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Alkali metal-assisted nucleation and growth of stable 1T/2H MoS₂ for the hydrogen evolution reaction

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The present work investigates the extent of mixed phase 1T/2H MoS₂ formation as the same precursor solution was maintained for 0 to 8 days in an alkaline medium. To confirm the mixed phase of MoS₂, powder X-ray diffraction (XRD), Raman spectroscopy, scanning electron microscopy, and transmission spectroscopy measurements were performed. The major XRD peak of the 1T phase for the 2θ value was at 13.8° and was indexed as the (002) plane along with the 2H phase, and Raman spectra $E_{2g}^1 = 381\text{ cm}^{-1}$ and $A_{1g} = 405\text{ cm}^{-1}$, including extra peaks J_1 , J_2 , E_{1g} , and J_3 . The high-resolution transmission electron spectroscopy interplanar spacing of 1T-MoS₂ and 2H-MoS₂ was 0.246 nm and 0.688 nm, respectively. Our studies indicate that the alkali metal Na⁺ plays a crucial role *via* intercalation, which expanded the inter-layer spacing and increased the catalytic surface area and conductivity, hence affecting the hydrogen evolution reaction (HER) efficiency. The HER showed an overpotential @10 mA cm⁻², and Tafel slope of 260 mV, 65 mV dec⁻¹ for the instantly prepared 0-MoS₂ sample compared to longer duration samples of 2 to 8 days.

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1. Introduction

The widespread reliance on fossil fuels has severely polluted the atmosphere, driving an urgent need for cleaner, renewable sources of energy. One promising alternative is hydrogen fuel cell technology.¹ These cells generate electrical energy through an electrochemical reaction between hydrogen (H₂) and oxygen (O₂) gases, and the amount of energy produced directly correlates to the quantity of these gases used.² Unfortunately, the naturally occurring concentration of hydrogen gas in the environment is insufficient to power large-scale hydrogen fuel cells, but electrochemical cells can be employed to produce the necessary H₂. In a typical setup, these cells comprise three electrodes: a reference electrode, a counter electrode, and a working electrode (where hydrogen generation occurs). Highly effective platinum (Pt) is traditionally used as the working electrode (cathode) to catalyze the hydrogen evolution reaction (HER), where protons are reduced to hydrogen.

The high cost of platinum, motivates the search for more affordable yet efficient alternatives,^{3,4} such as non-noble metals (*viz.*, metal-carbides, phosphides, chalcogenides),^{5,6} metal oxides (*viz.*, spinel oxide, perovskite oxide),⁷ and composite hybrid

materials.^{8,9} Two-dimensional transition metal dichalcogenides (2D-TMDs) are widely explored materials in this field because they can be used in the HER, and they are also commonly applied in sensors,¹⁰ electronic devices,¹¹ optoelectronics, and energy storage.^{12,13} 2D-TMDs (*viz.*, molybdenum disulfide (MoS₂), tungsten disulfide (WS₂), and niobium disulfide (NbS₂)¹⁴) are created by sandwiching a single atomic layer of transition metal atoms between two layers of sulphur atoms.¹⁵

In recent years, 2D-MoS₂-based materials have shown remarkable promise in lowering the cost of hydrogen production while demonstrating excellent efficiency and stability in the HER.¹⁶ Efficient catalysts are crucial for enhancing the kinetics of this reaction and enabling applications in energy conversion and storage systems. MoS₂ possesses several unique properties that make it attractive as an HER catalyst. Its layered structure provides a large surface area, which increases the availability of active sites for the reaction. The catalytic activity of MoS₂ stems from its electronic structure, which enables efficient charge transfer during the HER in comparison to traditional catalysts, *viz.*, platinum.

2D-MoS₂ exists in three distinct phases (2H, 1T, and 3R),^{17–19} as well as a mixed phase (1T/2H). The 2H and 3R phases of MoS₂ are thermodynamically more stable at high temperatures, which can be achieved by physical and chemical methods,^{20–22} but higher HER efficiency was achieved only in a limited synthesis process. Compared to the pure semiconductor 2H-MoS₂, whose active sites are primarily on its edges, the

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metallic 1T phase increases active sites on both edges and basal surfaces. Zhang *et al.* (2020) reported that an increase in the percentage of the 1T phase leads to a corresponding increase in the efficiency of the HER. This enhancement is attributed to the higher basal plane catalytic activity of the sulfur sites in the 1T phase for hydrogen binding.²³ The 1T phase exhibits several advantages, but its instability, makes attaining an octahedral structure challenging.

Researchers have rigorously worked to prepare the 1T phase using hydrothermal and traditional intercalation/exfoliation ($(\text{CH}_3)_4\text{N}^+$, Li) techniques. Hongqiang *et al.* created exfoliated 1T-MoS₂ nanosheets using sonication and $(\text{CH}_3)_4\text{N}^+$ ion intercalation. The intercalated ions expanded the interlayer spacing and decreased the van der Waals force between the layers, which exfoliated it into the 1T phase.²⁴ Mengyao *et al.* synthesized 1T-MoS₂ with superior ambient stability using a hydrothermal method *via* hydrazine intercalation.²⁵ In the mixed phase, the interaction with the surrounding 2H phase stabilized the 1T phase, allowing it to contribute its beneficial properties without compromising the structure. Browne *et al.* developed a partially converted 1T phase from 2H using a two-step chemical vapor deposition (CVD) method, followed by a five-day *tert*-butyllithium immersion treatment.²⁶ The presence of the metallic 1T phase within the mixed phase contributed to its overall high electrical conductivity compared to pure semiconducting 2H-MoS₂. This improves conductivity, which exhibits wide applications for acquiring efficient charge transfer in electrocatalytic hydrogen evolution.

