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Nicolas Goubard-bretesché, Olivier Crosnier, Camille Douard, Antonella Iadecola, Richard Retoux, et al.. Unveiling Pseudocapacitive Charge Storage Behavior in FeWO₄ Electrode Material by Operando X-Ray Absorption Spectroscopy. *Small*, 2020, 13, pp.2002855. 10.1002/sml.202002855 . hal-02902341

HAL Id: hal-02902341

<https://hal.science/hal-02902341>

Submitted on 6 Nov 2020

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Unveiling Pseudocapacitive Charge Storage Behavior in FeWO₄ Electrode Material by Operando X-ray Absorption Spectroscopy

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Keywords: FeWO₄, Operando, X-ray absorption spectroscopy, electrochemical capacitors, pseudocapacitance

In nano-sized FeWO₄ electrode material, both Fe and W metal cations are suspected to be involved in the fast and reversible Faradaic surface reactions giving rise to its pseudocapacitive signature. As for any other pseudocapacitive materials, to fully understand the charge storage mechanism, a deeper insight into the involvement of the electroactive cations still has to be provided. The present paper illustrates how operando X-ray absorption spectroscopy (XAS) has been successfully used to collect data of unprecedented quality allowing to elucidate the complex electrochemical behavior of this multicationic pseudocapacitive material. Moreover, these in-depth experiments were obtained in real time upon cycling the electrode, which allowed investigating the reactions occurring in the material within a realistic timescale, which is compatible with electrochemical capacitors practical operation. Both Fe K-edge and W L₃-edge measurements point out the involvement of the Fe³⁺/Fe²⁺ redox couple in the charge storage while W⁶⁺ acts as a spectator cation. The result of this study enables to unambiguously discriminate between the Faradaic and capacitive behavior of FeWO₄. Beside these valuable insights toward the full description of the charge storage mechanism in FeWO₄, this paper demonstrates the potential of operando X-ray absorption spectroscopy to enable a better material engineering for new high capacitance pseudocapacitive electrode materials.

1. Introduction

Electrochemical capacitors (ECs) have been introduced in the wide family of electrochemical energy storage devices 60 years ago, since the patent granted to Becker in 1957. ^[1] Unlike their battery counterpart, they can store or deliver energy within short periods of time, typically ranging from few milliseconds up to hundreds of seconds. Subsequently, they complement batteries when a short time scale is required for a specific use such as regenerative braking in hybrid or electric cars, fast charging electric buses or tramways,

powering the emergency opening of airplane doors, or supplying electricity in uninterrupted power supply systems.^[2] Almost all commercial ECs are manufactured using activated carbon as the active electrode material. The charge storage in such a device occurs through an accumulation of ions at the interface of both electrodes with the electrolyte.^[3] The intriguing dynamics of ions in the carbon meso or microporosity as well as at the carbon surface have been widely investigated by the use of *in situ* or *operando* techniques such as infrared spectroscopy,^[4] electrochemical quartz crystal microbalance,^[5] or X-ray absorption spectroscopy (XAS),^[6] among others.

On the other hand, pseudocapacitive materials, *i.e.* compounds whose peculiar charge storage mechanism involves surface and/or subsurface fast and reversible redox reactions, are of great interest in the field, as they allow reaching greater capacitance values than carbon materials, especially from a volumetric point of view.^[7] However, the origin of the electrochemical behavior of pseudocapacitive compounds^[8] is still under debate since only few experiments have confirmed its Faradaic origin. Even fewer studies aimed at investigating such a behavior while the cell is operated at a realistic cycling rate, *i.e.* within few seconds or few tens of seconds. While a better knowledge of the electrochemical phenomena occurring at/in pseudocapacitive electrodes will provide fundamental insights, it should also help to design new compounds for high capacitance devices.

Nowadays, tremendous efforts are being made to improve the energy density of ECs based on pseudocapacitive materials without compromising either their power density or cycling ability. As such, new materials are regularly emerging and, most recently, multicationic oxides were proposed as high volumetric capacitance electrode materials for ECs operated in aqueous electrolytes.^[9] The combination of an electroactive element with high density species can lead to attractive volumetric capacitance values without being detrimental to both cycling rate and stability. Iron tungstate (FeWO_4) has demonstrated a capacitive-like

electrochemical signature over long-term cycling operation,^[10] but the involvement of W and Fe cations in the charge/discharge mechanism has not been established yet.

