 21, 31, 36, 42 with corresponding indexes at (200), (101),
(211/020), (220), (202) either completely disappeared or shifted into broad peaks in the MoO$_x$Sy-CoPi composite. Most likely, it is due to new chemical interactions established between MoO$_x$Sy and CoPi. The role of effective ionic radii [38] cannot be ignored: elements of nearly close ionic radii in different oxidation states like Mo & Co and O & S can develop new chemical interactions and phases during the composite formation. Considering this, the appearance of some new crystal phases at peak intensities $2\theta = 28$ (021/201) and broad peaks at $2\theta = 13$ (020/200), 24 (110/101) & 46 (060/600) endorses the formation of MoO$_x$Sy-CoPi nanocomposite material.

The XRD patterns of MoO$_x$Sy-CoPi also showed three characteristic MoO$_x$ diffraction peaks at $2\theta$ angles $\sim 13$, 26, and 39 corresponding orthorhombic crystalline phase indexes (020), (040), and (060/600), respectively [18,39]. The slight shifting in these peaks is attributed to the formation of new composite material and the synthetic strategy adopted. In summary, the XRD patterns of MoO$_x$Sy-CoPi showed some new and combined peaks that arose due to the possible existence of sulfur contents and also the composite formation of MoO$_x$Sy with CoPi via a hydrothermal approach.

X-Ray photoelectron spectroscopy

The elemental composition of MoO$_x$Sy-CoPi composite and their corresponding oxidation states are investigated by X-ray photoelectron spectroscopy (XPS). The spectrums are presented in Fig. 5. A complete elemental XPS survey indicated the presence of Mo, S, Co, P, and O elements (Fig. 5a), which are present in the MoO$_x$Sy-CoPi and found in good agreement with the EDX and TEM findings. This not only comprehended the formation of MoO$_x$Sy-CoPi nanoflower composite but also supported the in-situ impregnation of CoPi in MoO$_x$Sy layers during a hydrothermal reaction. All the peaks are adjusted with the reference C 1s peak (284.6 eV) before further description. Fig. 5b shows the Mo (3d) spectra consisting of two spin–orbit triplet peaks characteristics of Mo-S (3d$_{5/2}$), Mo-S (3d$_{3/2}$), and a residual Mo-O (3d), respectively [18]. These Mo (3d) components at $\sim 229.7$ eV, 232.5 eV, and 235.5 eV can be assigned as Mo$^{4+}$, Mo$^{6+}$ and Mo$^{2+}$ oxidation states, respectively. The occurrence of reduced Mo oxidation states indicate that the d-orbital is partially filled; therefore, it showed the enhanced electrical conductivity[24], as can be seen in PEC-WS. Likewise, the S (2p) spectra, as presented in Fig. 5c, contains residual peaks with two dominant spin–orbit doublets, i.e., characteristics of S$^2$ 2p$_{3/2}$ at 161.8 eV and S$^2$ 2p$_{1/2}$ at 164.2 eV, respectively.
which specifies that Sulphur exists as a very stable oxidation state, i.e., (-2) in the MoO$_x$S$_y$-CoPi composite material [24]. However, some lower intensity doublet peaks characteristic of S$_2^-$ 2p$_{1/2}$ at 160.7 eV and 163.2 eV correspond to further oxidation of Sulphur in the MoO$_x$S$_y$ composite [40]. Benoist et al [40] reported a similar environment of Sulphur in MoO$_x$S$_y$ films. In light of their study, the S 2p peaks are split up into four components (160.7 eV, 161.8 eV, 163.2 eV, and 164.2 eV), as shown in Fig. 5c. The binding energies of these doublets are close to those determined for the S 2p peak of the sulfur-rich thin films having the composition and environment similar to that of MoS$_2$, this sulfur is obviously S$_2^-$, the real effective charge of which is more negative than the actual charge of S$_2^-$ in MoS$_2$. It is also observed that when the amount of oxygen increases in MoO$_x$S$_y$, the central doublet peak (163.2 eV) decreases. However, the doublets at higher and lower binding energies (peaks at 164.2 eV and 161.9 eV) equally increase (Fig. 5c). Thus, both sulfur and oxygen ions are associated with the new species of molybdenum oxy sulphide. Benoist et al [40] confirmed that their XPS data showed new “oxysulfide” in which molybdenum environment is composed of oxygen (O$^-$) and sulfur (S$^2$) in equal proportions. Their findings were also well aligned with electron spin resonance (ESR) studies conducted by Deroide et al. [41,42] for molybdenum sulphide compounds.