A wet chemical approach is one of the synthesis techniques studied to determine greater insight on the nucleation and development of 2D-MoS₂. It also provides an alternative and adaptable method to produce a mixed phase (1T/2H) of bulk and atomically thin layers of molybdenum disulphide.²⁷ This is a facile method that can be used to manufacture 2D-MoS₂ (1T/2H), in comparison with vapour deposition techniques, and thus, it is suited for extensive applications and large-scale production.^{28,29} In 2D-MoS₂ nucleation and growth by wet chemical methods, MoS₂ nuclei or seeds are formed in a solution. This is accomplished by combining precursors such as molybdenum salts (*e.g.*, ammonium molybdate) with sulphur sources (*e.g.*, thiourea) in a solvent, or by employing alternative techniques such as hydrothermal or solvothermal reactions.³⁰

These precursors undergo a reaction to generate MoS₂ nuclei, which serve as a basis for the growth of MoS₂ particles under optimized experimental parameters for bulk samples, as previously reported.³¹ The MoS₂ nuclei serve as seeds for the deposition of additional MoS₂ layers from solution, and these layers laterally expand on the nuclei as the reaction advances, resulting in a continuous 2D-MoS₂ film.³² Temperature, reaction time, and precursor concentrations can be optimized to influence the growth kinetics and quality of the final MoS₂ layers.^{33,34} Also, with thin films, a thin seeding layer (Na^+) is deposited prior to MoS₂ layer deposition. This Na layer affects the growth of thin films, and is advantageous and disadvantageous for MoS₂ samples.

Obtaining a mixture of 1T and 2H phases (1T/2H) of MoS₂ using chemical vapor deposition (CVD) is challenging due to

the high-temperature requirements for CVD (typically 600–800 °C), which destabilizes the desired 1T phase. Browne *et al.* (2019) achieved the synthesis of 1T/2H MoS₂ films through a dual CVD method followed by a lengthy (five-day) immersion treatment with *tert*-butyllithium, but this approach resulted in a low yield and extended processing time.²⁶ A more efficient method for 1T/2H MoS₂ synthesis was developed by Hongmei *et al.*, who employed a two-step hydrothermal approach. This involved a 20-hour reaction at 200 °C, followed by chemical reduction with sodium borohydride (NaBH_4) under an argon atmosphere.³⁵ Another approach, reported by Huanran Li *et al.* (2022), involved treating as-synthesized MoS₂ with sodium hydroxide (NaOH) to induce the formation of the mixed 1T/2H phase.³⁶ The hydrothermal method is conducted at a temperature of 130–250 °C in aqueous solution with significant vapor pressure (0.3–4 MPa).³⁷ This approach promotes improved crystallinity, which is required for catalytic activity.

Herein, the present work focused on determining (i) how variation of the synthesis parameters, *viz.*, time, temperature, and pressure regulates the HER activity, whereby the wet chemical synthesis approach was adopted for experimentation, and (ii) how alkali metal ion insertion affects the catalytic activity involved in the HER. Hence, we report a wet-chemical synthesis method at 90 °C for 10 min, followed by filtration and annealing under atmospheric pressure for the stable 1T/2H-MoS₂ phase. Mixed phase 1T/2H MoS₂ was instantly formed from the precursor solution under normal atmospheric pressure. For a qualitative understanding of the nucleation seeding and growth of the MoS₂ phase formation, the precursor was maintained in the same solution and then collected after regular intervals of 0, 2, 4, 6, and 8 days. Then, the solution was filtered, and the residue powder was annealed at 450 °C for 1 h in a sulphur atmosphere to form crystalline MoS₂.

MoS₂ phase samples were characterized by different structural and morphological techniques, and electrocatalytic HER applications were studied. We observed that crystallite size varied (increasing from 0-MoS₂ to 4-MoS₂, and then decreasing for 8-MoS₂) over time, thereby regulating the HER activity of the samples. Among all samples, the highest HER efficiency in terms of current density (2.07 mA cm⁻²), onset potential (96 mV), and Tafel slope (65 mV dec⁻¹) was for 0-MoS₂. Furthermore, Na⁺ ion insertion widened the interlayer spacing of metallic 1T/2H MoS₂, and thus enhanced the HER performance of the samples.

2. Experimental section

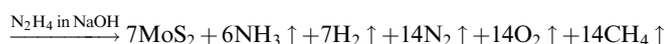
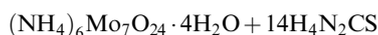
2.1 Synthesis of MoS₂ powder

The 2D-MoS₂ nanosheets were prepared using the wet chemical method^{38,39} and Mo and S as precursors, with reaction time intervals of 0, 2, 4, 6 and 8 days. The precursors were 30 mM of ammonium heptamolybdate ($(\text{NH}_4)_6\text{Mo}_7\text{O}_{24}\cdot\text{H}_2\text{O}$; 99% purity, AR, SRL) for the Mo source, and 0.5 M of thiourea ($\text{N}_2\text{H}_4\text{CS}$; 99% purity, AR, SRL) for the sulphur source, and these were added to 1 M of hydrazine hydrate (N_2H_4 ; 80% extrapure) in basic NaOH medium (pH 10). Initially, the precursor solution



was mixed at room temperature and heated to 90 °C (10 min) on a hot plate for the nucleation of MoS₂.

The prepared solution was filtered after varied reaction times for controlling the growth of MoS₂ (0, 2, 4, 6, and 8 days), with transfer from one filter to another filter solution. The instant filtrate, *i.e.*, the first powder residue product, was named 0-MoS₂ (0-day). The filtered solution was undisturbed and settled so that the next batch of MoS₂ powder could be created for different durations. After two days, the first solution was filtered, and the collected residue powder sample was named 2-MoS₂ (2-days). The same procedure was repeated after 4 days, 6 days, and 8 days, and the solutions were named 4-MoS₂, 6-MoS₂, and 8-MoS₂ respectively. All the residue powder samples (0-MoS₂, 2-MoS₂, 4-MoS₂, 6-MoS₂, and 8-MoS₂) were annealed under a sulphur atmosphere at 450 °C for 1 h in a tubular furnace with the flow rate of carrier Argon (Ar) gas at 100 standard cubic centimeters per minute (sccm). For annealing, the residue powder samples (in a ceramic boat), and 2 g sulphur (in a brass holder) were maintained 10 cm apart in a quartz tube, as illustrated in Fig. 1. The formation of MoS₂ can be written by chemical reaction as:



2.2 Characterization of MoS₂ powder

Powder X-ray diffraction (PXRD) measurement of MoS₂ powder samples was performed using a D8-Bruker (CSR-Indore) instrument with Cu-K α (1.54178 Å) radiation as the X-ray source in the range of 5–80° (two theta values). Raman spectra were recorded using a LabRam excited with a 473 nm laser source. The morphology of the sample was examined using field emission scanning electron microscopy (FESEM) with energy dispersive X-ray spectroscopy (EDS) mapping, carried out on a Nova Nano 450 with 1 µm resolution, and acceleration voltage

of 20 kV. The atomic structure of MoS₂ was examined by high-resolution transmission electron microscopy (HRTEM, JEM F-200).

