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Correlating Microstructure and Activity for Polysulfide Reduction and Oxidation at WS₂ Electrocatalysts

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The polysulfide reduction and oxidation activity of WS₂ electrocatalysts was studied. This was undertaken with a specific view to improve the efficiency of the polysulfide-bromine redox flow battery, for large scale energy storage. Using data from the literature, it is estimated that the catalysts would require exchange current densities of ~0.7 to 1 mA cm⁻² true microscopic surface area to enable them to become economically viable. The experiments were performed upon three different forms of WS₂: platelet-like particles of the 2H polymorph of WS₂ (2H-WS₂), ball milled WS₂ and inorganic fullerene-like WS₂ (IF-WS₂). The catalysts were characterized ex-situ using X-ray diffraction, scanning electron microscopy, transmission electron microscopy, energy dispersive X-ray analysis and N₂ gas adsorption methods. Electrochemical measurements were performed at 35 °C, in aqueous solutions of 1.8 M Na₂S₂.47 and 1 M NaOH, simulating the operating conditions of a half-charged polysulfide-bromine redox flow battery. The catalyst activity increased in the following order: IF-WS₂ < 2H-WS₂ < ball milled WS₂. We attempt to rationalize this trend on the basis of the morphological features observed ex-situ. The exchange current density of the ball milled WS₂, at ~0.012 mA cm⁻², falls short of the estimated target by a factor of ~60 to 70.

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There is an increasing need for energy storage on a large scale.1–4 This stems from a number of requirements: to improve power quality, improve economic efficiency and to enable more widespread use of renewable energy sources. At present, the most widely deployed means of storing energy on a large scale is hydroelectric power. However, the prerequisite for the construction of a new hydroelectric power station is the flooding of large areas of land. This is no longer permissible in Europe or North America due to social, political and environmental considerations. Consequently, new technologies are needed with the high efficiency and the low cost of hydroelectric power. The polysulfide bromine redox flow battery could represent a viable solution for the storage of energy on a large scale.5–9 Through the following reaction, electrical energy is stored in chemical bonds:

\[ xNa_2S_{x+1} + 3NaBr \rightarrow (x + 1)Na_2S_x + NaBr_3 \]  

 Upon charging, sodium tribromide is oxidized at the anode:

\[ 3NaBr \rightarrow NaBr_3 + 2Na^+ + 2e^- \]  

The sodium polysulfide species are reduced to lower polysulfide species at the cathode:

\[ xNa_2S_{x+1} + 2Na^+ + 2e^- \rightarrow (x + 1)Na_2S_x \]  

where (1 ≤ x ≤ 4). Alternatively, only taking into consideration the most abundant species in solution (the other species are in equilibrium with these species), the polysulfide reaction can be depicted as:10

\[ S_2^{2-} + 4H_2O + 6e^- \rightleftharpoons 4HS^- + 4OH^- \]  

The bromide and polysulfide reactions proceed in separate half cells, divided by an ion conducting membrane. Upon discharging the above reactions go into reverse.

The distinguishing feature of redox flow batteries in comparison to conventional, static batteries is that the electroactive species are stored in a separate tank.11,12 Conversely, in a static battery, these always remain within the actual cell. This means that the total energy stored in a redox flow battery system is independent of its power rating. The total energy capacity is a function of the size of the electrolyte storage tank. Consequently, in redox flow batteries, economies of scale come into place when large amounts of energy need to be stored.

One of the chief advantages of the S₂⁻-Br redox flow battery, relative to other redox flow batteries is that both S and Br are cheap and abundant. Polysulfide species, S₂⁻, have a high solubility in alkaline solutions,13 which gives the Na₂Sₓ electrolyte a favorable energy density in comparison to some other aqueous electrolytes. Nonetheless, the toxicity of the electrolyte in the Br half cell could be of some concern: when the Br electrolyte is in its charged state, the Br₋₂ anion will dissociate slightly to form Br⁻.14

In order to evaluate the viability of the polysulfide bromine redox flow battery for large scale energy storage, two recent studies have employed numerical models. The first of these, by Denholm and Kulcinski,16 estimates the “life cycle losses” of polysulfide bromine redox flow batteries, vanadium redox flow batteries, hydroelectric power stations and compressed air energy storage. Not only did they take into account the losses incurred directly by the electrical conversion efficiency, but also the indirect energy losses. These indirect losses were those incurred by construction, operations and maintenance, and in the case of compressed air energy storage, from the fuel and its delivery. According to their analysis, the overall life cycle efficiency is highest for hydroelectric power and vanadium redox flow batteries, at ~73%. In comparison, the polysulfide bromine redox flow batteries and compressed air energy storage were estimated to have an efficiency of ~65%. Although the energy conversion efficiency of compressed air systems is high, their use of fossil fuels markedly increases the effective energy input. In the case of the S₂⁻-Br system, the majority of the energy losses seem to be caused by the low electricity conversion efficiency. Nonetheless, an increase in the electrical conversion efficiency from 65% to 75% would provide it with an overall efficiency equal to that of hydroelectric power.16

The key to improving the efficiency of the polysulfide bromine redox flow battery can be found in a recent study by Scamman et al. They performed numerical simulations to evaluate the overall cost effectiveness of the polysulfide-bromide redox flow battery. In brief, Scamman et al.’s investigation predicted that Regenesys’ polysulfide-bromide redox flow battery would make a net economic loss. However, they also envisaged that the system would become profitable if an improvement was made to the kinetics of the system by increasing the rate constants of both half cells to 10⁻⁷ ms⁻¹. Improved kinetics would result in a lower volume of cells and electrolyte, decreased energy losses during charging and discharging and a higher power density. Since the rate constant for the polysulfide reaction is much lower than that of the bromide reaction, there is much more room for improvement in the kinetics of the polysulfide reaction. Arguably, the best way to achieve this would be to optimize the catalysis of the polysulfide half cell. Based on the rate constants stated above, this would require an exchange current density of ~0.2 A cm⁻² geometric surface area. In a redox flow battery, the electrodes are likely to
be three dimensional, and the catalysts would be dispersed on a high surface area, conducting support such as C-black. \(^9\) This signifies that the true catalyst surface area should be appreciably higher than the 2D geometric surface area. For example, in a state of the art polymer electrolyte membrane fuel cell, the true microscopic surface area of its Pt catalysts is 200 to 320 times greater than the 2D geometric surface area.\(^{20}\) Assuming that equivalent degrees of dispersion are achievable with the electrocatalysts of a S\(^2\)\(^-\)Br redox flow battery, the exchange current density required to achieve Scamman et al.’s targets would be \(< 0.7\) to 1 mA cm\(^{-2}\) true microscopic surface area.

