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Electrochemical sulfidation of WS₂ nanoarrays: Strong dependence of hydrogen evolution activity on transition metal sulfide surface composition



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ABSTRACT

The activity of transition metal sulfides for the hydrogen evolution reaction (HER) can be increased by sulfur-enrichment of active metal-sulfide sites. In this report, we investigate the electrochemical sulfidation of atmospherically aged WS₂ nanoarrays with respect to enhancing HER activity. In contrast to MoS₂, it is found that sulfidation diminishes HER activity. Electrochemical and XPS experiments suggest the involvement of insoluble tungsten oxides in the altered HER and electron transfer properties. This demonstrates the strong dependence of the transition metal dichalcogenide (TMD) composition with the successful sulfur incorporation and subsequent HER activity.

1. Introduction

The demand for sustainable sources of electrochemical hydrogen production [1] has triggered the development of the abundant and low-cost TMDs as substitutes to the best performing platinum group metal catalysts for the hydrogen evolution reaction (HER) [2–5].

To improve their HER activity, research has focussed on the preparation of S-rich TMD structures which surpass the 1:2 M:X stoichiometry found in bulk materials [6–8]. Such sulfur enrichment aims to incorporate more bridging S₂²⁻ and terminal S²⁻ moieties into the TMD structure; both consistently reported as being the active sites involved in proton adsorption and desorption [9,10]. Enhanced HER performance has been reported for S-rich structures such as amorphous MoS_{2+x} [11–15] and WS_{2.64} electrodeposited thin films [9], as well as on wet chemical synthesis-prepared MoS_{2+x} [16,17] or MX₃/MX₂ physical mixtures [18]. However, some of the proposed structures exhibit diminished HER performances after atmospheric or electrochemically-induced sulfur depletion [14,18] or impurities presence [19].

We report the use of a one step, room temperature electrochemical sulfidation method initially developed for MoS₂ [20], for sulfur-enriching WS₂. In particular, atmospherically-aged WS₂ nanocone arrays which are of interest due to their enhanced electrocatalytic properties [21]. Changes in the electrocatalytic behaviour are understood via

monitoring surface composition, morphology, and electron transfer properties over a one month period by XPS, SEM, and voltammetric experiments.

2. Materials and methods

2.1. Fabrication

The plasma-etch fabrication method used is based on a literature method [22–24], recently reported in TMDs for electrocatalytic applications [20,21,25].

In short, WS₂ (defect-free, 99.9995% purity, 2D Semiconductors USA) crystals cut into rectangles of approximately 1.5 × 5 mm were affixed to glassy carbon (GC) type 2 stubs (7 mm diameter, 2 mm thick, Alfa Aesar, UK) with carbon tape. A 20 μL mixture of a 216 ± 4 nm diameter polystyrene-latex nanosphere (NS) suspension (3000 Series Nanosphere, 1 wt% in water, Thermo Scientific, UK) with absolute ethanol in a 1:1 vol. ratio was transferred to a silicon wafer (previously cleaned with piranha solution and oxygen plasma) to form a self-assembled, hexagonal close-packed, NS monolayer. The NS monolayer was transferred onto the liquid interface of a water-filled Petri dish containing the TMD-modified GC stubs, and the supernatant extracted with a syringe to promote NS deposition onto the TMD surface.

NS-modified TMDs etching was carried out in an Oxford

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Instruments Plasmalab NGP 80 ICP/RF etcher. Isotropic oxygen plasma etching (40 s, 30 sccm O₂ flow rate, 100 W RF power) was performed as an NS shrinking step. Afterwards, exposure to an anisotropic plasma etching mixture of SF₆/C₄F₈ (20 sccm/30 sccm, 200 W ICP, 20 W RF power) for variable times (15–60 s) was carried out to obtain nanocone arrays with variable aspect ratios. Pressures of 15 mTorr and temperatures of 20 °C were used in both etching steps. SEM micrographs of the electrochemically tested samples before/after sulfidation were acquired to determine the aspect ratio (nanocone height/base diameter), interspacing, and dimensionless radial (*R*, domain radius/nanocone radius) and normal (*Z*, nanocone height/nanocone radius) coordinates [26] of the individual nanostructures in the array (XL 30 SFEG and JEOL 7100 FEG-SEM, 5 kV, tilt angles from 45° to 85°).