2.3 Electrochemical measurements

For HER characterization, the electrochemical measurements were studied using linear sweep voltammetry (LSV), Tafel slope, and electrochemical impedance spectroscopy (EIS) at ambient conditions in 0.5 M H₂SO₄ electrolyte solution using a CHI electrochemical workstation with a three-electrode cell system.⁴⁰ The MoS₂ catalyst was deposited on a glassy carbon electrode (GCE) and acted as the working electrode (WE), while platinum wire (Pt) and a calomel electrode were the counter electrode and reference electrode, respectively. The WE preparation was carried out using a 80:10:10 ratio of catalyst, graphene, and polyvinylidene difluoride (PVDF) binder dissolved in *N*-methyl pyrrolidone (NMP) to form a homogenous ink solution. Then, the black ink was deposited on a 3-mm-diameter GCE (loading 0.004 g cm⁻²) using a drop-cast method. At room temperature, the properties of the HER were evaluated using linear sweep voltammetry (LSV) from 0 to -0.8 V (V vs. RHE) at a scan rate of 5 mV s⁻¹. The following equation was applied to assign all potential contours to the reversible hydrogen electrode (RHE) according to the Nernst equation:

$$E(V) \text{ versus RHE} = E(V) \text{ versus Hg/HgCl} + 0.242 \text{ V} + 0.059 \text{ pH}$$

where $E(V)$ vs. RHE is equal to $E(V)$ vs. Hg/HgCl along with 0.242 V in 0.5 M H₂SO₄. EIS was used to characterize the liquid impedance and capacitance of the electrocatalysts. The impedance was measured with an induced voltage magnitude of 1 to 10⁶ Hz.

3. Results and discussion

Analysis of the powder X-ray diffraction (PXRD) pattern of mixed-phase 1T/2H-MoS₂ samples (as shown in Fig. 2(a))

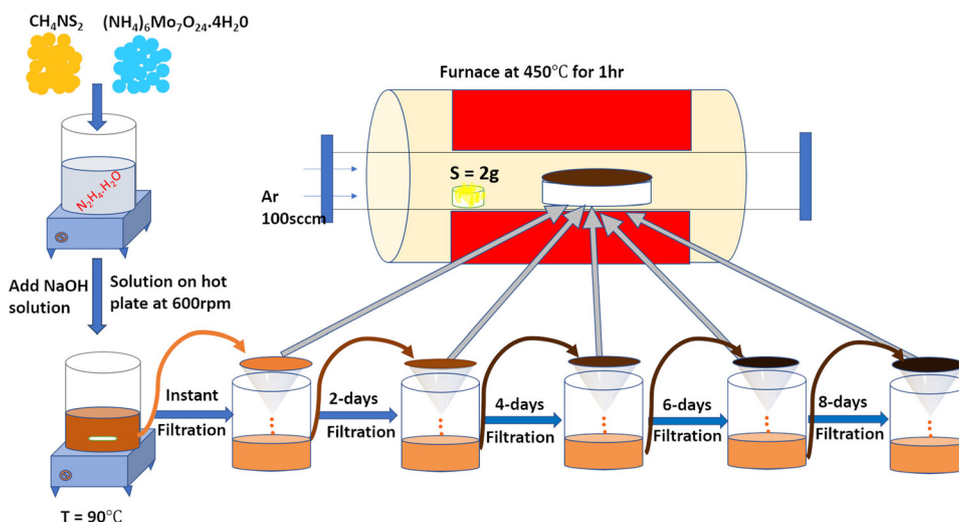


Fig. 1 Schematic representation of the experimental setup: A wet chemical method was used to prepare MoS₂ material by varying the duration time. Further annealing was conducted under a sulfur atmosphere at 450 °C for 1 hour.



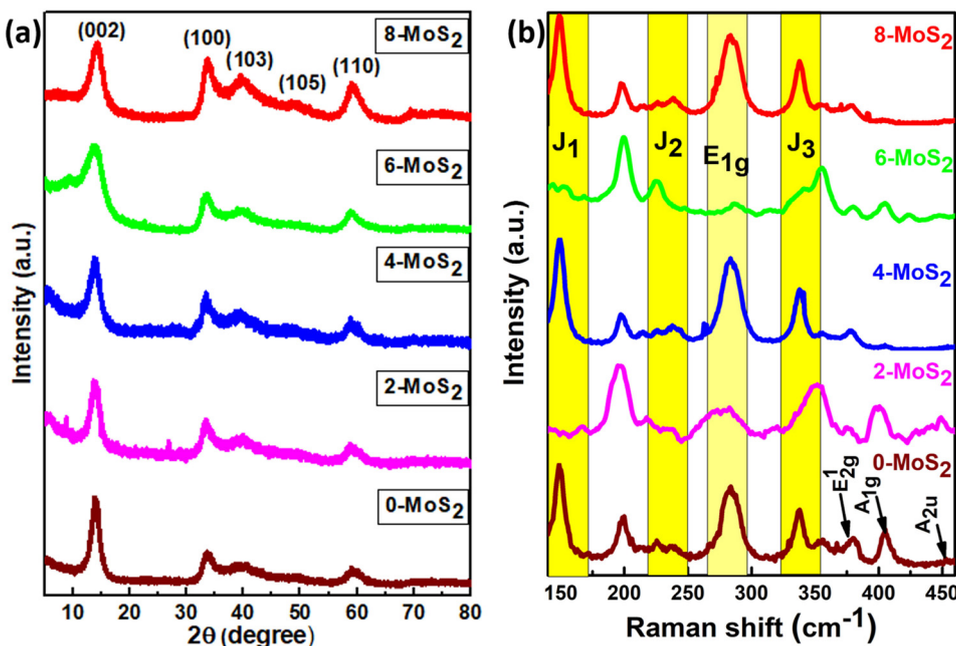


Fig. 2 (a) Corresponds to the X-ray diffraction pattern and (b) Raman spectra of the prepared MoS₂ samples with duration time of 0, 2, 4, 6, and 8 days.

reveals a key distinction compared to the pattern of bulk 2H-MoS₂. This difference lies in the behavior of the characteristic (002) peak at 14.41° (2θ) for pristine 2H-MoS₂. Also from the XRD data, the peak shifted to a lower 2θ angle (002) peak at 13.86°,¹⁸ and this signified a crucial structural change within the material. The (002) peak corresponds to the interlayer spacing of 0.61 nm between the MoS₂ sheets. When the peak shifts, this indicates that the interlayer spacing is no longer uniform.