There are numerous experimental studies which suggest that Scamman et al.’s proposed improvements to the kinetics of the polysulfide reduction and oxidation have been accelerated significantly through judicious use of electrode materials.\(^{7,21-27}\) However, very few, if any, studies have aimed to develop structure-activity relationships for electrocatalysts of polysulfide reduction and oxidation in particulate form.

Herein, we investigate the properties of several different forms of WS\(_2\) for polysulfide reduction and oxidation. Although bulk WS\(_2\) has not been studied previously for this reaction, several studies have reported that MoS\(_2\), which has a very similar chemistry to WS\(_2\), is a highly active catalyst for polysulfide reduction and oxidation.\(^{24,25,28}\) Allen and Hickling investigated a W electrode for polysulfide oxidation;\(^{29}\) even though its bulk composition was metallic, its surface composition was likely to be a WS\(_x\) phase (most metal surfaces have a tendency to form sulfide phases when exposed to sulfide containing electrolytes).\(^{21,27,30}\) Notably, the reduction of polysulfide occurs in nature via enzymes containingＷ;\(^{31}\) this could suggest that electrocatalysts based on W could also hold promise for man-made applications. Finally, the catalytic properties of layered transition metal sulfides have been studied extensively.\(^{32-36}\) Previous investigations of these catalysts have focused primarily on (non-electrochemical) catalytic applications in the petrochemical industry.\(^{32,33,37}\) and also more recently for the electrochemical or photoelectrochemical evolution of hydrogen.\(^{34-36,38-42}\) They have shown how it is possible to tailor the surface chemistry of these materials toward high catalytic activity. Consequently, capitalizing upon this knowledge base should enable the optimisation of layered transition metal sulfides for polysulfide reduction and oxidation.

MoS\(_2\) and WS\(_2\) take on an anisotropic layered structure, known as the “2H-polypeptide”, henceforth referred to as 2H-MoS\(_2\) or 2H-WS\(_2\).\(^{43}\) The hexagonal arrangements within each layer are similar to those of graphite. Also, similar to graphite, the layers are held together by weak Van der Waals forces, which allow the sulfides to be used as a solid state lubricant.\(^{44}\)

Fundamental investigations have afforded a deep understanding of the factors that govern the activity of these layered metal sulfide catalysts, enabling the improvement of their performance in large scale applications.\(^{32,34,35,37,38-48}\) Several studies have reported that the basal plane of these structures is inert, whereas the catalytically sites are located on the edge planes.\(^{46,49-51}\) This was clearly demonstrated by Jaramillo et al., who showed that the electrocatalytic activity of nanostructured MoS\(_2\) for hydrogen evolution was directly proportional to the length of its edge planes.\(^{52}\)

In this work, we aim to determine whether the activity of metal sulfides can also be tailored for polysulfide reduction and oxidation. We shall achieve this by studying the properties of WS\(_2\) in particulate form. The pronounced structure dependence of the catalytic properties of WS\(_2\) make it particularly amenable to such investigations. Moreover, changes to the structure can be characterized easily using ex-situ microstructural characterization techniques such as high resolution transmission electron microscopy (HR-TEM) and X-ray diffraction (XRD).

Our work compares two nanostructured variants of WS\(_2\) in comparison to micron-sized particles of 2H-WS\(_2\). In the first approach, edge planes and other defects are introduced into the 2H-WS\(_2\) through the process of ball milling. In the second approach, we investigate a form of WS\(_2\) which has very few edge planes: inorganic fullerenelike WS\(_2\) (IF-WS\(_2\)).\(^{52}\) Both these forms of metal sulfides have previously been investigated as gas phase heterogeneous catalysts, where they showed promising activity.\(^{7,35}\) Moreover, a recent investigation also showed that doped nanotubes of WS\(_2\) can efficiently photodecompose organics.\(^{46}\) However, to the best of our knowledge, they have not been investigated for their electrocatalytic properties.

### Experimental

**Microstructural characterization.—X-ray diffraction.**—X-ray diffraction (XRD) techniques were used for phase identification and the determination of crystallite size. A Phillips Vertical Diffractometer, model PW1050 was used, with the Bragg-Bretano geometry. This utilized Cu K\(_\alpha\) radiation (with wavelength, \(\lambda = 1.5418\) Å), a single bounce carbon monochromator, a 0.5 mm divergence slit, a 0.2 mm receiving slit, a 0.5 mm anti-scatter slit, a dwell time of 30 s and a step size of 0.02s. The powdered samples were placed on an oriented Si wafer for analysis.

From the peak width, it is possible to determine the crystallite size and microstrain in the crystal structure, although sometimes these two effects may be convoluted.\(^{49}\) The crystallite size was estimated from the peak width using the Debye-Scherrer equation:

\[
L_{hkl} = \frac{\lambda}{\beta_{hkl} \cos \theta_{hkl}}
\]

where \(L_{hkl}\) is the length of the crystallite perpendicular to the (hk1) crystallographic plane, \(\beta_{hkl}\) is the integral breadth of the peak, \(\theta_{hkl}\) is the peak position and \(\lambda\) is the wavelength of the incident radiation. In order to estimate microstrain (i.e. inhomogeneity in the lattice spacing) the following equation was used:

\[
\epsilon_{hkl} = \frac{\beta_{hkl}}{4\tan\theta_{hkl}}
\]

where \(\beta_{hkl}\) is the microstrain. The values of \(\beta_{hkl}\) were corrected for instrumental broadening by measuring a calibration sample of 10 wt% silicon powder and 90 wt% WS\(_2\) ball milled for 24 hours, using the method given by Langford et al.\(^{60}\)

**Surface area by gas adsorption.**—the surface area was determined using the Brunauer, Emmett and Teller (BET) method. This was achieved using a Micromeritics Tristar 3000 Instrument, using N\(_2\) as an adsorbate at 77.3 K. Each catalyst powder was weighed so that its total BET surface area would come to the recommended amount of 5 to 10 m\(^2\). Prior to the physical gas adsorption measurements, the samples were degassed overnight in a vacuum oven at 90 °C at 1 mbar total pressure. The decrease in mass as a result of the degassing procedure was always below 0.5%.

**Electron microscopy.—**the scanning electron microscopy (SEM) images were taken in a field emission gun SEM JEOL-6340F, in secondary electron imaging (SEI) mode at an acceleration voltage of 5.0 kV. A FEI Tecnai F20 microscope was used at an accelerating voltage of 200 kV. The catalyst powders were first dispersed in acetone for five minutes using an ultrasonic probe, and then dropped onto holey carbon Cu TEM grids. The TEM grids were supplied by Agar Scientific. Images taken in high resolution mode shall be designated as HR-TEM images, whereas lower resolution images are designated as TEM images.