As a matter of fact, early *in situ* ruthenium K-edge XAS was used to evidence the change in the ruthenium oxidation state upon polarizing a RuO₂ electrode at various potentials.^[11] The same technique shed light on the role of Mn cations in the charge storage mechanism in MnO₂ thin films,^[12] involving a continuous and reversible change in Mn oxidation state and supporting the results previously obtained by *ex situ* XPS analyses on these materials.^[13] More recently, similar conclusions were obtained with *in situ* XAS experiments at the titanium K-edge in the pseudocapacitive behavior of Ti₃C₂ MXene material in sulfuric acid.^[14] In all these materials, the role of structural cations was pointed out. However, there are still ambiguous concerns to be solved, namely (i) the evidence of changes in the cation oxidation state while operating the cell in a realistic timeframe and (ii) the role of each cation when multicationic oxides are considered such as in FeWO₄. Indeed, despite the importance of such findings obtained by *in situ* XAS, each data set was recorded after at least 15-20 minutes polarization of the electrode at a given potential. Unfortunately, the *in situ* approach does not allow to depict the dynamics of an electrode upon cycling and its (electro)chemical response to a realistic electrochemical solicitation. To the best of our knowledge, *operando* Raman spectroscopy^[15] is currently the only analytical technique that has been used on a pseudocapacitive material (MnO₂) and that reasonably fits with EC electrode cycling rates, but without giving access to the change in manganese oxidation state, unfortunately.

For the above-mentioned reasons, quick-XAS experiments performed under *operando* conditions (*i.e.* when the cell is in a realistic operation mode) is the best approach to investigate the fast processes occurring in pseudocapacitive EC electrode materials. To get a chance to evidence the studied phenomenon, it seems reasonable to assess the data collection time to be tens to hundreds times shorter than the charging/discharging time.^[16] As such, for

operando XAS measurements on battery electrodes, data acquisition times of a few minutes per spectrum for a charge or discharge at a C/10 rate (*i.e.* 1 mole of ion exchanged in 10 hours) have been used.^[17] When translated to the fast cycling rate of EC electrodes, the time scale is reduced to only a few seconds per spectrum and compatible only with the fast data acquisition capabilities achieved with the quick-XAS setup.

In the present work, we fully describe the electrochemical charge storage mechanism in FeWO₄, thanks to the analysis of high quality XAS data gained using *operando* quick-XAS measurements at both Fe and W K and L₃ -edges, respectively. Beside the elucidation of the respective roles of Fe and W metal cation centers in its electrochemical behavior, this study sets FeWO₄ as an example of how the contribution of cations can be unveiled during the cycling of the electrode by using this advanced spectroscopic *operando* technique.

2. Results and discussion

Nanosized FeWO₄ was synthesized using a microwave-assisted coprecipitation route with water as the solvent. A poorly crystallized material is obtained, as it can be seen from the XRD diagram (**Figure 1A**) and from the TEM observations (**Figure 1B** and **1C**) with 10-15 nm particle size and a specific surface area of 73 m².g⁻¹ (**Table S1**). After mixing the FeWO₄ powder with conductive carbon black and PTFE binder (60:30:10 weight ratios), thick self-supported electrodes (FeWO₄ mass loading = 10.0±0.3 mg.cm⁻²) were manufactured. Preliminary electrochemical investigations have demonstrated the capacitive-like signature of such an electrode (**Figure S1A**) as well as an impressive cycling ability (**Figure S1B**), thus confirming the choice of FeWO₄ nanopowder-based electrode as a typical potential pseudocapacitive compound.

Before running *operando* XAS measurements, these electrodes were placed in a specially designed *in situ* cell (**Figure S2**) and soaked in the electrolyte for several minutes at open-