In the current study, the presence of a lower angle peak shift suggests an expansion of the interlayer spacing, which probably resulted from the Na⁺ intercalation between the layers (discussed further in the HRTEM analysis) compared to pure 2H-MoS₂. The Na⁺ intercalated between the layers, as reported by Nuwan *et al.* (2017), expanded the gap between the layers to 1.2 nm, and as a result, the (002) peaks shifted to a lower angle at low-temperature synthesis, suggesting the 1T phase.⁴¹ Also, Huanran *et al.* (2022) reported that the concentration of Na⁺ decreased when the hydrothermal temperature at 180 °C increased. Hence, the gap was decreased to 0.65 nm causing the (002) peaks to shift to a higher angle, suggesting the emergence of a mixed phase (1T/2H).³⁶

In this work, the sample was annealed at the high temperature of 450 °C, and the gap between the layers was 0.688 nm. This expansion is a well-established signature of the presence of the 1T phase within the MoS₂ lattice.⁴² The overall XRD pattern exhibited the characteristic peaks associated with the hexagonal and trigonal crystalline phases of MoS₂ at 33.36° (100), 39.9° (103), 49° (105), and 59° (110) 2θ values consistent with JCPDS No. 37-1492. The lower angle shift of the (002) peaks provides crucial evidence for the successful incorporation of the 1T phase within the semiconductor 2H phase MoS₂ lattice,^{39,43} and no additional peaks were observed for all MoS₂

samples. The increased interlayer spacing and phase transformation due to the alkali metals increased the catalytically active surface area and movement of electrons within the MoS₂ lattice. This led to increased conductivity and greater catalytic activity. The crystallite size of the prepared MoS₂ nanoparticles was calculated using Scherrer's formula:⁴⁴

$$D = \frac{k\lambda}{\beta \cos \theta}$$

where β denotes the full width at half maximum (FWHM) value, θ denotes the angle of diffraction, and λ denotes the wavelength of an X-ray source of copper (Cu) metal. Table 1 shows that the calculated crystallite size of MoS₂ material varied over different time durations. The size for the instantly filtered (after being removed from the hot plate at 90 °C and annealed) material 0-MoS₂ was 17.7 nm. For the sample filtered after two days, 2-MoS₂, which was undisturbed at room temperature, the size increased to 20.3 nm compared to 0-MoS₂. The longer the reaction duration, the more time elapsed for intercalation of additional Na⁺ between the layers. Hence, the crystallite size increased to 37.97 nm after 4 days (4-MoS₂), but then decreased to 23.45 and 20.07 nm for 6-MoS₂ and 8-MoS₂, respectively. The nucleation seed of MoS₂ was developed at 90 °C on a hot plate,³¹ and because phase particles were in the solution, the crystal

Table 1 Crystallite size of prepared MoS₂

Sample	Average D (nm) value
0-MoS ₂	17.7
2-MoS ₂	20.3
4-MoS ₂	37.97
6-MoS ₂	23.45
8-MoS ₂	20.07



continuously grew over the reaction duration time. MoS₂ particles were further obtained as the number of days increased. According to the LaMer mechanism,⁴⁵ nucleation and growth depend on the concentration of the precursors, and the filtrate solution of mixed MoS₂ precursors assists in the formation of new MoS₂ particles.⁴⁶ Therefore, MoS₂ growth occurred in different batches of filtration of the same solution with varying time duration, until both precursors were exhausted of MoS₂. Hence, the crystallite size increased for 4-MoS₂.

The structural phase of all synthesized MoS₂ material was further analyzed by Raman spectra, and Fig. 2b shows Raman modes of the 1T/2H mixed phase of MoS₂. From the figure, Raman spectra were found in seven peaks at 148, 238, 283, 337, 381, 405, and 452 cm⁻¹ representing the J₁, J₂, E_{1g}, J₃, E_{2g}¹, A_{1g}, and A_{2u} phonon modes of all synthesized MoS₂, respectively. Out of seven phonon modes, two modes are the E_{2g}¹ in-plane vibration of Mo-S, and A_{1g} is an out-of-plane vibration of S-S at 381 cm⁻¹ and 405 cm⁻¹ of the 2H phase with the A_{2u} phonon mode at 452 cm⁻¹ (0, 4, 6, 8-MoS₂). However, for 2-MoS₂, the modes slightly shifted to the higher energy side (375 cm⁻¹ and 399 cm⁻¹). The Raman mode (A_{1g}) intensity of MoS₂ samples decreased at 405 cm⁻¹ as the number of days passed from 0 to 6 days and disappeared for 8-MoS₂. This is because of the absence of Raman phonon mode (A_{1g}), which is a relatively out-of-plane (S-S) bond vibration⁴⁷ for 8-MoS₂. The trigonal (1T) phase of all as-synthesized MoS₂ was significant because it exhibited additional peaks at 148, 238, 283, and 337 cm⁻¹, which corresponded to J₁, J₂, E_{1g}, and J₃, respectively.^{25,43} However, the 2- and 6-MoS₂ sample exhibited J₁, J₂, and J₃ peaks, which are broad with very weak intensity,^{48,49} while the J₃ peak of 2-MoS₂ slightly shifted to the lower energy side of 350 cm⁻¹. The E_{1g} band was attributed to the octahedral coordination of Mo in 1T MoS₂.

The surface morphologies and elemental distribution of MoS₂ samples were examined using FESEM and EDS mapping, respectively. In Fig. 3(a)–(c), a FESEM image shows all the

prepared MoS₂ samples, and the inset depicts the change in solution colour from 0 days to 4 days. There was a wine-red colour to the initial solution (after the first filtration, which changed to brown after two days, and darkened further), and with an increase in the number of days (4, 6, and 8 days), the solution changed to dark brown. A spherical nanoparticle shape was found for 0-MoS₂, 2-MoS₂, and 4-MoS₂, as shown in Fig. 4. The dimension compared to 0-MoS₂ increased with increasing reaction time duration (2 and 4 days), and can be said that the MoS₂ particles were involved in nucleation. This nucleation further increased while the reaction duration proceeded, and thus additional agglomeration occurred, which led to an increase in crystallite size.