**Catalysts.—**inorganic fullerenelike WS\(_2\) (IF-WS\(_2\)) was supplied by NanoMaterials Ltd, courtesy of Reshef Tennen. The synthesis of the IF-WS\(_2\) is described by Feldman et al.\(^{62}\) and involves the reaction of W\(_0\) nanoparticles with H\(_2\) and H\(_2\)S at elevated temperatures.
The 2H-WS₂ (99.8%) was supplied by Alfa Aesar. The ball milling experiments were carried out using a Spex 8000 vibratory mill. A custom made, hardened steel vial was used for the grinding. Prior to each experiment, the mill was cleaned in isopropyl alcohol using an ultrasonic probe. For each experiment, 7.30 g of material was placed inside the mill, along with \(2 \times \frac{1}{2} \text{inch}\) and \(4 \times \frac{1}{4} \text{inch}\) hardened steel balls. The powder occupied approximately 20% of the internal volume of the vial, as recommended by the mill’s manufacturer.\(^{63}\) The vial was closed using a screw top lid, and sealed using a Viton O-ring. Before the milling commenced, the mill was degassed and back filled with Ar (at a pressure of 2 bar) at least 6 times via a ball valve. A piece of filter paper was placed between the ball valve and the pipe leading to the main body of the vial in order to prevent the powder from being evacuated from the vial. After milling, the sample was left to cool, before being opened to air and then transferred to a vacuum desiccator.

**Electrochemical measurements.**—the water was purified using a Milipore water system, where the resistivity of the water was measured to be greater than 18 MΩ cm. All electrochemical glassware was washed and sonicated in this water, prior to usage.

Na₂S.9H₂O (ACS Reagent, >98%), powdered S (99.98%), NaOH (>97%), KCl (99.5% to 100.5%) and paraffin wax were all supplied by Sigma Aldrich. Isopropyl alcohol (>99.5%) and acetone (>99%) were supplied by Fisher Scientific. Acetylene black (99.9%) was supplied by Alfa Aesar.

The electrolyte was prepared as follows: the NaOH was dissolved in water in a flask placed on a magnetic heater/stirrer and deaerated for 30 minutes by bubbling it with Ar. An overpressure of Ar was maintained at all times. The Na₂S.9H₂O was then dissolved in the deaerated NaOH solution at 40°C. Once the Na₂S.9H₂O had completely dissolved, the S was added and stirred until it had also dissolved. The solution was made up to the required volume with de-aerated water in a volumetric flask.

The catalysts were immobilized on the electrode using the "sticky carbon" method\(^{64,65}\). This sticky carbon composite was made of a mixture of 35 wt% acetylene black and 65 wt% paraffin wax. The working electrode body was supplied by BioAnalytical Systems, model MF-2010. It was made from chlorotrifluoroethylene (CTFE), and incorporated an empty inset, 4 mm in diameter, 3 mm deep. A heat gun was then used to raise the temperature of the wax to around 50°C, so that the composite could be pressed further into place, albeit protruding out of the inset slightly. The surface of the electrode was then rubbed against a clean sheet of weighing paper, until it was smooth and planar.

A small amount of catalyst (≈0.2 mg) was then dispersed using an analytical sieve onto a glass optical microscopy slide. The electrode was weighed on an analytical balance, before being pressed lightly against the catalyst. The catalyst loaded electrode was then weighed to determine the amount of catalyst loaded on the electrode. The electrode was then transferred to the electrochemical cell.

All electrochemical measurements were controlled using a PG-STAT 30 from Autolab. All impedance spectroscopy measurements were taken from 1 MHz to 0.1 Hz, with a peak to peak amplitude of 10 mV. The impedance spectra were fitted using Autolab’s FRA software.

A two compartment, three electrode cell was used. A built in water jacket was used to maintain the cell at 35°C; this is the operating temperature of the polysulfide bromine redox flow battery.\(^{17,26}\)

All experiments were carried out under an inert, purified atmosphere of Ar, with a slight overpressure. The electrolyte was bubbled with Ar prior to the commencement of the experiments, and blanketed during the course of the experiment.

A Pt mesh of about 2 cm² in geometrical surface area, was used as a counter electrode. The reference was a 3.5 M KCl/calomel electrode. All potentiostatic measurements were cycled potentiodynamically for one hour at 10 mVs⁻¹. Each scan proceeded in the following order: \(\eta = 0 \text{ V} \rightarrow -0.27 \text{ V} + 0.22 \text{ V} \rightarrow 0 \text{ V}.\) (d) The potentiostatic steady state current measurement then commenced, proceeding in the following order: \(\eta = -0.275 \text{ V} \rightarrow +0.23 \text{ V},\) increasing in 10 mV intervals. The current was measured using a chronocoulometric technique, for 50 s at each potential (apart from the initial potential, where it was held for 100 s). The steady state current was evaluated after 50 s, immediately before the electrode was stepped to the next potential. (e) The potentiostatic measurement was also interspersed with impedance spectroscopy measurements at selected potentials.

**Results.**

**Microstructural characterization.**—The X-ray diffraction traces of the various WS₂ samples are shown on Figure 1. The traces have been plotted on a logarithmic scale; this scale was chosen simply because the (002) reflection is so much more intense than the other reflections that otherwise it would be difficult to view the smaller peaks, especially in the case of the 2H-WS₂. The predominance of the (002) reflection is a reflextion of the anisotropic nature of the 2H-WS₂ material. Clearly, all the traces correspond quite closely to the reference pattern for 2H-WS₂. With increased milling time, the peaks become significantly broader, which (apart from instrumental broadening) could be either attributed to a decrease in crystallite size or increase in microstrain (as described in the Experimental section). In principle, these effects can be deconvoluted using Williamson Hall plots.\(^{68,69}\) However, since the peaks on the XRD traces are broad and convoluted, it was not possible to converge to a meaningful fit using this type of analysis.

Nevertheless, Table I provides a rough approximation of the crystallite size and extent of microstrain, through analysis of the (002) peak, centered at around 14°. The integral value of the peak breadth, \(\beta_{(002)}\), was evaluated using a Lorentzian fit. Using equation 5 and

![Figure 1. X-ray diffraction traces of WS₂ based electrocatalyst powders. Starred peaks denote those that are not present in powder diffraction file 08–0237: \(\dagger\) denotes an unknown peak, \(\bigstar\) denotes the peak attributed by Zhu et al. to the (203) reflection of 2H-WS₂.\(^{14}\) The relative intensities are shown on a logarithmic scale, with each trace offset vertically from the other.](https://example.com/figure1.png)
Table I. Analysis of (002) peak broadening in WS$_2$ samples, from XRD results.