circuit potential (OCP) (reference electrode Ag/AgCl (3M NaCl); counter electrode: platinum wire). Under such *in situ* conditions, X-ray absorption spectra were acquired at both Fe K-edge and W L₃-edge to determine the valence of iron and tungsten, respectively, in the pristine material at the OCP. The corresponding spectra are presented in Supporting Information (**Figure S3A** and **S4**). Since the oxidation state of an absorbing element varies linearly with the energy of its absorption edge, it is therefore possible to monitor the evolution of the oxidation state during cycling by following the edge energy shift. For a higher oxidation state of the studied element, a corresponding edge shift at higher energy values is expected, and *vice versa* for the lower oxidation state. At OCP, the edge position of the Fe K-edge X-ray absorption near edge structure (XANES) spectrum of as-prepared FeWO₄ overlaps with that of hematite (Fe³⁺), which is consistent with the Fe oxidation state determined by Mössbauer spectroscopy (Fe³⁺, see **Figure S3B**). One can see that Fe₂O₃ spectrum displays a shoulder (between ca. 7122 and 7125 eV) after which both FeWO₄ and Fe₂O₃ spectra are superimposed. This feature is common for iron-containing materials and can be attributed to spurious electronic effect related to the different local geometries (typically 1s → 4s transitions).^[18] Moreover, the presence of the pre-peak features at about 7114 eV in the pristine sample confirms the distorted FeO₆ octahedral environment typical of Fe³⁺ oxidation state. To evaluate the W oxidation state at OCP, a WO₃ powder was used as a reference, and the maximum of the derivative of its W L₃-edge overlaps with that of the pristine sample, confirming that tungsten is at a +6 oxidation state. This observation is consistent with previous XAS and XPS studies performed on nanosized iron tungstates,^[19] and suggests that cationic (Fe or W) vacancies should occur in the material to compensate the anionic charge. First-principles DFT+U calculations performed on the stoichiometric FeWO₄ phase clearly point iron rather than tungsten as the deficient metal in this structure (see **Figure S5**). The Density of States computed for FeWO₄ displays a very narrow band at the Fermi level which

corresponds to highly localized Fe(d)-states and suggests that polarons can easily form in this system. This result is fully consistent with a thorough study recently reported on the electronic structure of FeWO₄ in which the pseudocapacitive behavior of FeWO₄ was predicted to be linked to the formation of hole polarons at the Fe-sites together with dominant native point defects on the iron sites. [20] From now on, the non-stoichiometric Fe_{1-x}WO₄ will then be written as $\square_y Fe_{(1-3y)}^{II} Fe_{2y}^{III} WO_4$, \square_y being the content of Fe-vacancies in the material. A simple electron count shows that y varies from 0 to 1/3 when going from the pure Fe^{II} phase (Fe^{II}WO₄) to the pure Fe^{III} phase $\square_{1/3} Fe_{2/3}^{III} WO_4$, all intermediate corresponding to mixed Fe^{II}/Fe^{III} valence. Combining these results with Mössbauer and XANES data (100% Fe³⁺ in our sample), the iron tungstate material presented herein would display the formula $\square_{1/3} Fe_{2/3}^{III} WO_4$. To experimentally confirm this theoretical statement, an elemental analysis of the material was carried out using ICP-OES spectroscopy (see [Table S2](#)) and confirmed a Fe/W molar ratio of 0.68, close to expected for a 100% Fe³⁺-containing iron tungstate.

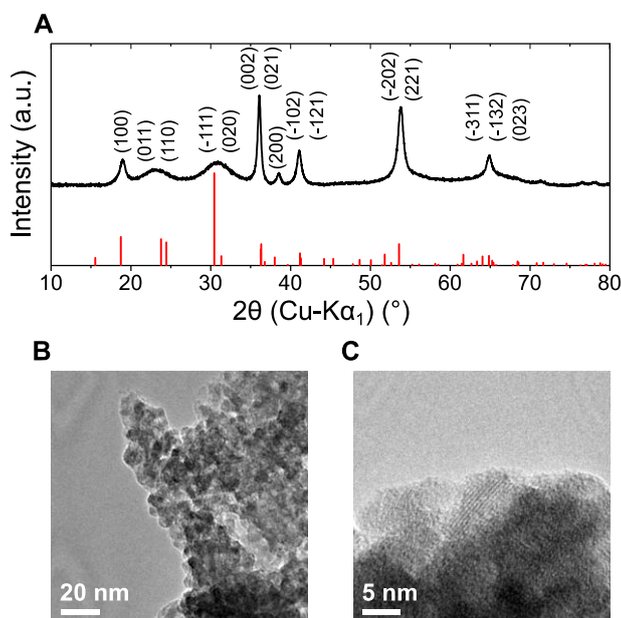


Figure 1: (a) XRD pattern of FeWO_4 synthesized *via* micro-wave assisted coprecipitation route. The main peak reflections are indexed; (b) TEM and (c) HRTEM images of FeWO_4 particles.