The elemental composition of MoS₂ was determined using EDS, as demonstrated in Fig. 4(a)–(h). Extracted images show strong peaks, confirming the presence of Mo, S, and weak peaks of Na and O. However, the elemental mapping of 0-MoS₂ shown in Fig. 4(a)–(e) indicates that Mo and S were uniformly dispersed, as shown in Fig. 4(b) in red, and Fig. 4(c) in green, respectively. Nevertheless, some O and Na were detected (shown in Fig. 4(d) and (e)), where Na⁺ was observed due to the addition of NaOH to the precursor solution to maintain the pH value, while O comes from the atmosphere.⁵⁰ Each Mo and S element was evenly distributed in the 0-MoS₂, 2-MoS₂, and 4-MoS₂ compounds (0, 2, and 4 days). However, when the reaction duration time advanced, the elemental mapping of Na wt% increased. The Na elements were dense in the case of the larger crystallite size for 4-MoS₂. As a result of this, Na supports enhanced the crystallinity of the samples.⁵¹ However, as the particle growth proceeded, the saturation of the Na bonds decreased, which lowered the energy barrier and increased the growth rate.^{52,53} The increase in the number of days provided the solution sufficient time to create a greater yield with large particle sizes.

HRTEM analysis was carried out on 0-MoS₂, to examine phase crystallinity and dimensions, as shown in Fig. 5(a). The nanosheets were formed by MoS₂ layer by layer, as shown in Fig. 5(b), and the *d*-spacing between the interplane was 0.688 nm and corresponds to the (002) plane of 0-MoS₂. Generally, bulk 2H-MoS₂ has an inter-layer distance of 0.61 nm.^{38,39} However, the interplanar spacing of the prepared 0-MoS₂ sample was slightly higher (0.688 nm) than that of the bulk 2H-MoS₂, which is consistent with our XRD data, as displayed in Fig. 2(a). The 1T/2H-MoS₂ crystal surface was studied to determine discrimination between the two phases using the high magnification of HRTEM, as shown in Fig. 5(c).

The enlarged image of the MoS₂ 2H phase shown in Fig. 5(d) (represented by the yellow ellipses in Fig. 5(c)) displays lattice fringes as a honeycomb structure (trigonal prismatic structure). Also in Fig. 5(e), the magnified white ellipse region in Fig. 5(c) shows the 1T phase of the lattice fringes with a trigonal structure (octahedral structure).^{25,35} The interplanar spacing of the 1T phase is 0.246 nm, as shown in Fig. 5(c), and indexed as the (102) plane. Selected area electron diffraction (SAED) was performed to evaluate the material's crystallinity, and also support the matching crystallinity plane,⁶ as shown in Fig. 5(f). It consisted of four blurred diffraction rings with multiple bright spots

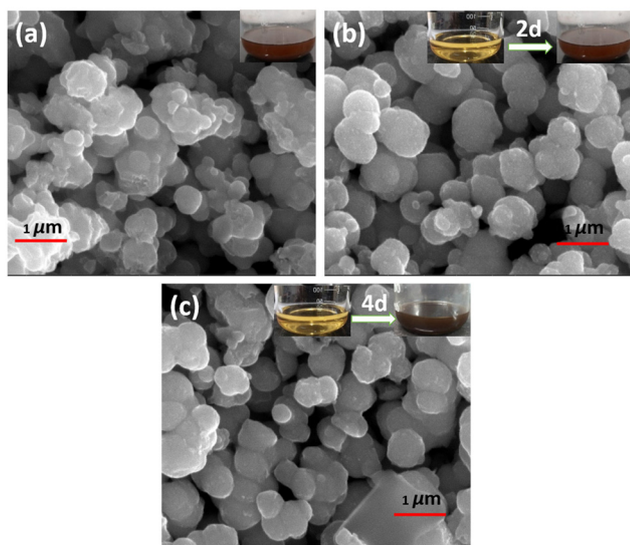


Fig. 3 FESEM images of (a) 0-MoS₂, (b) after 2 days, and (c) after 4 days. The inset shows solution colour changes as time increased from 0 to 4 days.