<table>
<thead>
<tr>
<th></th>
<th>2H-WS$_2$</th>
<th>WS$_2$ ball milled for 2 hours</th>
<th>WS$_2$ ball milled for 12 hours</th>
<th>WS$_2$ ball milled for 24 hours</th>
<th>IF-WS$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\beta_{(002)}$ (°)</td>
<td>0.1</td>
<td>0.2</td>
<td>1.3</td>
<td>2.2</td>
<td>0.7</td>
</tr>
<tr>
<td>Minimum crystallite size (nm)</td>
<td>79</td>
<td>38</td>
<td>7</td>
<td>4</td>
<td>14</td>
</tr>
<tr>
<td>Maximum microstrain</td>
<td>0.4%</td>
<td>0.8%</td>
<td>4.7%</td>
<td>7.9%</td>
<td>2.3%</td>
</tr>
</tbody>
</table>

assuming that the broadening of the (002) peaks is entirely due to decreased crystallite size, we estimate the lowest possible value of the average crystallite size. Conversely, by using equation 6 and assuming that the broadening is entirely attributable to microstrain, we estimate the maximum degree of microstrain. The data in Table I show that the milling brings about a large increase in microstrain or decrease in crystallite size, which develops further with milling time. There are some minor shifts in the peak positions between the different samples, suggesting a change in the size of the unit cell.

There is a significant unidentified peak on the WS$_2$ samples that were milled for 12 and 24 hours, centered at $\sim 24$°. Most probably this is an artifact of the milling process which resulted in the formation of a second phase. This could either be due to contamination from the balls or the vials, or otherwise the segregation of a separate phase within the WS$_2$. However, its intensity is still only $\sim 5\%$ of the (002) peak, and given that there are no other unidentified peaks of significant magnitude, the concentration of this second phase is probably low. There are some small additional peaks on the traces shown on Figure 1 (e.g. at $\sim 27$° on the unmilled 2H-WS$_2$ and the sample that was milled for 2 hours); however, bearing in mind the logarithmic scale, their intensity is negligible, and comparable to the background noise.

**Electron microscopy.**—FEG-SEM, TEM and HR-TEM images of the 2H-WS$_2$ particles are shown on Figures 2–4. Figure 5 shows FEG-SEM images of the WS$_2$ ball milled for 24 hours. Based upon these images, there is a marked difference between the milled and unmilled 2H-WS$_2$. Conversely, the different milled samples (not shown) are largely indistinguishable from each other. The milling process has resulted in a roughened microstructure, and the particles have lost their platelet-like morphology. It would be difficult to specify the dimensions of the particles, as they are of such an ill-defined shape and distribution that it is meaningless to distinguish an individual particle from the agglomerated mass to which it is attached. The agglomerates themselves are also fairly polydisperse, their size ranging from $\sim 100$ nm to $\sim 20 \mu$m.

The HR-TEM images shown on Figure 6 and Figure 7 provide further evidence of the effect of ball milling on the microstructure of WS$_2$. The HR-TEM images of the WS$_2$ ball milled for 2 hours are clearly distinguishable from those of the as received 2H-WS$_2$. The particles are no longer shaped like platelets, and the edges are particularly distorted. However, it is still possible to observe some ordered, uninterrupted lattice fringes of the (00$l$) plane that span tens of nanometres. This is similar in order of magnitude to the minimum...
average crystallite size which was estimated in Table I from the (002) peak in the XRD data. Notably, Kouzu et al.’s HR-TEM images of ball milled 2H-MoS2 resemble those shown here.55

On the basis of the HR-TEM images, the WS2 that was milled for 24 hours, shown on Figure 7, seems more distorted than the sample that was milled for 2 hours, shown on Figure 6. Although it is still possible to see the lattice fringes from the (00) reflection on the WS2 ball milled for 24 hours, they are much distorted and there are many dislocations present. Moreover, the extent of each ordered domain does not span more than about 4.5 nm, which is similar to the minimum crystallite size predicted from the XRD data, 4 nm (c.f. Table I).

The typical size and morphology of the IF-WS2 particles are depicted on the SEM and TEM images shown on Figures 8–10. These images are similar to those previously taken by Tenne and co-workers.52,53,71 The TEM images of the milled WS2 are qualitatively similar to those of recently reported “double-gyroid” MoS2, a catalyst engineered to have a high proportion of edge sites at its surface.74 In summary, it seems that the abundance of the edge planes and similar surface sites increases in the following order IF-WS2 < 2H-WS2 < milled WS2.

EDX analysis.— In order to determine the elemental composition of the WS2 samples, SEM-based EDX analyzies were performed. One of the primary motivations for doing this was to determine whether milling procedure had introduced metallic impurities into the material through its interaction with the balls and vial.

Typical spectra of the 2H-WS2, the WS2 milled for 24 hours, and the IF-WS2 are shown on Figure 11. The spectra of the 2H-WS2 and the IF-WS2 are close to identical. W and S are the most prominent peaks, as expected from the XRD data. There are some additional peaks due to Pt, C and O. The Pt is from the conductive coating sputtered onto the surface of the catalysts to prevent charging. The C is probably due to the backing tape which was used to support the particles. The O peak is so small that it can be neglected.

The WS2 that was milled for 24 hours exhibits the same peaks as the other two samples, with the exception of Pt. The S peak is slightly diminished whereas the O peak has increased in magnitude. Notably, no other metallic elements were detected that could have been introduced from the hardened steel vial and balls of the ball mill, such as Fe, Cr or Ni. However, it is possible that these elements are present in concentrations lower than ∼0.1%, the limiting sensitivity of EDX.75

Surface area by gas adsorption.— The BET surface area of all the catalysts is relatively low, ranging between 2.3 and 6.2 m² g⁻¹, as summarized in Table II. The ball milling effects an initial increase in the surface area, but then passes through a maximum before decreasing again. These values seem reasonable: a very rough estimation can be made of the expected surface area of the IF-WS2: assuming that the particles are 100 nm in diameter, completely spherical and have a
filled core with the theoretical density of 7.73 g cm\(^{-3}\), yields a value of \(\sim 8 \text{ m}^2 \text{ g}^{-1}\). Alternatively, by using the measured density of a pellet of IF-WS\(_2\), at 4.5 g cm\(^{-3}\) (which is presumably lower than the theoretical value as result of the hollow core)\(^7\)\(^6\) the surface area is \(\sim 13 \text{ m}^2 \text{ g}^{-1}\).

Electrochemical characterization.— All of the electrodes tested, including the carbon substrate, and the Pt counter electrode reached the same potential, \(-0.754 \text{ V}\) with respect to an SCE, within a few minutes of immersion within the cell. This is consistent with reports in the literature,\(^1\)\(^0\),\(^7\)\(^2\),\(^7\)\(^3\) and suggests that this potential is the reversible Nernst potential for polysulfide reduction and oxidation. Moreover, we assume, unless otherwise specified that all voltammetric and potentiostatic dc currents are faradaic in nature, and approximates to its steady state value. The hysteresis is slightly more pronounced on the IF-WS\(_2\), but only at low cathodic overpotentials. On all WS\(_2\) samples there is a limited degree of hysteresis at anodic overpotentials. The anodic response of the unmilled WS\(_2\) has been omitted from Figure 12 (as well as subsequent electrochemical data), as the catalyst loaded electrode was generally less active than the blank sticky carbon substrate.