The electrodes were then electrochemically tested by cyclic voltammetry in a 5M LiNO_3 aqueous electrolyte (pH = 6.5) at a scan rate of $10 \text{ mV}\cdot\text{s}^{-1}$, from -0.6 to 0 V *vs.* Ag/AgCl. ^[10a] Interestingly, the electrode material exhibits a capacitance of $100 \text{ F}\cdot\text{g}^{-1}$ for the first cycle. This capacitance decreases upon the first 1000 cycles and stabilizes at $50 \text{ F}\cdot\text{g}^{-1}$ which is *ca.* 1.5 times larger than previously reported by us for $\square_y \text{Fe}_{(1-3y)}^{\text{II}} \text{Fe}_{2y}^{\text{III}} \text{WO}_4$ electrode prepared with a polyol-mediated synthesis ^[10a] which may be due to a larger amount of structural defects as depicted from XRD analysis. As demonstrated later on, the pseudocapacitance experimentally measured is directly related to the concentration of native point defects in the $\square_y \text{Fe}_{(1-3y)}^{\text{II}} \text{Fe}_{2y}^{\text{III}} \text{WO}_4$ (FWO) system which explains the discrepancies in the capacitance values displayed by iron tungstate depending on its synthesis route. During this potentiodynamic cycling, XAS spectra were continuously collected at both tungsten L_3 -edge and iron K-edge. To increase the signal-to-noise ratio of the XAS spectra, only an "average" spectrum calculated by averaging 20 consecutive spectra is considered. This translates into one average spectrum every 100 mV (*i.e.* every 10 s) when cycling at $10 \text{ mV}\cdot\text{s}^{-1}$ as depicted in **Figure 2**. Therefore, this procedure allows to accurately tracking changes in both Fe and W absorption spectra induced in potentiodynamic conditions. Further information concerning the XAS data collection procedure can be found in the experimental section.

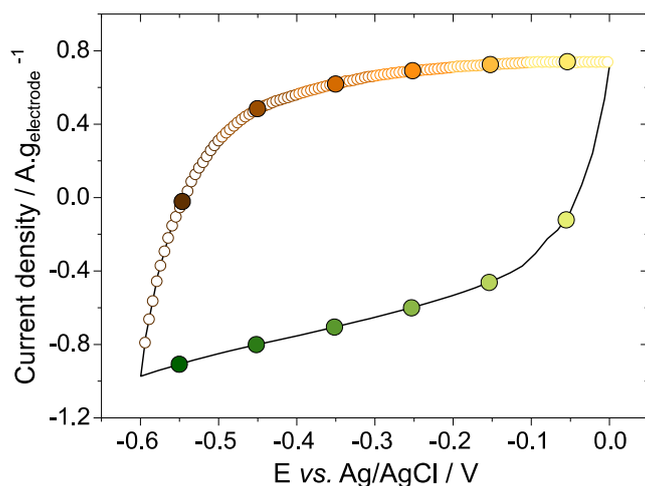


Figure 2: Cyclic voltammogram of the iron tungstate electrode cycled at $10 \text{ mV}\cdot\text{s}^{-1}$ in 5M LiNO_3 aqueous electrolyte (reference electrode Ag/AgCl; counter electrode: platinum wire). The open dots represent the XAS spectra acquisition rate (1 spectrum every 0.5 s), the full dots corresponding to the average of 20 spectra for clarity purpose (1 averaged spectrum every 10 s).

Figure 3A shows the *operando* XANES spectra obtained at Fe K-edge for the iron tungstate electrode during the acquisition of a cyclic voltammogram. A closer insight to the edge region reveals a significant shift of the Fe K-edge toward lower energy values upon cathodic scan of the electrode (**Figure 3B**), and a shift toward higher energy upon anodic scan. Moreover, the pre-peak feature, which is the signature of distorted Fe^{3+}O_6 geometry in the pristine material, progressively disappears upon electrode reduction in favor of a regular of Fe^{2+}O_6 coordination (inset, **Figure 3A**). As well established for the 3d metals, ^[18] the energy shift of the edge position is directly related to the Fe oxidation state change, which, in our case, ranges between Fe^{3+} to Fe^{2+} . Since the electrodes were first cycled toward a cathodic scan, these observations can be summarized as follows: when the potential of iron tungstate electrodes is decreased from 0.0 V vs. Ag/AgCl to lower values, part of the Fe^{3+} in is reduced to Fe^{2+} leading to a shift of the Fe K-edge toward lower energy values. Upon subsequent anodic scan,

while going upward from -0.6 V vs. Ag/AgCl to more oxidative potentials, Fe^{2+} cations are oxidized back to Fe^{3+} . Such a behavior was also observed over several cycles, as depicted in **Figure 4A**, where the Fe K-edge XANES spectra are depicted for the first 9 cycles, confirming the robustness of our finding. A slight irreversibility is observed in the first cycle, probably due to some Li^+ or H^+ trapped in the material. On the other hand, the W L_3 -edge position does not show any significant energy shift under the same *operando* conditions, as can be seen from **Figure 4B**.