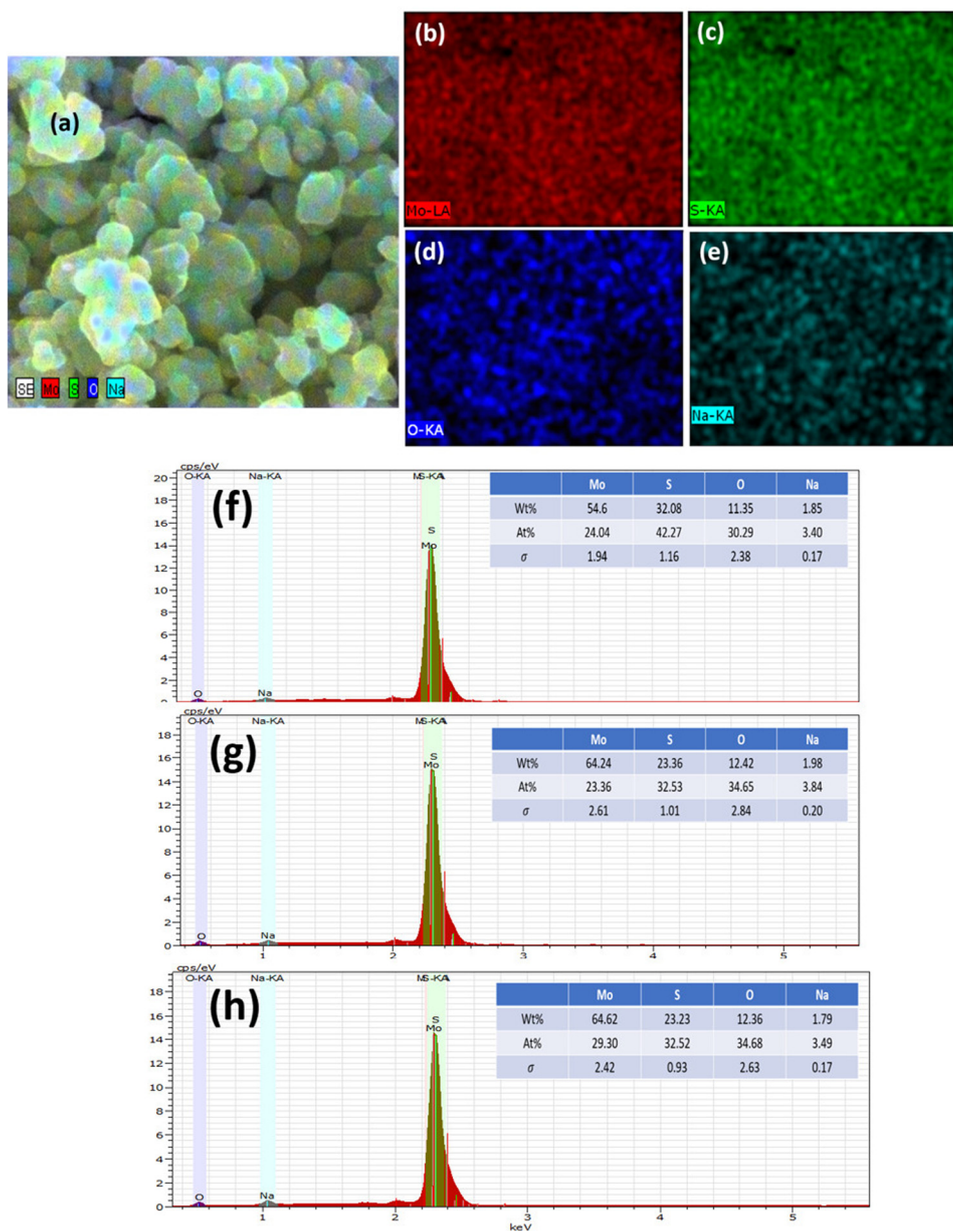


Fig. 4 Elemental mapping. (a) FESEM image of the 0-MoS₂ sample, (b) molybdenum element in red, (c) sulphur element in green, (d) oxygen in blue, and (e) sodium in light blue. Energy dispersive X-ray spectroscopy (EDS) study of (f) 0-MoS₂, (g) 2-MoS₂, and (h) 4-MoS₂ samples.

that were indexed as (100), (103), (110), and (108) accordingly (JCPDS No. 37-1492).⁵⁴ The SAED image confirmed the polycrystalline behaviour of the 0-MoS₂ sample.⁵⁵

3.1 Electrocatalytic application

The sample polarization curves were examined using linear sweep voltammetry (LSV) with a scan rate of 5 mV s⁻¹ in 0.5 M of H₂SO₄ electrolyte solution under ambient conditions, as shown in Fig. 6(a). The Pt electrode (0.5 mm in diameter and 32 mm in length) acted as a reference electrode, and all MoS₂ samples showed catalytic activity towards the HER with a high current density up to 110 mA cm⁻² and low overpotential. For the performance of samples in the HER, onset potentials were

between -90 to -270 mV, which can be attributed to the nanoparticle structure assembled by the nanosheets. It is widely accepted that the HER active sites are located on the edge S-sites, and hence, changing the microstructure of MoS₂ to obtain additional edge planes enhances the HER activity. Among the prepared set of catalyst samples of the present work, the 0-MoS₂ shows the best catalytic activity due to the smaller crystallite size towards the HER, with positive onset potential at -96 mV and a current density of 10 mA cm⁻² at -260 mV overpotential. This indicates that Na⁺ intercalation can promote the stabilization of the 1T phase, and this phase enhancement is crucial for the HER. However, it also increases the overall electrical conductivity and facilitates electron transport during the HER process.



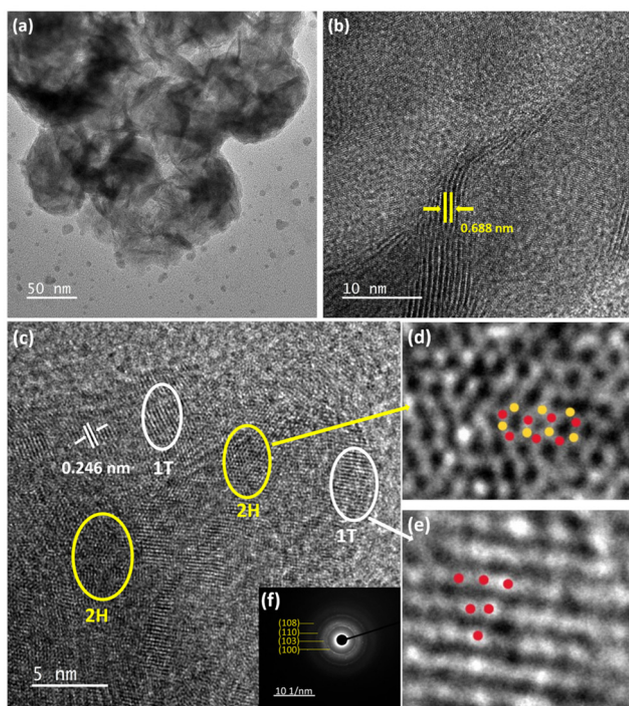
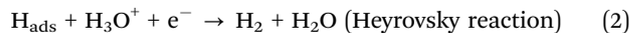


Fig. 5 (a) Transmission electron microscopy image, (b) high-resolution transmission electron microscopy (HRTEM) image of 0-MoS₂, (c) low magnification HRTEM of 1T/2H-MoS₂, (d) HRTEM image of region magnified by the yellow ellipse in (a) and (e) white region magnified in (a) and (f) selected area electron diffraction pattern of 0-MoS₂.

The LSV value of the characterized 0-MoS₂ sample is equal to earlier reported values for MoS₂-based catalytic materials determined using CVD, intercalation, and hydrothermal techniques, as listed in Table 2. However, materials with different reaction duration times, *i.e.*, 2-MoS₂, 4-MoS₂, 6-MoS₂, and 8-MoS₂, exhibited different catalytic activity due to their intrinsic structure and the synthesized experimental parameters. Fig. 6(a) shows all MoS₂ samples producing HER activity. The overpotential values @10 mA cm⁻² current density were -260 mV, -744 mV, and -560 mV for 0-MoS₂, 2-MoS₂, and 4-MoS₂, respectively. Although there was an increase in the number of days the current density decreased to -8.3 mA cm⁻² and -3.8 mA cm⁻², corresponding to 6-MoS₂ and 8-MoS₂ at -746 mV overpotential, respectively, and 8-MoS₂ sample exhibited the least catalytic activity towards the HER.