2.2. Electrochemical characterization

Electrochemical measurements were carried out with a PC-controlled PGSTAT128N potentiostat (Metrohm Autolab B-V, Netherlands) in a thermostatted three-electrode electrochemical cell (23 ± 2 °C). The electrodes used were: a bright Pt mesh counter (Alfa Aesar Ltd., UK), a double junction saturated Ag/AgCl reference (electrochemical sulfidation experiments, Sigma-Aldrich), a saturated calomel (SCE) reference electrode (HER experiments, BAS Inc., Japan) and TMD-modified GC stubs connected to a rotating disk working electrode (OrigaLys ElectroChem SAS, France).

HER experiments were carried out in a 2 mM HClO₄ (ACS ≥ 70%, Sigma-Aldrich), 0.1 M NaClO₄ (ACS ≥ 98%, Sigma-Aldrich) solution using a range of voltage scan rates (2–1200 mV s⁻¹). Preconditioning of TMD electrodes prior to HER experiments was via 10 cycles from -0.045 to -1.645 V (vs SCE) at a voltage scan rate of 50 mV s⁻¹. Additional capacitance (voltage range -0.2 to 0.2 V vs. NHE, scan rates 10–500 mV s⁻¹) and impedance measurements (voltage range 0 to -1.645 V vs. SCE, frequency range 10⁻¹ to 10⁵ Hz, voltage amplitude 10 mV) were performed alongside all HER measurements to apply roughness factor and iR compensation corrections. HER potentials are referenced versus the normal hydrogen electrode (NHE) by means of Nernstian shift correction ($E_{\text{NHE}} = 0.242 \text{ V} + 0.059 \text{ pH}$).

Investigation of heterogeneous electron transfer (HET) rates was performed under the same conditions as of Pumera et al. for ease of reference [5,13], by acquiring five CVs in 10 mM K₄Fe(CN)₆/K₃Fe(CN)₆ (BioUltra ≥ 99.5%, Sigma-Aldrich) electrolyte supported by a pH 7.2 phosphate buffer (50 mM potassium phosphate monobasic/potassium phosphate dibasic trihydrate, ≥ 99%, Sigma-Aldrich) at scan rates from 10 to 300 mV s⁻¹. Experimental HET rate constants (k_{app}°) were calculated by correlating the peak-to-peak separation of the Fe(CN)₆⁴⁻/Fe(CN)₆³⁻ redox couple with the dimensionless parameter ψ , using the literature methods of Nicholson and Shain ($\psi \approx 0.1$ and $\Delta E_p < 210 \text{ mV}$) [27], and Klinger and Kochi ($\psi < 0.1$ and $\Delta E_p > 210 \text{ mV}$) [28]. A diffusion coefficient of $7.26 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$ for the [Fe(CN)₆]^{4-/3-} redox pair was used in the calculation [29], assuming $\alpha = 0.5$. For the nanoarrays under study, Case 4 diffusion behaviour (1D) was observed [30]. All electrolytes were freshly prepared with ultrapure water (resistivity not < 18.2 MΩ·cm, Millipore Milli-Q Direct 8), and thoroughly purged with N₂ (Oxygen-free grade, BOC Gases plc), and experiments run under a N₂ atmosphere. All glassware was cleaned with a dilute solution of KMnO₄ in concentrated H₂SO₄ followed by rinsing with ultrapure water.

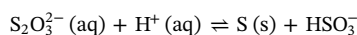
2.3. XPS measurements

A Kratos Axis HSi X-ray photoelectron spectrophotometer (Aston University) fitted with a charge neutraliser and operated using a Mg Kα (1253.6 eV) achromatic radiation, was used to record spectra at a pressure of < 1 × 10⁻⁹ Torr using a spot size of 100 μm. Pass energies used were: 160 eV for survey spectra, and 20 eV for high resolution scans of specific energy regions. Data processing was performed using