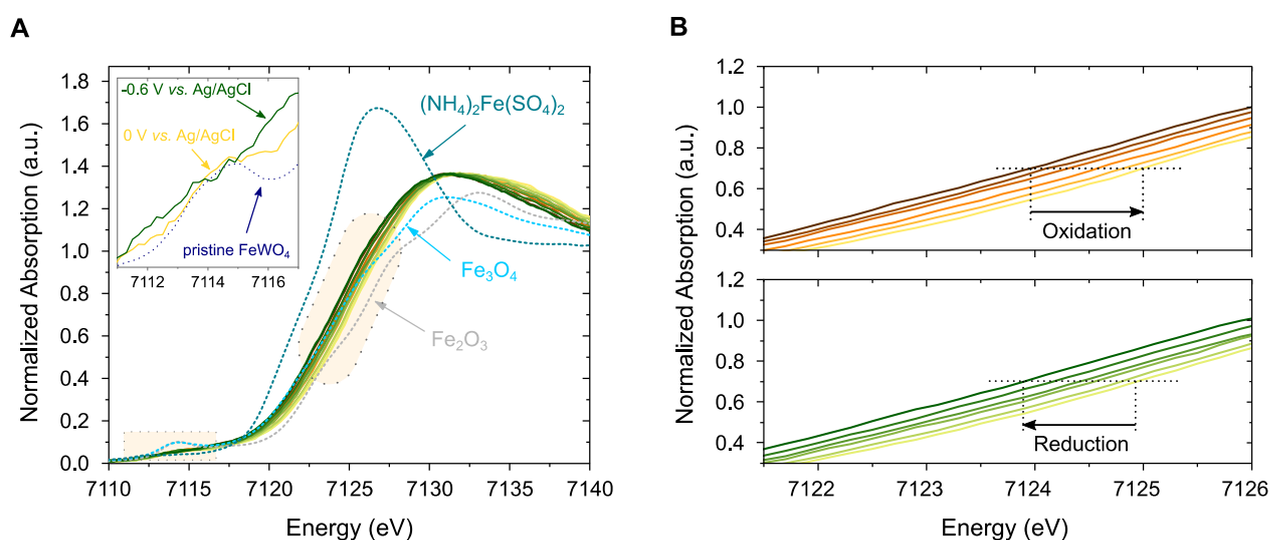


Figure 3: (a) Operando Fe K-edge XANES spectra for the FWO electrode, compared to reference samples $(\text{NH}_4)_2\text{Fe}(\text{SO}_4)_2$ (Fe^{2+} , teal), Fe_3O_4 ($\text{Fe}^{2.67+}$, light blue) and Fe_2O_3 (Fe^{3+} , grey). Inset: zoom in the pre-edge region showing the spectra of the FWO electrode in reduction at -0.6 V (dark green) and in oxidation at 0 V vs. Ag/AgCl (yellow). The pristine electrode (dashed dark blue line) is also shown for comparison.

(b) Enlargement of the normalized absorption evidencing the change in Fe K-edge energy while cycling the electrodes between 0 V vs. Ag/AgCl down to -0.6 V vs. Ag/AgCl at $10 \text{ mV}\cdot\text{s}^{-1}$. The color code of the spectra is the same as the cyclic voltammogram presented in Figure 2.