The unmilled 2H-WS\(_2\) and the milled WS\(_2\) show very little hysteresis between scans taken in the positive and negative directions, under cathodic polarization. This suggests that the current is faradaic in nature and approximates to its steady state value. The hysteresis is attributable due to an adsorption pseudocapacitance, reconstruction of the surface, or the formation of elemental S.

The initial activity of the different samples toward polysulfide reduction and oxidation is shown on Figure 12. The inset shows the response of 0.16 mg unmilled 2H-WS\(_2\) in comparison to the blank substrate.

Table II. Summary of BET surface area and analysis for WS\(_2\) catalysts tested.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>BET surface area (m(^2)/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2H-WS(_2)</td>
<td>2.80 ± 0.01</td>
</tr>
<tr>
<td>WS(_2) milled 2 hours</td>
<td>6.12 ± 0.07</td>
</tr>
<tr>
<td>WS(_2) milled 12 hours</td>
<td>5.74 ± 0.03</td>
</tr>
<tr>
<td>WS(_2) milled 24 hours</td>
<td>2.35 ± 0.01</td>
</tr>
<tr>
<td>IF-WS(_2)</td>
<td>6.23 ± 0.08</td>
</tr>
</tbody>
</table>

Figure 11. Typical EDX spectra of 2H-WS\(_2\), WS\(_2\) ball milled for 24 hours and IF-WS\(_2\).

Figure 12. Initial potentiodynamic sweep of different WS\(_2\) samples. Normalized according to BET surface area. Taken in 1.8 M Na\(_2\)S\(_2\)\(_{4.7}\) and 1 M NaOH at 35°C. The scan proceeded in the following order: \(\eta = 0 \text{ V} \rightarrow -0.27 \text{ V} \rightarrow -0.22 \text{ V} \rightarrow 0 \text{ V}\), sweep rate: 10 mV \cdot s\(^{-1}\). The anodic response of the IF-WS\(_2\) has been omitted, as it was lower than that of the sticky carbon substrate. The inset shows the response of 0.16 mg unmilled 2H-WS\(_2\) in comparison to the blank substrate.

order: unmilled 2H-WS\(_2\) < WS\(_2\) milled for 2 hours < WS\(_2\) milled for 12 hours < WS\(_2\) milled for 24 hours.

It also appears that the IF-WS\(_2\) shows a lower activity than the 2H-WS\(_2\), especially at high cathodic overpotentials. The anodic response of the IF-WS\(_2\) has been omitted from Figure 12 (as well as subsequent electrochemical data), as the catalyst loaded electrode was generally less active than the blank sticky carbon substrate.

The unmilled 2H-WS\(_2\) and the milled WS\(_2\) show very little hysteresis between scans taken in the positive and negative directions, under cathodic polarization. This suggests that the current is faradaic in nature and approximates to its steady state value. The hysteresis is slightly more pronounced on the IF-WS\(_2\), but only at low cathodic overpotentials. On all WS\(_2\) samples there is a limited degree of hysteresis at anodic overpotentials. This hysteresis could possibly be attributed due to an adsorption pseudocapacitance, reconstruction of the surface, or the formation of elemental S.

Figure 13 shows that after cycling the electrodes for one hour, the same general trends in activity persist as those for the first sweep. However, there is a notable decline in activity over this period for the WS\(_2\) that was ball milled for 24 hours. The cathodic current decreases by 15 to 20%, whereas the anodic current drops by 40%.
to 50%. Conversely, the other samples retained their activity upon cycling, within a ~5% margin.

The data presented thus far suggest that the microstructural and electrochemical characteristics of the samples milled for 2 hours and 12 hours seem to be intermediate between those of the unmilled sample 2H-WS2 and the sample which was milled for 24 hours. Consequently, for the purpose of brevity, we shall focus on the unmilled sample and the sample that was milled for 24 hours, which shall henceforth be referred to as the milled WS2.

Following the potentiodynamic measurements, the steady state current was also measured potentiostatically. These data are shown on Figure 14. Neither the milled WS2 nor the IF-WS2 show a marked change in cathodic activity between the potentiodynamic and potentiostatic measurements. In contrast, the 2H-WS2 exhibited a three-fold increase in activity toward polysulfide reduction when measured potentiostatically.

On the other hand, under anodic polarization the potentiostatic activity of the milled WS2 was up to ~50% lower than the activity measured potentiostatically. In order to investigate whether this anodic deactivation of the ball milled sample was due to mass transport induced passivation, the experiment was also carried out in the presence of a magnetic stirrer. However, the activity of the unstirred sample was identical to that of the stirred sample. This suggests that the polarization of the milled WS2 is not influenced by mass transport.

Table III summarizes the Tafel parameters evaluated from the potentiostatic steady state polarization data shown in Figure 14, fitted to the exponential approximation of the Butler-Volmer equation:

\[ i_{SS} = i_0 10^\eta/b \]  

where \( i_{SS} \) is the steady state current, \( i_0 \) is the exchange current density, \( b \) is the Tafel slope, and \( \eta \) is the overpotential. The Tafel slopes alone can provide no additional information in this particular case, as the samples are all bimetallic catalysts. Consequently, we used impedance spectroscopy was used to shed further light upon the underlying electrochemistry of these materials.

**Impedance spectroscopy.** The same general features were present on the impedance spectra of all samples. However, for the purpose of brevity, we will focus the discussion here on the impedance spectra of the most active sample, the milled WS2.

Typical impedance spectra of the WS2 that was milled for 24 hours are shown on Figure 15 and 16, taken under anodic and cathodic polarization, respectively. There are two peaks: at high frequencies, between 10 and 100 kHz, and at low frequencies, between 10 and 100 Hz. Both relaxations are also clearly distinguishable from each other on the Nyquist plot, and are manifested as depressed, intersecting semicircles.

When fitting these data, the equivalent circuit shown on Figure 17 provided the best fit for all samples. Both relaxations in the impedance spectra can be modeled by parallel combinations of a resistance and a constant phase element, \( R_1 \) and CPE1 for the low frequency relaxation and \( R_2 \) and CPE2 for the high frequency relaxation. The fitting parameters for these spectra are summarized on Table IV.

With increasing cathodic overpotential, the low frequency intercept on the real axis, \( Z' \), decreases, as shown on Figure 15, due to the lowering of kinetic barriers as the system is drawn away from equilibrium. On the other hand, Figure 16 shows that under anodic...
polarization, the low frequency intercept on the real axis is virtually unchanged between the η = 0 V and η = +0.108 V.