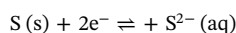
CASA XPS version 2.3.18PR1.0, with spectral energy corrected to the adventitious C 1 s peak at 284.6 eV. Shirley backgrounds were applied to high resolution peaks before being fitted with individual components. W 4f spectra were fit using a FWHM of 0.98 eV, peak area ratios of 4:3, doublet separations of 2.17 eV and Gaussian-Lorentz (30) lineshape, with W 4f_{7/2} WS₂ 2H (32.7 eV), W 4f_{7/2} WS₂ 1 T (31.7 eV) and W 4f_{7/2} WO₂ (33.1 eV) components, whilst the overlapping W 5p_{3/2} feature was fitted with a FWHM of 2 eV, a Gaussian-Lorentz (30) lineshape and a binding energy of 38.1 eV. Sulfur 2p peaks were fitted with a FWHM of 1.03 eV, peak area ratios of 2:1 and a binding energy for S 2p_{3/2} WS₂ of 162.3 eV [31,32].

3. Results and discussion

For reinstating, or improving, the initial HER activity of atmospherically-exposed WS₂ samples, a solution-phase method previously demonstrated for MoS₂ nanoarrays [20] was used. This requires voltage cycling of the TMD samples in a pH 3 solution containing 10 mM Na₂S₂O₃ and 0.1 M Na₂SO₄ whereby S₂O₃²⁻ spontaneously decomposes to form colloidal sulfur [33].



An anodic sweep to fully oxidize the TMD surface, was followed by a cathodic scan to maximize sulfur incorporation onto the TMD surface by reduction



For MoS₂, electrochemically-induced surface oxidation (at $E > +1 \text{ V}$ vs Ag/AgCl) yields the acid-soluble MoO₄²⁻ species [34]. Sulfur incorporation after surface oxidation suggests that MoO₄²⁻ species assist in the overall sulfidation mechanism. For WS₂, the cyclic voltammogram obtained during the sulfidation treatment is similar to that of MoS₂ [20].

The application of this method to WS₂ was evaluated by monitoring HER performance, oxidation state, and electron transfer properties over a one month period following this sulfidation treatment on previously tested, atmospherically aged WS₂ samples. This provided the following observations: (i) freshly sulfidated samples did not necessarily present enhanced HER performances compared with pre-sulfidated samples, and (ii) the samples' HER peak current, after correction for roughness factor, was inferior after a 3-week environmental exposure compared to the pre-sulfidated, atmospherically-exposed state.

Both phenomena can be understood by changes in oxidation state revealed by XPS. For the 31 ± 1 s (*R* = 2, *Z* = 6.4) plasma-etched WS₂ sample, the peak current decays to half of its initial value following sulfidation (Fig. 1a). This is correlated to a decrease in the total S:W ratio (from ca. 2:1 to 1.5:1, see Fig. 2e), and the appearance of WO₂ at the crystal surface up to ca. 24% (W 4f_{7/2}/W 4f_{5/2} doublet lies at binding energies of ca. 33 and 35.2 eV, respectively; Figs. 1c, 2a) [31]. Previous reports on bulk and chemically-exfoliated WS₂ crystals suggest that incorporation of WO₂ is detrimental for the HER [18,35]. In the case of the 31 ± 1 s sample, this is supported by the changed HER kinetics (Tafel slope increase from 100 to 185 mV dec⁻¹, Fig. 3c) and higher onset potentials ($|j_{\text{onset}}|$ from 173 to 207 mV).

Conversely, the 61 ± 1 s plasma-etched WS₂ sample presented higher peak currents (Fig. 1b) and kinetics (Tafel slope 130 vs. initial 210 mV dec⁻¹, Fig. 3d) following sulfidation, despite the decay in the total S:W ratio (from ca. 2:1 to 1.88:1, see Fig. 2f) and the 14% increase in surface WO₂ content (Figs. 1d and 2c). This initially non-linear trend is found to be linked to the S:W ratio, if calculated solely using the W⁴⁺ XPS components characteristic of WS₂. Sulfur-rich S:W ratios promote enhanced HER performance and vice versa. Maximum peak currents coincide with the highest sulfur-to-metal ratios for both 31 ± 1 s (S:W = 2.08:1, $j_p \approx 9 \text{ mA cm}^{-2}$, day 8) and 61 ± 1 s (S:W = 2.18:1, $j \approx 1.6 \text{ mA cm}^{-2}$, freshly sulfidated) samples. After these peak values, both post-sulfidated 31 ± 1 s and 61 ± 1 s etched samples exhibited