The Tafel plots of all samples were derived from the polarization curves mentioned in Fig. 6(b). The Tafel slope is determined by fitting the linear portion of the Tafel plot to the Tafel equation $\eta = b \log |j| + a$, where η denotes overpotential, j denotes current density, b denotes the Tafel slope, and a denotes the intercept. The lower value of the Tafel slope indicates a more optimal chemical kinetic reaction for HER applications with low potential. In Fig. 6b, the smallest Tafel slope of 65 mV dec⁻¹ was obtained for the 0-MoS₂ sample, and the Tafel slopes of 2-MoS₂, 4-MoS₂, 6-MoS₂, and 8-MoS₂ were 303 mV dec⁻¹, 136 mV dec⁻¹, 290 mV dec⁻¹, and 322 mV dec⁻¹, respectively. From this analysis, the smaller Tafel slope of 0-MoS₂ as compared to the others indicates that it

performs the HER with greater efficiency at a lower overpotential. The HER reaction occurred on the surface of the MoS₂ electrode in acidic (0.5 M H₂SO₄) medium, and can be explained by the following mechanism of the Volmer–Heyrovsky reaction, as shown in Fig. 7:



Furthermore, EIS was used to investigate the interface reactions and electrode kinetics during the catalytic HER reaction. Fig. 6c depicts the Nyquist plots of the prepared MoS₂ samples. The Nyquist plot shown in the figure is divided into two sections: a semicircle and a straight line, and these portions relate to the high and low frequency bands, respectively.⁵⁹ The point at which the figure intersects the *x*-axis at higher frequencies denotes the electrolyte resistance (*R*_s) that is potentially independent. The semi-circular arc at the lower frequency end is potentially dependent and reflects the resistance-to-charge transfer (*R*_{ct}) across the electrode–electrolyte interface.⁶⁰ This resistance is connected to the ease with which charges may be separated during the electrochemical process. The charge-transfer resistance for an electrocatalyst with a smaller semicircle diameter is decreased, which indicates increased electrocatalytic activity.⁶¹ It was found that the lowest semicircle was for 4-MoS₂, in comparison to the other samples (0-, 2-, 6-, and 8-MoS₂). The calculated (*R*_{ct}) values of prepared samples 0, 2, 4, 6, and 8-MoS₂ were 6.1, 35.4, 5.6, 14.2, and 18.4 Ω, respectively. 4-MoS₂ conveyed unusually low values due to its exceptional intrinsic conductivity, although the second-lowest *R*_{ct} value was for 0-MoS₂. The most inherent measure of activity for the HER is the exchange current density, *I*_{ex}, which is determined from the EIS.⁶²

$$\text{Exchange current density } (I_{\text{ex}}) = RT/nF\theta$$

where *R* denotes the universal gas constant, *T* denotes the reaction temperature (room temperature), *n* denotes the number of electrons transferred (2), and θ denotes the resistance calculated from the EIS. Exchange current density is a measure of the intrinsic rate of electron transfer at the interface between the electrode and the analyte, or reacting species. Satisfactory electrocatalytic activity is indicated by a high exchange current density for a reaction on a given electrode material. The exchange current density values for all the samples are listed in Table 3. The *I*_{ex} values for all MoS₂ samples are lower than that of 4-MoS₂ (2.28 mA cm⁻²), which has the lowest *R*_{ct} value. However, the *R*_{ct} value increased, and the *I*_{ex} value decreased. Hence, the largest exchange current density is for 4-MoS₂ and 0-MoS₂, and the order of *I*_{ex} is 4-MoS₂ > 0-MoS₂ > 6-MoS₂ > 8-MoS₂ > 2-MoS₂.

Besides HER activity, stability is a crucial parameter for evaluating an advanced electrocatalyst's longevity. A stability test in an acidic medium was performed to assess durability. The time-dependent curve, which was obtained by applying a constant overpotential of 96 mV for 20 000 s of data, as