This anomalous behavior can be traced back to the high frequency relaxation, due to R2 and CPE2. The depressed semicircle corresponding to this relaxation actually becomes more prominent as the electrode is polarized in the positive direction. It seems likely that the origin of this feature is a solid-state contact resistance. Evidence for this is provided by the capacitances, C2, for this feature in Table IV, estimated using the relation.81,82

\[ C = Q^{1/3} R^{(1/n-1)} \]  

Where Q and n are the fitting parameters of the CPE. The values are roughly the same for all samples, at ~20 to 30 nF cm\(^{-2}\); these values are far too low for an electrochemical double layer capacitance, but more likely to be related to a solid state contact.79,83 The notion that a solid state contact resistance could play such a significant role is unsurprising; the difficulties of making “Ohmic contacts” to layered semiconductor electrodes are well documented.83,84 A very similar feature was also observed by Conway and Liu upon their CoO2 electrocatalysts, due to the interface with a Ti substrate.85 The low frequency capacitance, C1, of the milled WS2 is around an order of magnitude higher than the equivalent values for the unmilled WS2 (not shown) This could suggest that the milled WS2 has an intrinsically higher electrochemically active surface area or a larger concentration of active sites.

Figure 18 compares the resistances R1 and R2 for the different WS2 catalysts. Clearly, the difference between R2 and R1 is much smaller for IF-WS2 and the milled WS2 than for the unmilled 2H-WS2. Indeed, in the case of the milled WS2, at η > + 150 mV, the contact resistance, R1, is greater than the charge transfer resistance R2. This suggests that at these high anodic potentials, an increase in overpotential, Δη1, is applied to a greater extent over the solid-solid interface than the electrochemical interface.

Using R1 and R2, we may estimate the relative contribution of the contact resistance, and the kinetic resistance, R1, to the overpotential. Since:

\[ \frac{dE}{d\eta} = Z_F (f \rightarrow 0) = R_1 + R_2 \]  

Figure 17. Equivalent circuit used to fit impedance spectra.

---

Table IV. Fitted parameters from impedance response of WS2 milled for 24 hours, in 1.8 M Na2S2,47 and 1 M NaOH, at 35°C. The surface areas were measured using the BET method.

<table>
<thead>
<tr>
<th>η (V)</th>
<th>R0 (Ω cm(^2))</th>
<th>R1 (Ω cm(^2))</th>
<th>10(^6)Q1 (F cm(^{-2}))</th>
<th>n1</th>
<th>10(^6)C1 (F cm(^{-2}))</th>
<th>10(^{-2})R2 (Ω cm(^2))</th>
<th>10(^5)Q2 (F cm(^{-2}))</th>
<th>n2</th>
<th>10(^5)C2 (F cm(^{-2}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>-0.256</td>
<td>49 ± 1</td>
<td>86 ± 2</td>
<td>151 ± 8</td>
<td>0.59 ± 0.01</td>
<td>7.5 ± 0.966 ± 0.02</td>
<td>164 ± 9</td>
<td>0.63 ± 0.01</td>
<td>0.9</td>
<td>10 ± 0.2</td>
</tr>
<tr>
<td>-0.211</td>
<td>65 ± 1</td>
<td>162 ± 4</td>
<td>106 ± 10</td>
<td>0.60 ± 0.02</td>
<td>7.4 ± 1.146 ± 0.04</td>
<td>79 ± 8</td>
<td>0.73 ± 0.02</td>
<td>2.6</td>
<td>20 ± 0.4</td>
</tr>
<tr>
<td>-0.168</td>
<td>68 ± 1</td>
<td>309 ± 8</td>
<td>78 ± 8</td>
<td>0.62 ± 0.02</td>
<td>8.3 ± 1.716 ± 0.06</td>
<td>72 ± 7</td>
<td>0.74 ± 0.02</td>
<td>2.8</td>
<td>30 ± 0.5</td>
</tr>
<tr>
<td>-0.142</td>
<td>66 ± 1</td>
<td>446 ± 8</td>
<td>74 ± 6</td>
<td>0.61 ± 0.02</td>
<td>8.6 ± 2.0 ± 0.1</td>
<td>59 ± 4</td>
<td>0.75 ± 0.01</td>
<td>3.0</td>
<td>40 ± 0.6</td>
</tr>
<tr>
<td>-0.104</td>
<td>68 ± 1</td>
<td>686 ± 15</td>
<td>64 ± 5</td>
<td>0.61 ± 0.02</td>
<td>8.7 ± 2.7 ± 0.1</td>
<td>49 ± 4</td>
<td>0.76 ± 0.01</td>
<td>3.1</td>
<td>50 ± 0.7</td>
</tr>
<tr>
<td>-0.076</td>
<td>76 ± 2</td>
<td>1000 ± 30</td>
<td>54 ± 6</td>
<td>0.62 ± 0.02</td>
<td>8.8 ± 3.5 ± 0.1</td>
<td>43 ± 5</td>
<td>0.77 ± 0.02</td>
<td>3.0</td>
<td>60 ± 0.8</td>
</tr>
<tr>
<td>-0.047</td>
<td>74 ± 2</td>
<td>1340 ± 40</td>
<td>49 ± 5</td>
<td>0.61 ± 0.02</td>
<td>8.5 ± 4.3 ± 0.2</td>
<td>35 ± 3</td>
<td>0.78 ± 0.02</td>
<td>3.1</td>
<td>70 ± 0.9</td>
</tr>
<tr>
<td>-0.009</td>
<td>65 ± 1</td>
<td>2030 ± 40</td>
<td>41 ± 3</td>
<td>0.63 ± 0.02</td>
<td>9.4 ± 5.3 ± 0.2</td>
<td>34 ± 2</td>
<td>0.79 ± 0.01</td>
<td>3.5</td>
<td>80 ± 1.0</td>
</tr>
<tr>
<td>0.000</td>
<td>65 ± 1</td>
<td>2070 ± 50</td>
<td>40 ± 3</td>
<td>0.63 ± 0.02</td>
<td>9.2 ± 5.4 ± 0.2</td>
<td>33 ± 2</td>
<td>0.79 ± 0.01</td>
<td>3.5</td>
<td>90 ± 1.1</td>
</tr>
<tr>
<td>0.040</td>
<td>67 ± 1</td>
<td>2180 ± 50</td>
<td>32 ± 3</td>
<td>0.62 ± 0.02</td>
<td>6.4 ± 7.2 ± 0.2</td>
<td>23 ± 2</td>
<td>0.82 ± 0.01</td>
<td>3.3</td>
<td>100 ± 1.4</td>
</tr>
<tr>
<td>0.050</td>
<td>68 ± 2</td>
<td>2140 ± 50</td>
<td>31 ± 3</td>
<td>0.62 ± 0.02</td>
<td>5.7 ± 7.7 ± 0.2</td>
<td>21 ± 2</td>
<td>0.83 ± 0.01</td>
<td>3.3</td>
<td>110 ± 1.6</td>
</tr>
<tr>
<td>0.089</td>
<td>69 ± 1</td>
<td>1910 ± 50</td>
<td>28 ± 2</td>
<td>0.60 ± 0.02</td>
<td>4.0 ± 9.0 ± 0.3</td>
<td>15 ± 1</td>
<td>0.85 ± 0.01</td>
<td>3.1</td>
<td>140 ± 1.7</td>
</tr>
<tr>
<td>0.108</td>
<td>70 ± 1</td>
<td>1710 ± 60</td>
<td>28 ± 3</td>
<td>0.59 ± 0.02</td>
<td>3.6 ± 9.8 ± 0.4</td>
<td>12 ± 1</td>
<td>0.87 ± 0.01</td>
<td>3.0</td>
<td>150 ± 1.9</td>
</tr>
<tr>
<td>0.118</td>
<td>72 ± 1</td>
<td>1650 ± 60</td>
<td>28 ± 3</td>
<td>0.59 ± 0.03</td>
<td>3.3 ± 10.6 ± 0.4</td>
<td>11 ± 1</td>
<td>0.87 ± 0.01</td>
<td>2.9</td>
<td>160 ± 2.1</td>
</tr>
<tr>
<td>0.147</td>
<td>73 ± 1</td>
<td>1280 ± 60</td>
<td>36 ± 4</td>
<td>0.56 ± 0.03</td>
<td>3.3 ± 10.9 ± 0.4</td>
<td>9 ± 1</td>
<td>0.89 ± 0.01</td>
<td>2.8</td>
<td>170 ± 2.3</td>
</tr>
<tr>
<td>0.186</td>
<td>76 ± 2</td>
<td>780 ± 50</td>
<td>68 ± 8</td>
<td>0.52 ± 0.03</td>
<td>4.4 ± 10.6 ± 0.4</td>
<td>6 ± 1</td>
<td>0.91 ± 0.01</td>
<td>2.4</td>
<td>180 ± 2.5</td>
</tr>
<tr>
<td>0.216</td>
<td>75 ± 2</td>
<td>490 ± 30</td>
<td>150 ± 20</td>
<td>0.45 ± 0.03</td>
<td>6.5 ± 9.2 ± 0.3</td>
<td>5 ± 1</td>
<td>0.93 ± 0.01</td>
<td>2.2</td>
<td>190 ± 2.7</td>
</tr>
</tbody>
</table>
assuming linear behavior, for a small change in steady state current density, $\Delta i_s$, there is a corresponding change in potential across the interface:

$$\Delta \eta = \Delta i_s (R_1 + R_2) = \Delta \eta_1 + \Delta \eta_2$$  \[10\]

where $\Delta \eta_1$ and $\Delta \eta_2$ are the contributions to the change in potential from $R_1$ and $R_2$, respectively, when the current is increased by an amount $i_s$. At each point where an impedance spectrum is taken, there will be a steady state DC current density, $i_s$. By summing up the $\Delta \eta$ for each incremental increase in $i_s$, it is then possible to estimate $\eta_1$ and $\eta_2$, the contributions to the overpotential from the $R_1$ and $R_2$, as a function of $i_s$. A similar approach was taken by Wagner to estimate the individual contributions to the total overpotential of a polymer electrolyte membrane fuel cell.86

The results of this analysis are shown on Figure 19a, 19b and 19c for the unmilled WS$_2$, the milled WS$_2$ and the IF-WS$_2$, respectively. For the unmilled sample only a small part of the overpotential seems to arise from the contact resistance. However, for the IF-WS$_2$ and the milled WS$_2$ it is much more significant, especially at high overpotentials; up to $\sim 30\%$ of the overpotential is due to the contact resistance on the milled WS$_2$, and $\sim 25\%$ for the IF-WS$_2$.

The insets on Figure 19a, 19b and 19c compare the polarization curves simulated from the impedance data, and the actual polarization curves from the potentiostatic data shown in Figure 14. In general, the simulated curves correspond closely to the actual polarization curves, validating this approach whereby impedance spectroscopy is used to describe the polarization of the WS$_2$ catalysts. Using the data shown from Figure 19, it is now possible to estimate the hypothetical activity of the different catalysts in the absence of the contact resistance, in which case the current should be controlled by kinetics. The simulated kinetic polarization plots of the catalysts are shown on Figure 20. In the case of the milled WS$_2$, the plots are linear in both the anodic and cathodic regions, and show no inflection, similar to those of the 2H-WS$_2$. The simulated curve of the IF-WS$_2$ is not as linear, but nevertheless a single Tafel slope can be fitted over the entire cathodic range more satisfactorily than in the polarization plots shown in Figure 17.

The Tafel parameters from these plots are summarized in Table V. The simulated exchange current densities, and some of the Tafel slopes are very similar in magnitude to the low overpotential values from Table III. The simulated Tafel slopes all range between 100 and 135 mV decade$^{-1}$. These values are within the typical range for a kinetically controlled current,80 and much lower than the apparent Tafel slopes for these materials at high overpotentials on Table III.

Finally, it is also of interest to compare the kinetic activity of the different catalysts, using the data presented in Figure 20. The milled WS$_2$ is 30 to 40 times more active than the unmilled 2H-WS$_2$. Conversely, the 2H-WS$_2$ shows a 4 to 10 fold increase in activity over the IF-WS$_2$. 

![Figure 18. Comparison of resistances $R_1$ and $R_2$ for different WS$_2$ samples, determined from fits of impedance data. Taken at 35°C in 1.8 M Na$_2$S$_2$O$_7$ and 1 M NaOH. Filled squares represent the charge transfer resistance, $R_1$; hollow diamonds represent the contact resistance, $R_2$ with lines connecting data points to aid visualization.](image)

![Figure 19. Comparison of $\eta_1$ and $\eta_2$, for (a) unmilled WS$_2$, (b) the WS$_2$ milled for 24 hours, and (b) IF-WS$_2$, determined from impedance data. Taken at 35°C in 1.8 M Na$_2$S$_2$O$_7$ and 1 M NaOH. Shaded gray area represents kinetic contribution to overpotential, $\eta_1$; shaded red area represents contribution from contact resistance, $\eta_2$; black squares show simulated total overpotential, $\eta_2$, with lines connecting data points to ease visualization. The inset compares the polarization curve simulated from the impedance data ( ), to the actual polarization curve measured from the potentiostatic data ( ).](image)
Discussion

Structure dependence of electrocatalytic activity.— Evidently, the materials under study exhibit a wide range of electrocatalytic activity for polysulfide reduction and oxidation, spanning over two orders of magnitude. These can be explained by considering the differences in structure between each material.