The XPS data presented in Fig. 5 (c & d) disclosed an interesting observation; the spectra concluded that MoO$_x$S$_y$ and CoPi remain intact in the MoO$_x$S$_y$-CoPi nanoflower composite material [18]. Fig. 5 (d) further elaborates on the XPS insights of CoPi in MoO$_x$S$_y$-CoPi nanocomposite material. The XPS spectra of Co demonstrate a doublet of 2P splitting, i.e., 2P$_{3/2}$ and 2P$_{1/2}$ peaks with analogous shakeup satellites peaks, i.e., each splitting correspond to three spectral lines. Thus the Co (2p$_{3/2}$) spectral lines observed at 777.9 eV, 781 eV, and 784 eV can be assigned to Co, Co-O, and Co-P [43–45], respectively. However, Co (2p$_{1/2}$) spectral lines observed at 793.0 eV, 796 eV, and 799 eV can be assigned to Co, Co-O, and Co-O(sat) [44,46], respectively. Fig. 5 (e) displays the P (2p) spectra that exhibit characteristic 2p$_{3/2}$ doublets of Phosphorous and (f) demonstrates the spectra of O (1 s), i.e., low energy metal-oxygen (M–O) and higher energy nonmetal-oxygen (NMO) photoelectron peaks.

Fig. 5. XPS spectra of the constituent elements of MoO$_x$S$_y$-CoPi nanoflowers, where (a) shows complete elemental XPS survey indicated the presence of Mo, S, Co, P, and O elements, (b) shows the Mo (3d) spectra consisting of two spin–orbit triplet peaks characteristics of Mo-S (3d$_{5/2}$), Mo-S (3d$_{3/2}$), and a residual Mo-O (3d), respectively, (c) contains residual peaks with two dominant spin–orbit doublets S$^2^-$ 2p$_{3/2}$ & S$^2^-$ 2p$_{1/2}$, (d) elaborates XPS spectra of Co doublet 2P splitting, i.e., 2P$_{3/2}$ and 2P$_{1/2}$, (e) displays the P (2p) spectra that exhibit characteristic 2p$_{3/2}$ doublets of Phosphorous and (f) demonstrates the spectra of O (1 s), i.e., low energy metal-oxygen (M–O) and high energy nonmetal-oxygen (NMO) photoelectron peaks.

Fig. 6. Band energy estimation (a) diffused reflectance spectrum for band-gap estimation of MoO$_x$S$_y$-CoPi nanoflowers (d) diffused spectrum for band-gap calculation of CoPi.
529.5 eV, and 531 eV, i.e., for Mo-O and Co-O, respectively. Furthermore, a higher energy nonmetal-oxygen (NMO) photoelectron peak corresponding at > 531 eV, i.e., for O-S and O-P, which can be attributed to the phosphate-oxygen and adsorbed moisture, i.e., O-H bond [49,50]. These XPS findings confirm that MoO$_3$S$_n$ and CoPi retained their chemical identity in the MoO$_x$S$_y$-CoPi nanocomposite, as the binding energy of the constituent elements agrees with reported values for Mo$_2$ and CoPi, markedly [18].

**Optical properties of CoPi and MoOxSy-CoPi nanomaterials**

**UV–VIS spectroscopy and bandgap**

Diffused reflectance spectra of pure MoO$_x$S$_y$-CoPi and CoPi are presented in Fig. 6. Their optical properties are investigated using Cary 4000 series UV–VIS-NIR spectrophotometer. The blend of CoPi within MoO$_x$S$_y$ nanostructure changes the optical characterization of the nanocomposite material. The band-gap energy of CoPi (inset plot in Fig. 6) is estimated from the diffused reflectance spectrum, and it is found to be ~2.6 eV. The literature revealed MoS$_2$ and MoO$_x$ exhibit band-gap energies ~1.2–2.2 eV in its monolayer and multilayer/bulk form, respectively [25]. In our studies, the band-gap energy observed for MoO$_x$S$_y$-CoPi composite from Tauc’s Plot method [51,52] is 2.44 eV, as can be perceived in Fig. 6 (a). A noticeable difference in the band-gap energies from CoPi and MoO$_x$S$_y$ is observed. This shift in band gap energies could be attributed to changing phase and crystal structure upon the incorporation of CoPi to MoO$_x$S$_y$. Besides several factors such as morphology, variation in crystallographic phases and structure, particle size, annealing temperature, etc. can influence the optical properties of semiconducting materials [53].