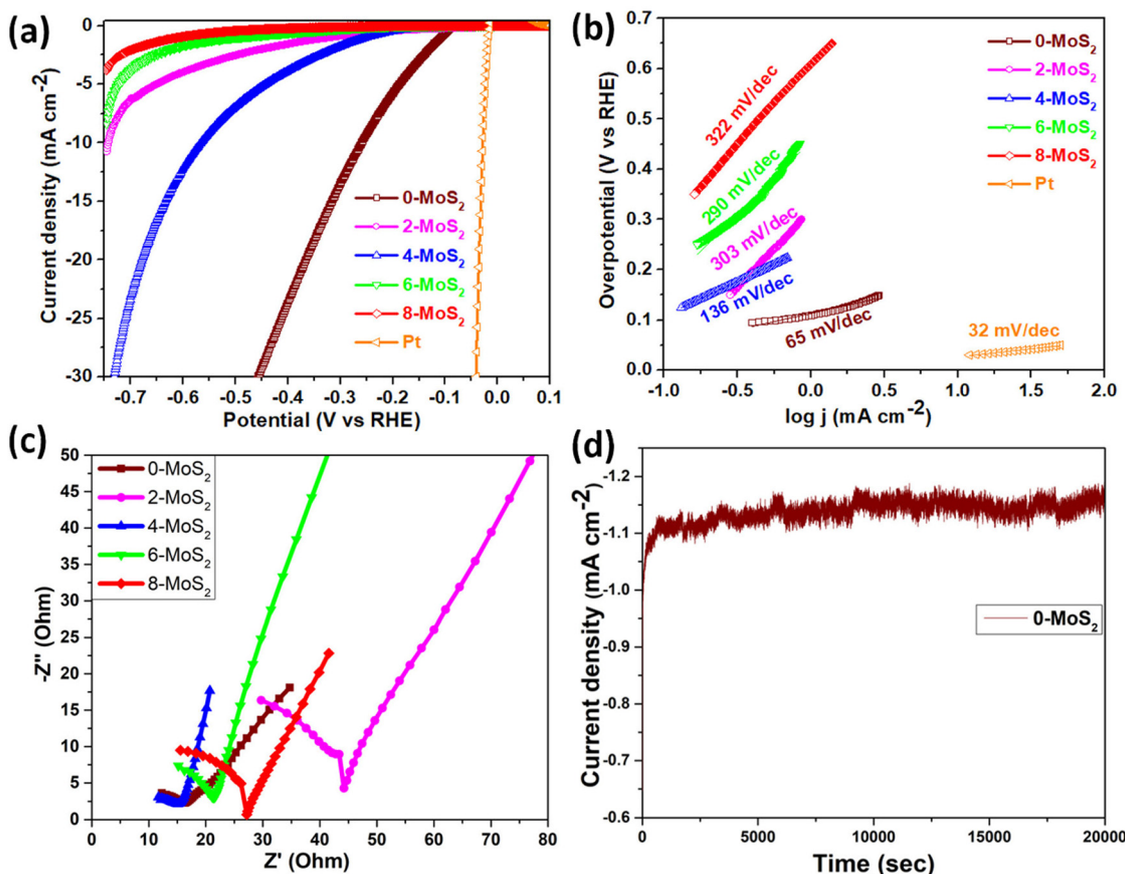


Fig. 6 Electrochemical performance of Pt- and MoS₂-based material. (a) Linear sweep voltammetry (LSV) curve, (b) Tafel slope, (c) electrochemical impedance spectroscopy (EIS) of 0-MoS₂, 2-MoS₂, 4-MoS₂, 6-MoS₂, and 8-MoS₂, and (d) stability test of 0-MoS₂.

Table 2 List of 1T/2H phases of MoS₂-based catalysts for HER applications

Name of the catalyst	Synthesis method	Overpotential (mV) η@ 10 mA cm ⁻²	Tafel slope (mV dec ⁻¹)	Ref.
1T/2H-MoS ₂	CVD, immersion treatment	320	70	26
1T-MoS ₂ (Na-MoS ₂ at 160 °C)	Hydrothermal	255	44	36
1T/2H-MoS ₂ (Na MoS ₂ at 180 °C)	Hydrothermal	271	49	36
1T/2H-MoS ₂ (HF treated)	Hydrothermal	317	90	36
1T@2H-MoS ₂	Exfoliation, and dispersion in PVP	330	70	56
(1T/2H) MoS ₂ scrolle@pt	Exfoliation, rolling up	251	57	57
(1T/2H) MoS ₂ @pt scrolle	Exfoliation, rolling up	125	39	57
1T@2H-MoS ₂ nanosheet 8 h C ⁻¹	Hydrothermal	64	49	27
1T@2H-MoS ₂ -4 h C ⁻¹	Hydrothermal	99	57	27
1T/2H-MoS ₂	Hydrothermal	296	59.6	58
1T-2H-MoS ₂	Solid vapour reaction	280	65	39
1T@2H-MoS ₂	Hydrothermal	270	88	30
1T-MoS ₂ (MoS ₂ -4HZ)	Hydrothermal	230	64	25
1T/2H-MoS ₂ (MoS ₂ -1HZ)	Hydrothermal	284	74	25
1T/2H-MoS ₂	Wet chemical method	260	65	Current work

displayed in Fig. 6d, revealed that a continuous HER process occurred to create molecular hydrogen without any current deterioration.

4. Conclusion

In summary, this work successfully synthesized mixed phase 1T/2H-MoS₂ via a wet chemical synthesis method. To study

nucleation seeding and growth, the reaction was carried out at various durations of time, *i.e.*, 0, 2, 4, 6, and 8 days. It was observed that the crystallite size of MoS₂ increased from 17.7 to 37.9 over a duration of 0 to 4 days, and subsequently decreased to 23.45 and 20 nm after 6 and 8 days, respectively. This observation may be attributed to the growth of MoS₂ particles with a precursor concentration, which gradually decreased with time. This observation was further validated by XRD data using



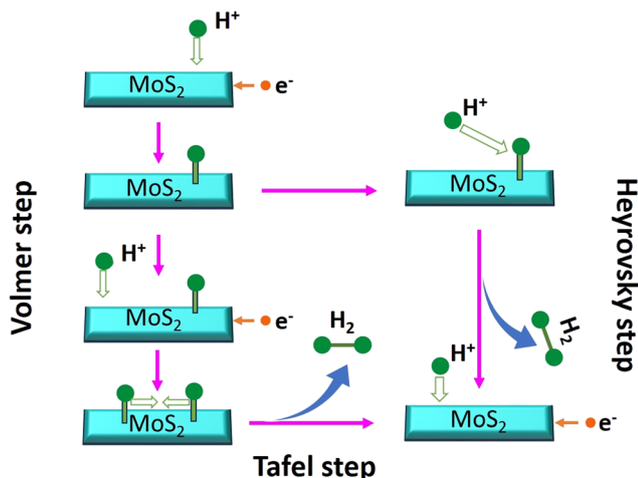


Fig. 7 Schematic representation of the reaction pathway diagram for a catalytic process.

Table 3 Comparison of catalytic parameters of different HER catalysts

Sample	Onset potential (mV)	Overpotential (mV) @ 10 mA cm ⁻²	Tafel (mV dec ⁻¹)	R _s (Ω)	R _{ct} (Ω)	I _{ex} (mA cm ⁻²)
0-MoS ₂	96	260	65	9.543	6.175	2.07
2-MoS ₂	133	744	303	3.124	35.36	0.36
4-MoS ₂	270	560	136	7.917	5.631	2.28
6-MoS ₂	251	—	290	6.144	14.29	0.8
8-MoS ₂	251	—	322	5.94	18.47	0.6

Scherrer's formula. Among the materials studied, 1T/2H 0-MoS₂ exhibited the smallest crystallite size of 17.7 nm and the highest catalytic activity with an onset potential of -96 mV, an exchange current density of 2.07 mA cm⁻², a low Tafel slope of 65 mV dec⁻¹, and a charge transfer resistivity of 6.175 Ω. Moreover, the presence of Na⁺ ions expanded the space between the layers, creating a larger catalytically active surface area and increasing the material's electrical conductivity. Both improved conductivity and increased surface area were beneficial for enhanced electrocatalytic activity. The developed mixed 1T/2H phase 0-MoS₂ exhibited superior performance as an instantly prepared catalyst and can be used in the scalable synthesis of MoS₂ nanocrystals, with promising use in acid electrolytes. Furthermore, the intercalated sodium ions (Na⁺) hold promise for future research in sodium-ion batteries and hydrogen cells. Our data could significantly contribute to the advancement of renewable energy technologies.

Conflicts of interest

There are no conflicts to declare

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