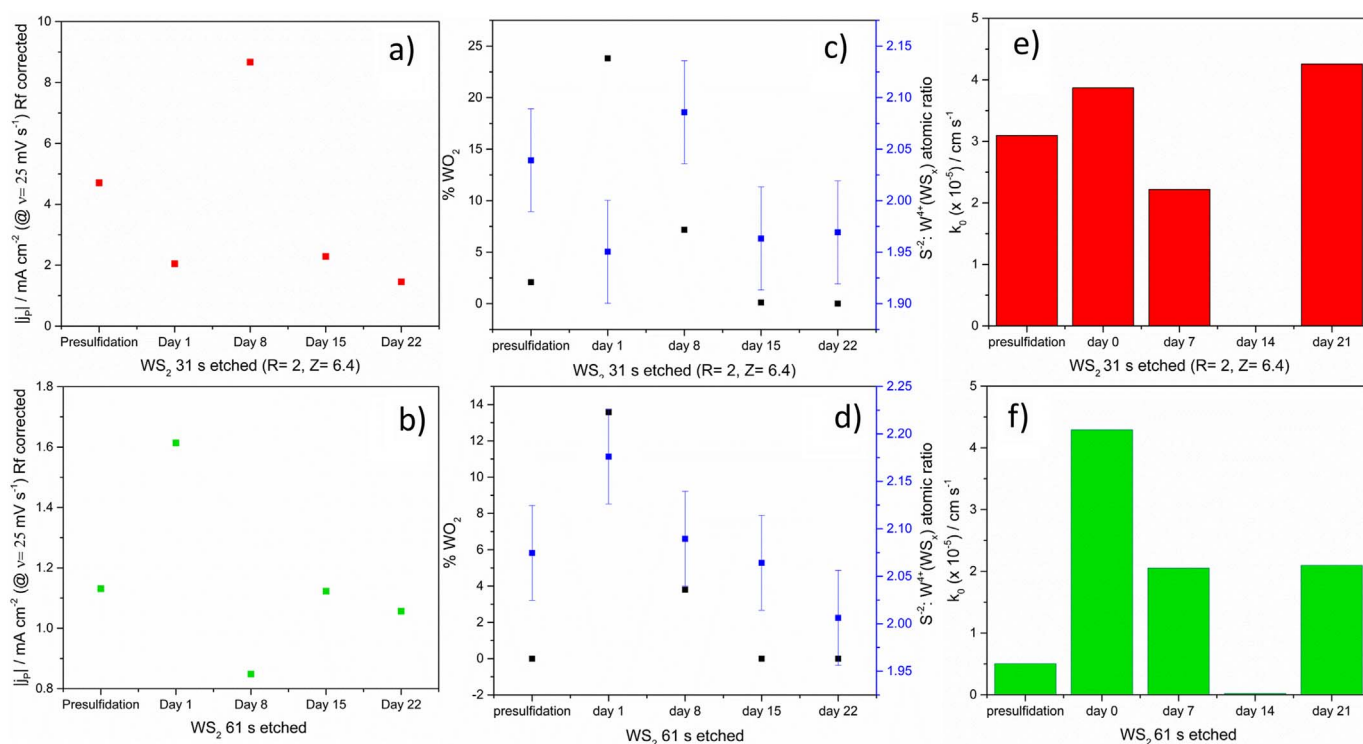


Fig. 1. Comparison of: a)–b) roughness-factor corrected HER peak current densities at $v = 25 \text{ mV s}^{-1}$, c)–d) WO₂ surface content (left axis) and S²⁻:W⁴⁺ XPS atomic photoemission ratios (right axis), and e)–f) k_{app}^0 values for the $[\text{Fe}(\text{CN})_6]^{4-}/[\text{Fe}(\text{CN})_6]^{3-}$ redox probe, for atmospherically aged, sulfidation treated plasma-etched WS₂ samples $31 \pm 1 \text{ s}$ ($R = 2$, $Z = 6.4$, first row) and $61 \pm 1 \text{ s}$ (second row), after weekly electrochemical testing over a three-week ambient exposure period.

an HER current decrease in subsequent electrochemical testing to values lower or comparable with the freshly sulfidated state, due to lower S:W ratios. This accords with previous investigations which correlated higher sulfur content in TMDs with improved hydrogen turnover frequencies [12,14], and sulfur-depleted W-edge sites of electro-oxidised WS₂ with poor catalytic activity [35]. We hypothesize that the electrochemically-induced restructuring gradually depletes the WO₂ phase, initially exposing underlying WS₂ with high active site densities which are later reconstructed during atmospheric and experimental conditions to a more homogeneous nanostructure (Fig. 2g–h).