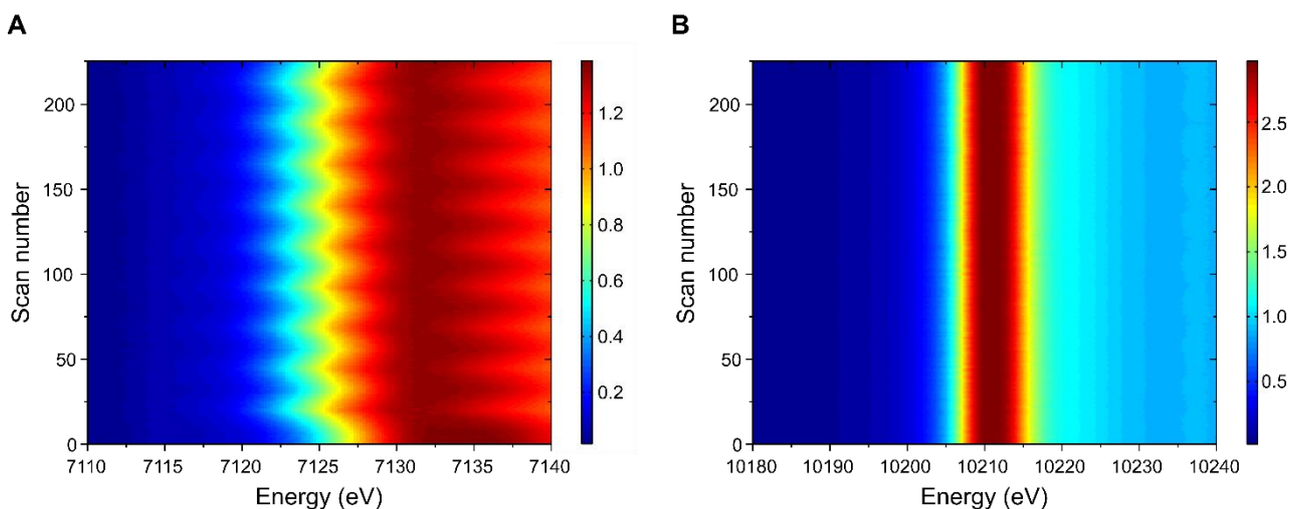


Figure 4: Evolution of the operando XANES spectra for the FWO electrode at (a) Fe K-edge and (b) W L₃-edge during 9 cycles between 0 V vs. Ag/AgCl down to -0.6 V vs. Ag/AgCl at 10 mV.s⁻¹.

For a quantitative and comparative analysis, the edge position corresponding to the energy at 0.7 normalized absorption was considered. In this way, the linear relationship between the edge position and the oxidation state can be used, avoiding the spurious electronic effect related to the different local geometries. [18] The edge positions during the cyclic voltammogram are reported for Fe and W in **Figure 5A and 5B**, respectively. Using Fe²⁺ ((NH₄)₂Fe(SO₄)₂ · 6H₂O), Fe^{2.67+} (Fe₃O₄) and Fe³⁺ (Fe₂O₃) as references, changes in the Fe oxidation state could be quantitatively extracted. Thus, for the iron tungstate electrode, a noticeable and linear change in Fe mean oxidation state is evidenced upon potentiodynamic cycling, corresponding to Fe^{2.55+} oxidized to Fe^{2.80+} upon anodic scan while a reversible change is measured upon cathodic one (**Figure 5A**). It can also be noted that, at a given potential, either in reduction or in oxidation, the energy shift of the Fe K-edge is almost the

same, at less than 0.1 eV, corresponding to a change of ± 0.02 in Fe mean oxidation state. This finding supports the pseudocapacitive behavior hypothesized for this multicationic oxide.^[10a] At such a high measurement rate, the observed change in the iron mean oxidation state is not related to any long polarization time at a constant potential value, as it could be argued from previous *in situ* studies. Even at such a fast cycling rate, the role of Fe cation redox centers is thus unambiguously evidenced. Regarding the redox activity of W, the *operando* W L₃-edge XANES spectra do not show any edge shift during cycling, as can be observed from **Figures 4B** and **5B**, thus emphasizing that W⁶⁺ cations act as spectator cations which do not participate in the charge storage mechanism of the electrode.