**Vibrational spectroscopy**

**FTIR-Spectroscopy**

The structural properties of MoO$_x$S$_y$-CoPi and CoPi are further expounded via Raman and FTIR spectroscopy. The FT-IR analysis of both CoPi and MoO$_x$S$_y$-CoPi nanoflowers composite material are presented in Fig. S5-A and S5-B respectively of supplemental information. The FT-IR for CoPi showed peaks at vibrational frequencies 620 cm$^{-1}$, 755 cm$^{-1}$, 860 cm$^{-1}$, 1050 cm$^{-1}$, 1440 cm$^{-1}$, 1650 cm$^{-1}$, 2750 cm$^{-1}$, 3240 cm$^{-1}$ and 3380 cm$^{-1}$. These are peak positions that are in good agreement with the reported FT-IR for CoPi in literature [36,54,55] i.e., 605 cm$^{-1}$, 956 cm$^{-1}$, 1008 cm$^{-1}$, 1045 cm$^{-1}$, 1632 cm$^{-1}$, 2930 cm$^{-1}$ and 3450 cm$^{-1}$. The most plausible reason of this shifting is the synthetic strategy and morphology of the material synthesized. On the other hand MoO$_x$S$_y$-CoPi shows IR peaks at 510 cm$^{-1}$, 780 cm$^{-1}$, 860 cm$^{-1}$, 960 cm$^{-1}$, 1080 cm$^{-1}$, 1140 cm$^{-1}$, 2015 cm$^{-1}$, 2255 cm$^{-1}$, 3120 cm$^{-1}$ and 3240 cm$^{-1}$ respectively. These bands corroborate with the characteristics peaks of both MoO$_2$ [56–59] and CoPi at their respective vibrational modes as reported in the literature [36,54,55]. The IR band at 510 cm$^{-1}$ corresponds to $\nu_{as}$ (Mo–S) vibrations. However, the bands above 600 cm$^{-1}$ correspond to Mo–O type of vibrations due to reaction presence of oxygen in the Mo–O bond, thus different peaks as labeled in Fig. S5-B indicate that oxygen is present in similar types of coordination environment as observed for MoO$_3$ [56]. We allocated the bands at 860 cm$^{-1}$ and 960 cm$^{-1}$ to the $\nu_{as}$(Mo–O) vibrations and 780 cm$^{-1}$ to (Mo–O) vibrations. Some IR peaks observed in CoPi or MoO$_x$S$_y$, respectively either disappeared or shifted to lower intensities and some become intense and sharp in MoO$_x$S$_y$-CoPi. This shifting of IR bands is attributed to formation of new composite that is mainly depending upon the synthetic methodology adopted, crystalline phases and morphology.

**Raman spectroscopy**

The Raman spectrum indicates characteristic phononic peaks in the fingerprint regions from 0 cm$^{-1}$ to 1200 cm$^{-1}$. Fig. S5-C in supplementary information, presents the Raman scattering of both MoO$_x$S$_y$-CoPi (red color line) and CoPi (blue color line), respectively. All the peaks are assigned according to their modes of vibrations comparable with the literature [60,61]. The vibrational peaks for CoPi are observed in region 50–200 cm$^{-1}$ and 940 cm$^{-1}$ respectively. However, these peaks appeared sharper and intense after hydrothermal reaction for MoO$_x$S$_y$-CoPi formation, i.e., peaks recorded at 50 cm$^{-1}$, 100 cm$^{-1}$, 140 cm$^{-1}$ and 980 cm$^{-1}$ respectively. Furthermore, the characteristics Raman scattering of MoO$_x$S$_y$ in MoO$_x$S$_y$-CoPi nanoflowers composite are shifted to lower values, i.e., the phononic peaks observed at 310 cm$^{-1}$ ($E_{2g}$) and 355 cm$^{-1}$ ($A_{1g}$), respectively. These peaks are assumed to be first-order Raman vibration modes within S–Mo–S layers [61]. Also, the appearance of new sharp and broad phononic peaks at 225 cm$^{-1}$, 260 cm$^{-1}$, 650 cm$^{-1}$, and 800 cm$^{-1}$ are considered due to interactions between MoO$_x$S$_y$ and CoPi atoms, e.g., P–Co–S or P–Mo–S or O atom to P, S, and Mo or vice versa. Nonetheless, the vibrational studies authenticate the formation of MoO$_x$S$_y$-CoPi that have potential photoelectrochemical applications.