The cathodic feature appearing at E ca. -0.4 V vs NHE in the HER experiments (Fig. 3a and b) is ascribed to the diffusion decay peak profile of proton reduction catalysed by the WS₂ active sites, characteristic of the fully-supported, low proton concentration electrolyte used [36,37]. Indeed, the resolution of this peak also seems correlated with the S²⁻:W⁴⁺ ratio, and consequently to the active sites present.

With regard to the electron transfer kinetics, both samples exhibit higher k_{app}^0 values ($\approx 4 \times 10^{-5} \text{ cm s}^{-1}$) after undergoing the sulfidation treatment (Fig. 1e–f). This agrees with literature reports which found enhanced electrical conductivities of WO_x species vs. WS₂ [38], beneficial for mediating in the redox chemistry of surface sensitive species such as $[\text{Fe}(\text{CN})_6]^{4-}/[\text{Fe}(\text{CN})_6]^{3-}$. A gradual decrease in k_{app}^0 values after two weeks of testing correlates with reduced WO₂ content at the crystal surface, whereas restored k_{app}^0 values after three weeks can be attributed to the higher WS₂ metallic 1T polymorph contents (ca. 10–15%).

These results suggest that this sulfidation method does not incorporate sulfur into the atmospherically aged WS₂ samples. Instead, it promotes the appearance of WO_x moieties at the WS_x surface which are reduced in the cathodic sweep. We hypothesize that in general, the sulfur incorporation is only effective when the electro-oxidative step of TMDs forms acid-soluble species, as sulfur incorporation into atmospherically-aged MoS₂ crystals was optimal when the cathodic voltage vertex surpassed the reduction potential of the TMD oxidised species (MoO_4^{2-}) [20]. In the case of WS₂, the oxidised WO_x species generated