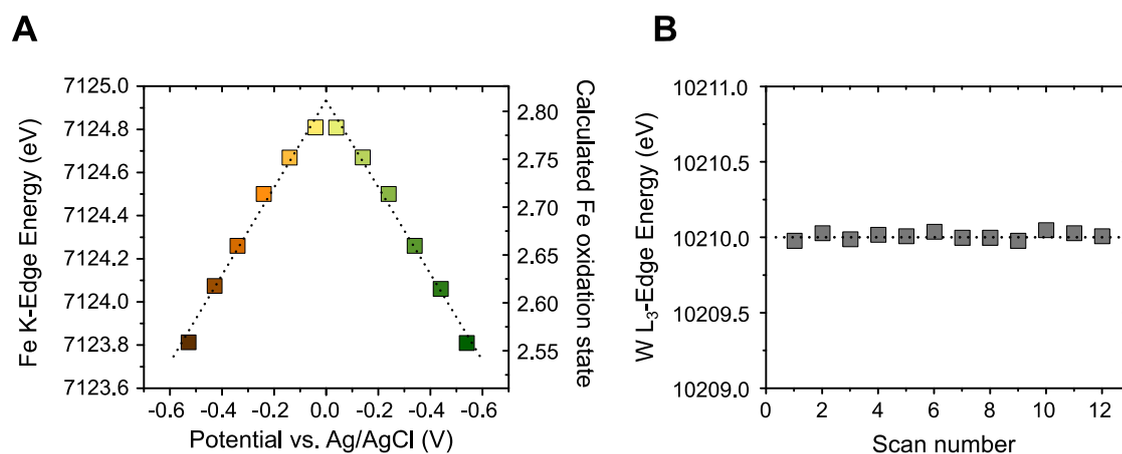


Figure 5: Evolution of the (a) Fe K-edge energy and mean Fe oxidation state and (b) W L₃-edge as a function of electrode potential when cycled between 0 V vs. Ag/AgCl down to -0.6 V vs. Ag/AgCl at 10 mV.s⁻¹.

To estimate the Faradaic contribution to the capacity, the change in the iron oxidation state upon cycling the FWO electrode between -0.6 V and 0 V vs Ag/AgCl was considered to correspond to an electronic transfer of *ca.* 0.25 electron per iron center (a variation from

Fe^{2.55+} to Fe^{2.80+}, **Figure 5A**). This number of exchanged electrons can then be translated into a capacity using equation (1):

$$Q = \frac{F \times n}{M} \quad (1)$$

where Q is the calculated gravimetric capacity of the sample (C.g⁻¹), F is the Faraday constant (96 485 C.mol⁻¹), n is the number of electrons exchanged, and M is the molar mass of the material (g.mol⁻¹). Considering the formula $\square_{1/3}Fe_{2/3}WO_4$ of our sample and the corresponding values for n and M ($n = \frac{2}{3} \times 0.25 \approx 0.167$ and $M = 285$ g.mol⁻¹), a capacity of 56 C.g⁻¹ can then be deduced from the *operando* XAS experiment, which is consistent with the value obtained from the cyclic voltammetry experiments ($Q_{CV} = 60$ C.g⁻¹).

All the results obtained by XANES are also confirmed using the chemometric approach described in detail in the Supporting Information (see **Figure S7** and **S8**). To get further insights about the structural evolution of the iron tungstate under *operando* conditions, the extended X-ray absorption fine structure (EXAFS) regions at Fe K- and W L₃-edges were analyzed (**Figure 6**). **Figure 6a** shows the magnitude of the Fourier transforms $|FT(k^2\chi(k))|$ of the EXAFS oscillations corresponding to the pristine and the reduced electrode at -0.6 V vs Ag/AgCl. The main peak at ~ 1.5 Å is due to Fe-O bond lengths (4 O at 1.96 Å and 2 O at 2.01 Å), while the peak at ~ 2.5 Å corresponds to the Fe-Fe and Fe-O bond lengths (2 Fe atoms at ~ 3.04 Å and 4 O at 3.36 Å) and the peak at ~ 3.2 Å corresponds to the Fe-W bond length. When the electrode is polarized at -0.6 V vs Ag/AgCl, the FeO₆ octahedron gets regular with 6 oxygen at 2.03 Å, as also shown by the disappearance of the pre-peak. The shortening of Fe-O bondlengths (**Table S3**) confirms the Fe³⁺/Fe²⁺ redox activity. On the other hand, no structural evolution could be detected from EXAFS data at W L₃-edge upon cycling the electrode (**Figure 6c, d**), which is also consistent with the XANES analysis.