**Photoluminescence’s spectroscopy**

Photoluminescence (PL) spectroscopy of MoO$_x$S$_y$-CoPi and CoPi is carried out at room temperature with an excitation wavelength of 350 nm, as described in Figs. S6-A and S6-B in supplementary information, respectively. Fig. S6-B presents the first onset of the PL curve, which lies in a visible light region with PL peak maxima observed at ~425 nm. The onset initiates at 370 nm (~3.35 eV) and prolonged till 550 nm (~2.26 eV) as of Fig. S6-B (blue line baseline fitted PL spectrum). The maximum excitation is observed at ~425 nm (~2.92 eV) in the MoO$_x$S$_y$-CoPi. However, this peak for CoPi (Fig. S6-A in Supplementary information) is recorded between 410 and 420 nm (~3.03 eV-2.95 eV). This small shift of PL peak for CoPi in MoO$_x$S$_y$-CoPi is attributed to the appearance of some defects and crystalline phases in the growing MoO$_x$S$_y$-CoPi during the hydrothermal reaction. The PL spectra also displayed low-intensity peaks between 580 and 680 nm, as shown in Fig. S6-B (red PL spectra in onset). This broader peak corresponds to MoO$_x$S$_y$ as a result of direct excitonic transitions (A1 and B1) in the range of 620–650 nm (2.0–1.9 eV). These excitations are following the literature values [61,62], where A1 and B1 transitions are recorded at 630 nm and 670 nm (1.97 eV-1.85 eV), respectively. The shifting of PL emission peaks for CoPi and MoO$_x$S$_y$ in our synthesized MoO$_x$S$_y$-CoPi nanomaterials are attributed to slower electronic relaxation. Furthermore, this shifting could be due to crystalline structure and nanoflowers like morphology of MoO$_x$S$_y$-CoPi as compared to CoPi and MoO$_x$S$_y$ independently. The presence of structural defects on the surface of nanopetals of MoO$_x$S$_y$-CoPi nanoflowers composite materials is also responsible for different PL spectra. That ultimately enhanced the disordered arrangement among Mo, Co to Sulphur, oxygen, and phosphorus and shifted excitations little away from the band-gap energies [61,62].

**Photoelectrochemical performance of MoOxSy-CoPi, MoOxSy, and CoPi Photocatalysts**

**Chronoamperometry**

MoO$_x$S$_y$-CoPi/FTO, MoO$_x$S$_y$/FTO, and CoPi/FTO Photocatalysts are prepared and used for PEC measurements in a standard three-electrode electrochemical cell. The cell contains an FTO coated working electrode, a reference standard calomel electrode (SCE), and a platinum counter electrode in 0.5 M Na$_2$SO$_4$ electrolyte solution at neutral pH (7). The photoresponse of each photocatalyst...
exposed to 1 SUN light source is recorded after regular intervals of time. The chronoamperometric measurements are presented in Fig. 7(a-d), which shows photocurrent density vs. time (Jp vs. t) profiles. The photocurrent generated from MoOxSy-CoPi/FTO photocatalyst is in the range of nano amperes, i.e., ~150 nA at the at ~0.0 V. This is 4–5 times higher than photocurrent observed for CoPi at ~0.0 V (Fig. 7b), which is only ~30 nA. We further extended our chronoamperometry studies at higher applied potentials. Fig. 7 (c & d) shows chronoamperometry measurements of both CoPi and MoOxSy-CoPi at 0.2 V applied potential. Fig. 7c presents the highest photocurrent density, i.e., 3.5 mA.cm⁻² from MoOxSy-CoPi nanoflowers composite. A photoresponse further explicit significant stability of MoOxSy-CoPi coated on FTO over a longer period. A stability measurement for MoOxSy-CoPi/FTO photocatalyst further confirms this recital upon continuous exposure to 1 SUN solar simulated light without any significant degradation within a given time, as shown in Fig. S7 (supplementary information). The photocurrent density recorded from CoPi is in the range of 500 μA.cm⁻² as of Fig. 7d. While performing PEC measurements for both MoOxSy-CoPi nanoflowers composite material and CoPi, we exhibited reversible response profiles with adequate stability in ON/OFF light modes. That, in turn, reflects the successful performance of MoOxSy-CoPi nanoflowers composite material for water splitting processes. The linear sweep voltammetry experiments further complimented this efficient PEC performance, where the photocurrent density is enhanced to mA, i.e., ~3.5 mA at higher potential values recorded for MoOxSy-CoPi nanoflowers composite material. The morphology, reduced bandgap, and facile synthetic strategy are advantageous features of our approach that we have implemented in the fabrication of MoOxSy-CoPi nanoflowers composites to substantiate better PEC properties.