The microstructural characterization showed that each catalyst has roughly the same bulk composition and crystal structure. However, distinct morphological differences were observed between each catalyst, especially with respect to the predominance of edge planes on their surfaces. The proportion of edge planes increases in the same magnitude. These can be explained by considering the differences in activity for polysulfide reduction and oxidation, spanning over two orders of magnitude. These can be explained by considering the differences in structure between each material.

Nonetheless, more complicated explanations could also account for the trends we observed. We acknowledge that the sites that dominate the catalytic activity may not be equivalent on each material.

Reactivity kinetic—It is interesting to note the similarities between the electrochemical response of all the different samples. The same equivalent circuit could be used to model each material’s impedance spectra. In particular, there only seems to be one relaxation between the electrochemical response of all the different samples.

Solid state contact resistance.—Impedance spectroscopy elucidated the influence of the contact resistance on the polarization response of the catalysts tested. Non-metallic, poorly conducting electrocatalysts have received increasing attention in recent years for energy conversion processes, in particular those based on sulfides.34–36,97

Mass transport and anodic passivation.—Several authors have observed the ‘passivation’ of their electrodes during polysulfide oxidation due to the formation of a S film.7–9,29,107–109 Zhou et al. propose that efficient mass transport can prevent this anodic passivation.9 Presumably, mass transport alleviates the passivation as a result of the autocatalytic effect that polysulfides seem to exhibit upon the dissolution of the S layer.29,107–109

On the catalysts tested here, there were no obvious signs that the current was controlled by mass transport, as the electrochemical response was unaffected by the absence of a stirrer. Indeed, a conventional mass transport limitation should manifest itself as a Warburg feature in the impedance spectra.110 It is possible that mass transport effects could be observable in more dilute solutions, or at higher current densities or potentials.

Table V. Tafel parameters for different WS2 samples, using simulated kinetic polarization plots from data presented in Figure 20. The errors shown are the standard deviation in each fit.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>η (V)</th>
<th>10^6b (V decade⁻¹)</th>
<th>10^6i₀ (A mg⁻¹)</th>
<th>10^6i₀ (A cm⁻² BET area)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cathodic region</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unmilled 2H-WS2</td>
<td>−0.253 to −0.059</td>
<td>103 ± 2</td>
<td>7.7 ± 0.4</td>
<td>28 ± 2</td>
</tr>
<tr>
<td>WS2 milled for 24 hours</td>
<td>−0.176 to −0.061</td>
<td>112 ± 1</td>
<td>283 ± 8</td>
<td>1200 ± 30</td>
</tr>
<tr>
<td>IF-WS2</td>
<td>−0.214 to −0.060</td>
<td>125 ± 10</td>
<td>5 ± 1</td>
<td>7 ± 2</td>
</tr>
<tr>
<td>Anodic region</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2H-WS2</td>
<td>+ 0.044 to + 0.080</td>
<td>104 ± 4</td>
<td>5.8 ± 0.3</td>
<td>21 ± 1</td>
</tr>
<tr>
<td>WS2 milled for 24 hours</td>
<td>+ 0.052 to + 0.181</td>
<td>135 ± 4</td>
<td>240 ± 10</td>
<td>1020 ± 50</td>
</tr>
</tbody>
</table>

Figure 20. Simulated kinetic polarization response of different WS2, based on i−η data, determined from impedance spectroscopy measurements. Taken at 35°C in 1.8 M Na₂S₂(47) and 1 M NaOH. The straight lines show the fitted Tafel slopes.
Although we find no evidence to suggest that there are mass transport restrictions under our experimental conditions, we speculate that some other kind of passivation, unrelated to mass transport, could be the cause of the steady decrease in anodic activity of the milled WS$_2$, with cycling. This effect can be observed by comparing Figure 12 and Figure 13. The lower anodic activity of the potentiostatic measurement, relative to the potentiodynamic measurement, on Figure 13 and Figure 14, respectively, could also be symptomatic of static measurement, relative to the potentiodynamic measurement, on the catalyst. This was beneficial; it would be difficult to make a fair comparison between different studies. However, previous studies did not attempt to determine the inherent activity of particulate electrocatalysts for polysulfide reduction and oxidation, normalized according to their ‘true’ surface area. This makes it difficult to make a fair comparison between different studies. Moreover, a large number of nanostructured sulfide catalysts have been tested for use in the petrochemical industry, whereas very few of these have been tested for polysulfide reduction and oxidation. Fruitful strategies could be to study the activity of WS$_2$ and other metal sulfides by incorporating more than one metallic element into the catalysts.

Each of the catalysts tested here had a uniformly low surface area, ranging from $\sim 2$ to $\sim 6 \text{ m}^2\text{g}^{-1}$. For their experimental characterization, this was beneficial; it would be difficult to make a fair comparison of catalyst activity if they exhibited widely different surface areas. Conversely, for their industrial application, a much higher surface area would be necessary in order to maximize the catalyst utilization. For instance, C-supported Pt electrocatalysts in a polymer electrolyte membrane (PEM) fuel cell have typical surface areas of $\sim 70$ to $\sim 80 \text{ m}^2\text{g}^{-1}$. Notably, the investigations of Kouzu et al. suggest that such high surface areas are also obtainable using the type of sulfide catalysts investigated here. Their ball milled MoS$_2$ had a surface area of $\sim 80 \text{ m}^2\text{g}^{-1}$. The primary difference between their synthesis method and that which was used here is their addition of butanol to the mill, which presumably acted as a stabilizing agent. Improved dispersion should also be achievable using a high surface area C support such as those used in PEM fuel cells. A viable one-step process to produce a supported catalyst could be to co-grind the WS$_2$ with a high surface area C support (possibly with a stabilizing agent such as butanol).

The solid state contact resistance observed here would result in significant efficiency losses in a S$_2^2$-Br redox flow battery, especially at high current densities. Methods of avoiding such poor contacts are usually found empirically, either by doping the semiconducting electrode or changing the support. Indeed, it is conceivable that this problem is a characteristic of the sticky carbon composite, and would not manifest itself on a support with a different composition.

Conclusions

In this study, a combination of ex-situ characterization methods and electrochemical experiments were used to study the factors controlling the activity of particulate WS$_2$ electrocatalysts. We found that the catalyst activity increased in the following order: IF-WS$_2$ < unmilled 2H-WS$_2$ < milled WS$_2$, in correlation with the number of edge planes at the catalyst surface. This trend is consistent with other investigations of similar metal sulfides but for different reactions. This suggests that the factors governing their activity toward polysulfide reduction and oxidation are similar to those for other catalytic systems. We conclude our work by proposing several promising avenues toward the optimal electrocatalysis of polysulfide reduction and oxidation.

Acknowledgments

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References