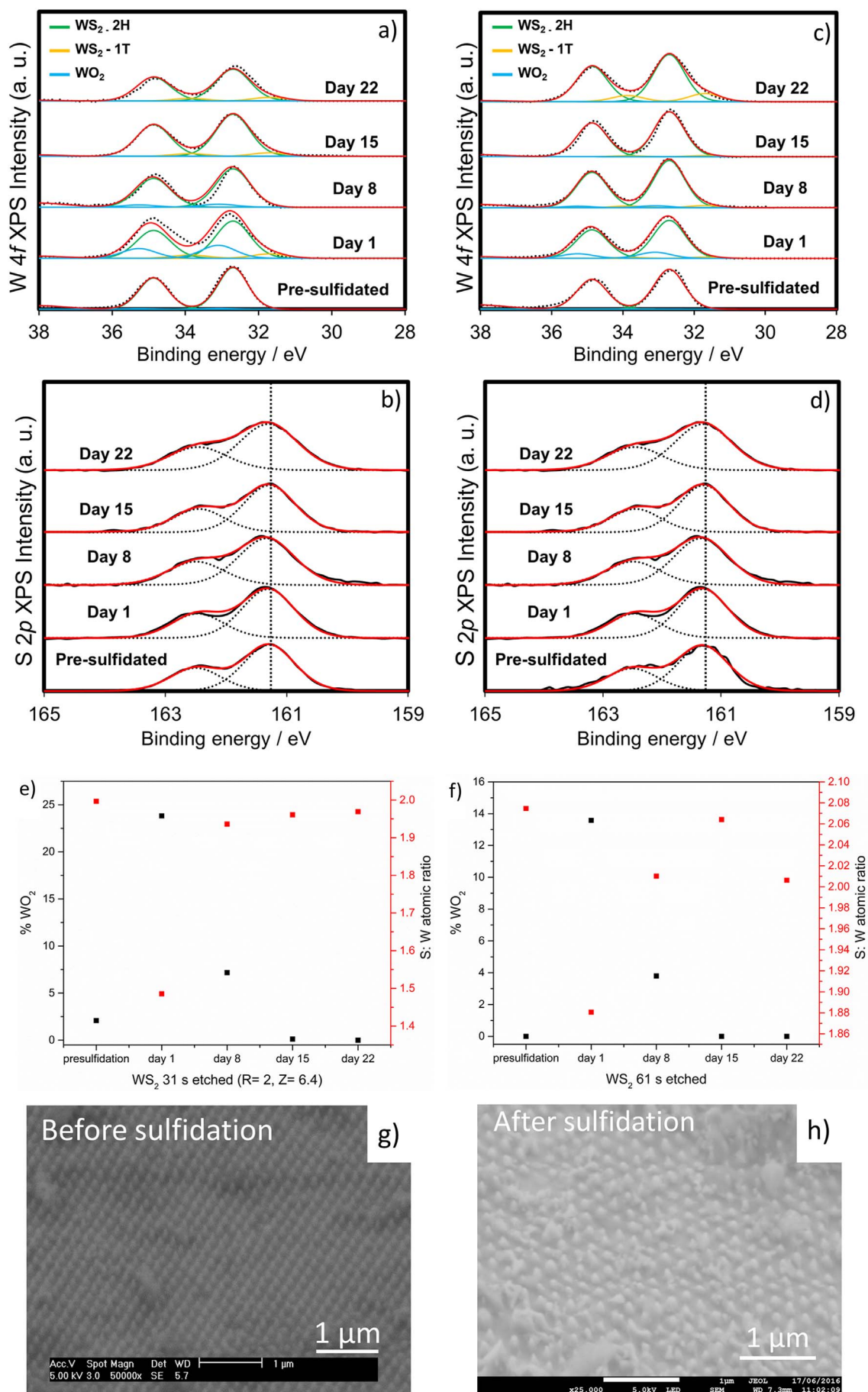
during the electro-oxidative step are insoluble at $\text{pH} \leq 3$ [39], coinciding with the optimized pH value for the sulfidation electrolyte (pH 3). Consequently, the electroreduced sulfur cannot be incorporated into the WO_x structure, and would dissolve under acidic conditions [40]. Hence, we predict that the electrochemical solvent-phase sulfidation method is only suited for MoX₂ ($X = \text{S}, \text{Se}$) rather than for WX₂ ($X = \text{S}, \text{Se}$).

4. Conclusions

In contrast to MoS₂, the application of the solution-phase, room-temperature electrochemical sulfidation method to obtain S-rich structures did not lead to S-rich WS_x but to S-deficient WS_x structures with high WO₂ surface content. The inferior HER performances but improved electron transfer properties are in agreement with the detrimental effect reported after WO₂ incorporation into WS₂ for the HER catalysis. The unsuccessful incorporation of electroreduced sulfide in the WS_x structure is suspected to arise from the nature of the sulfidation mechanism: redeposition of acid-soluble MoO_4^{2-} species for MoX₂ improves S²⁻ incorporation onto the surface, which is not possible in the case of WX₂ as the WO_x compounds formed are acid insoluble. This demonstrates the key role of the nature of the TMDs in the successful electrochemical incorporation of sulfur in their structure, and reveals that an electrochemistry-based sulfidation method universally applicable for any TMDs remains to be developed.

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(caption on next page)

Fig. 2. Stacked high-resolution XPS spectra of W 4f and S 2p for a)–b) 31 ± 1 s ($R = 2$, $Z = 6.4$) and c)–d) 61 ± 1 s atmospherically aged, sulfidation treated, plasma-etched WS₂ samples over a three-week ambient exposure period. e)–f) Comparison of total S:W XPS atomic photoemission ratios. Representative SEM micrographs g) before and h) after solution phase-sulfidation.