Fig. 7. Chronoamperometric measurements of MoOxSy-CoPi (a & c) and CoPi (b & d) at applied potentials of 0 V and 0.2 V respectively.

Fig. 8. Linear sweep voltamograms of blank FTO, MoOxSy/FTO, CoPi/FTO, and MoOxSy-CoPi/FTO photocatalysts at applied potential range of –1.20 V to 1.20 V under 1 SUN solar light in 0.5 M Na₂SO₄ electrolyte (pH 7).
**Linear sweep voltammetry (LSV)**

Fig. 8 shows a linear sweep voltammetry measurement of MoO$_x$S$_y$-CoPi, MoO$_x$S$_y$, CoPi, and blank FTO substrate, respectively. The measurement cell is developed by a standard three-electrode configuration using 0.5 M Na$_2$SO$_4$ electrolyte at pH 7. The potential range is selected between for −1.20 V to 1.20 V versus SCE under simulated solar light (1 SUN) at a scan rate of 100 mV/Sec. Fig. 8 shows that the pristine MoO$_x$S$_y$ and CoPi photoelectrodes bear very low current density. In contrast, the enhanced current density of ~0.6 mA/cm$^2$ is observed for MoO$_x$S$_y$-CoPi nanoflowers composite corresponding to a hydrogen evolution reaction (HER), which is 3–4 times more when compared with a current density of CoPi (~0.15 mA/cm$^2$) and MoO$_x$S$_y$ material. While looking at a positive potential, the oxygen evolution reaction (OER) is more preferred, and we observed a photocurrent density of ~ 3.5 mA/cm$^2$ from MoO$_x$S$_y$-CoPi. Whereas, the photocurrent density from CoPi and MoO$_x$S$_y$ is recorded around ~0.5 and 0.12 mA/cm$^2$, respectively. Besides this, we tested blank FTO under 1 SUN solar simulated light. The photoresponse of blank FTO is negligible during LSV measurements for the same selected potential range of ~1.20 V to 1.20 V. In overall LSV experiments, we observed dominance of OER by MoO$_x$S$_y$-CoPi. It endorses photochemical oxygen evolution preference of MoO$_x$S$_y$-CoPi nanocomposites material. Hence, these PEC characteristics are the most important achievements acknowledged. Which further supports our in-situ impregnation strategy of co-catalyst CoPi in growing MoO$_x$S$_y$ nanoflowers via facile hydrothermal approach. Consequently, we also observed better morphology, crystallinity, improved optical properties, and excellent photocatalyst current efficiencies. The curved morphology in nanopetals of MoO$_x$S$_y$-CoPi helped in better electron mobility and marginal electron-hole recombination[11,63].