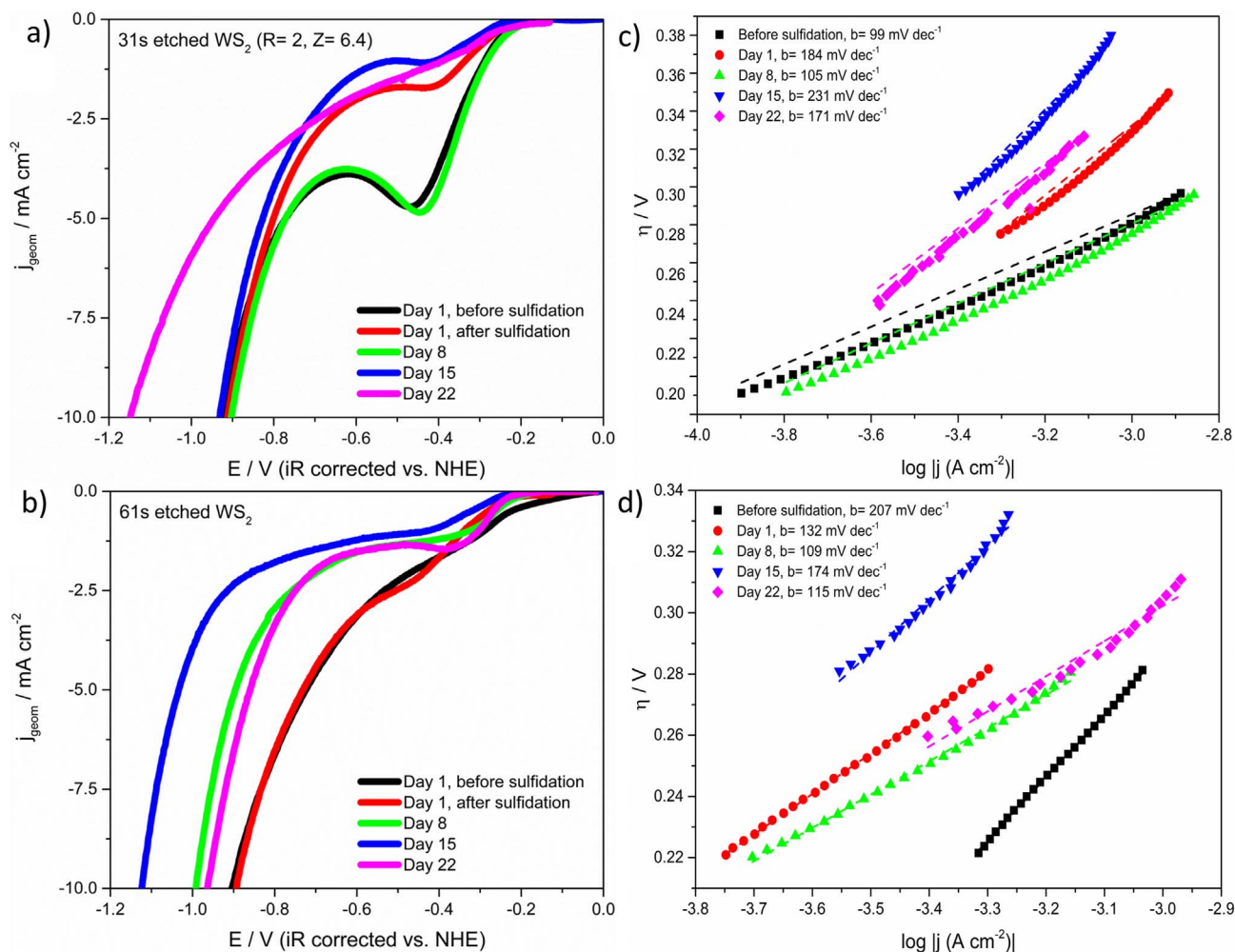


Fig. 3. Left column: Linear sweep voltammograms in the 0 to -1.2 V voltage range of a) 31 ± 1 s ($R = 2$, $Z = 6.4$) and b) 61 ± 1 s atmospherically aged, sulfidation treated plasma-etched WS₂ samples over a three week ambient exposure period. Right column: Tafel plots (η vs. $\log |j_{geom}|$) of c) 31 ± 1 s ($R = 2$, $Z = 6.4$) and d) 61 ± 1 s atmospherically aged, sulfidation treated plasma-etched WS₂ samples over a three week ambient exposure period. Labels: pre-sulfidated (black), post-sulfidated (red), 8-day atmosphere exposed (green), 15-day atmosphere exposed (blue) and 22-day atmosphere exposed (magenta). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

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