**MoO$_x$S$_y$-CoPi charge transfer mechanism**

Fig. 9 explains the most plausible charge transfer mechanism that systematically based on improved interfacial hole transfer in-between FTO substrate and material. The integration of CoPi in MoOxSy resulted in suppressing the electron-hole recombination within MoOxSy-CoPi composite material [60]. As holes are serving oxygen evolution from water, therefore improved oxidation or OER is observed in comparison to reduction or HER processes from as engineered MoOxSy-CoPi photocatalyst in solar-driven water splitting. “Furthermore, significant photocurrent densities as a result of CoPi incorporation are observed. CoPi is a well-known co-catalyst used for charge separating networks at the catalyst’s interface. Literature revealed that an oxygen evolution catalyst like Co-Pi can catalyze the water electrolysis process to succeed at a low overpotential by changing the reaction pathway [64,65]. When it combines with the MoOxSy, the Co-Pi uses the valence band photo holes of the metal oxides/oxysulphide to initiate multiple catalytic reactions [66]. This is accompanied by fast output of photoholes from metal oxide/ oxysulphide to water molecule and undergoes water oxidation. Thus, the relieved charge accumulation at the electrode/electrolyte decreases the electron-hole recombination rate and leads to enhanced photo electrochemical performance. It is also known that additional band bending at the electrode/electrolyte interface may possibly be created by the Co-Pi catalyst, which can also reduce the surface charge recombination [66]. Thermodynamic studies further explained that one-photon and one electron charge separation of photocatalyst is accommodated by the co-catalyst to attain four equivalent electron/hole processes (as shown below), which resulted in enhanced photoresponse during solar-driven water splitting by MoOxSy-CoPi photocatalyst [9,29,67].

\[
\begin{align*}
H_2O + \text{hv} & \rightarrow \text{O}^*_2 + e^- + H^+ \\
\text{O}^*_2 + \text{H}_2O & \rightarrow \text{OOH}^*_2 + e^- + H^+ \\
\text{OOH}^*_2 & \rightarrow \text{O}_2 + e^- + H^+ \\
\text{H}_2O + \text{hv} & \rightarrow \text{H}_2 + \text{O}^*_2
\end{align*}
\]

(where* represents, the active surface sites where OH and O species can interact during PEC-WS process)

These observations indicate better PEC-WS recital of MoOxSy-CoPi nanoflowers composite material. Where MoOxSy showed effective light sensitivity by the integration of CoPi with it via a hydrothermally approach that ultimately resulted in improved photocurrent densities [9,29,67]. These observations indicate better PEC-WS recital of MoOxSy-CoPi nanoflowers composite material. Where MoOxSy showed effective light sensitivity by the integration of CoPi with it via a hydrothermal approach that ultimately resulted in improved photocurrent densities.

**Conclusion and future perspectives.**

Herein, we presented a facile synthesis, characterization, and PEC-WS of novel MoO$_x$S$_y$-CoPi photocatalyst. The key features of our strategy include in situ hydrothermal reactions between CoPi and growing MoO$_x$S$_y$ nanoflowers. The MoO$_x$S$_y$-CoPi photocatalyst showed enhanced PEC-WS properties in comparison to bare CoPi and MoO$_x$S$_y$ XRD studies of MoO$_x$S$_y$-CoPi photocatalyst and CoPi materials confirm orthorhombic crystalline phases. However, XPS analysis predicted the most plausible elemental states of each constituent present in MoO$_x$S$_y$-CoPi photocatalyst. The FTIR and Raman spectroscopy’s further comprehended its XRD findings by identifying specific fingerprint regions for particular elements and groups present in the photocatalyst. The results of XPS and Raman are supported by HR-TEM, which confirms the successful incorporation of CoPi into the flakes of MoO$_x$S$_y$ nanoflowers in the MoO$_x$S$_y$-CoPi nanocomposite. FE-SEM studies revealed nanoflowers like morphology showing curved nanopetals with the thickness (>10 nm) and variable internal spaces (200–400 nm). The comparable optical properties, i.e., diffused reflectance spectroscopy, showed a variation in band-gap energy (2.44 eV) for MoO$_x$S$_y$-CoPi compared to bare MoO$_x$S$_y$ and CoPi. The MoO$_x$S$_y$-CoPi/FTO showed significant photocurrent density in
comparison to MoOxSy/FTO and Co/Pt/FTO. The positive photocurrent is related to photoinduced oxygen evolution reaction (OER). The enhancement of the photocurrent in the case of MoOxSy-CoPi is attributed to the improved charge separation and transfer, low electron-hole recombination, photo corrosion stability, and better reaction kinetics. Henceforth, MoOxSy-CoPi nanocomposite is considered as a promising material for solar-driven applications. Nonetheless, its PEC-WS and optoelectrical characteristic can be further tuned by doping it with different transition metals or co-catalysts. Thus, the MoOxSy-CoPi nanoflowers composite material is a potential candidate to target efficient solar-driven water splitting reactions in similar energy harnessing applications.

Compliance with Ethics Requirements

This article does not contain any studies with human or animal subjects.

Declaration of Competing Interest

The authors declare that they have no competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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