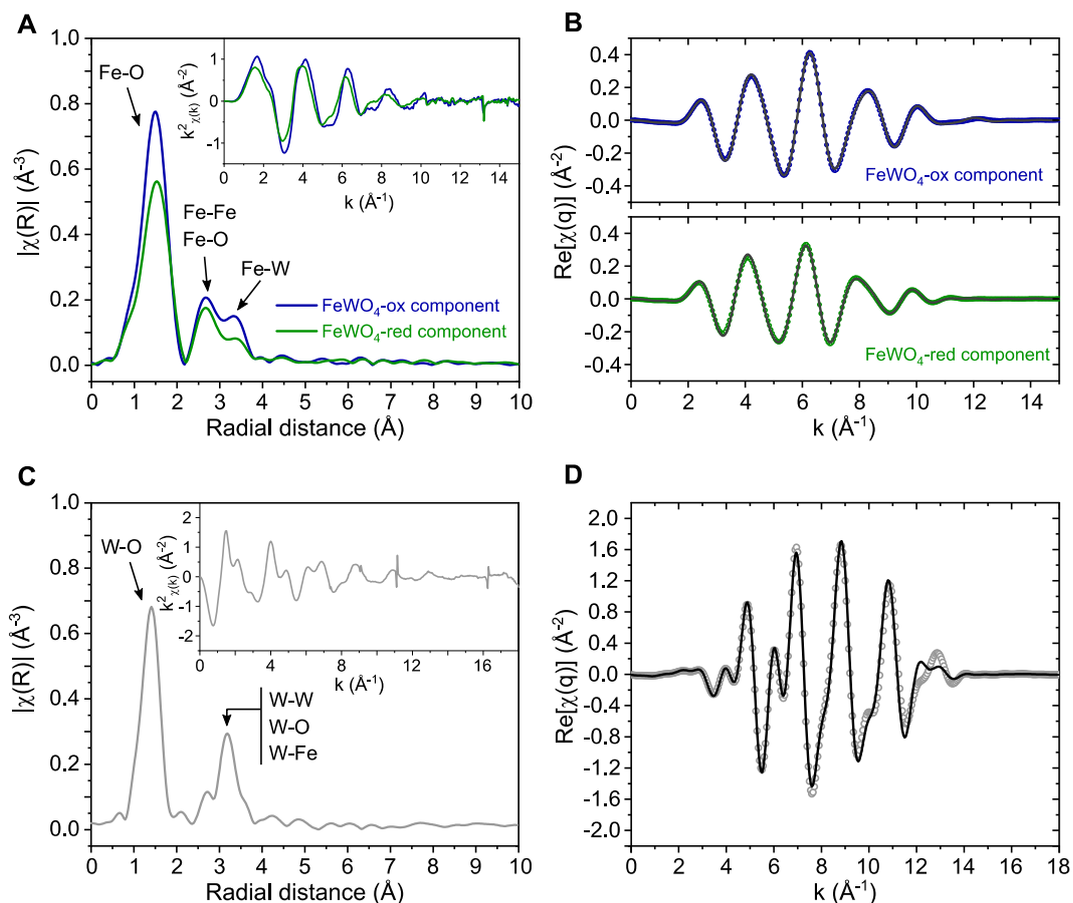


Figure 6: Magnitude of Fourier transforms of k^2 -weighted EXAFS oscillations for FeWO_4 's first cycle along with fitting results. (a, b) Principal components analysis (PCA) at Fe K-edge showing the two reconstructed principal components, *i.e.* FeWO_4 -ox (dark blue, pristine sample) and FeWO_4 -red (dark green, reduced electrode at -0.6 V vs. Ag/AgCl); c) W L_3 -edge spectrum, along with the fitting result (d). No structural evolution is detected at W L_3 -edge during cycling.

It can thus be concluded that the tungsten in the iron tungstate structure acts as a spectator cation and not as an active redox center. Such a spectator cation helps stabilizing the structure even if electrochemical changes affect the other cations of the multicationic oxide. Indeed, the WO_6 octahedra backbone of FWO is not affected by the electrochemical cycling while FeO_6 octahedra does. As a result, only a weak amorphization of the surface of FeWO_4 particles is depicted after 10000 charge/discharge cycles (**Figure S9**). Moreover, first-principles DFT+U

calculations performed on the bulk phases confirm that the tungsten atomic charge does not change with and that the DOS projected on Fe and W invariably displays Fe(d)-bands below and above the Fermi level (see Supporting Information for details). This clearly supports that iron should be the main redox center of both the anodic and cathodic scans, as deduced from *operando* XAS spectra. To definitely assess this statement, Fukui functions were computed for Fe-defective bulk phases. The Fukui function is a pertinent electrochemical descriptor as it measures the response of the system electron density to an electronic charge (electron or hole), including polarization effects. ^[21] Adding a fraction of electron or hole to the system therefore allows following how the electron density dynamically adapts to a cathodic or anodic scan, respectively. As shown in Supporting Information (**Figure S6**), the oxidation (resp. reduction) of iron is confirmed for the anodic (resp. cathodic) scan. The results also reveal that in the wolframite structure, the tungsten is likely to participate to the charge storage mechanism, but only if the material is a perfect crystal (no defect). This is therefore not the case for the Fe-deficient material studied herein. However, one should note that even in the case of a natural wolframite crystal, small amounts of Fe³⁺ (and therefore defects) can be detected, as shown by MacKenzie *et al.* ^[22] Therefore, triggering the redox activity of W in iron tungstate seems unfortunately highly unlikely, especially considering the fact that nanosized particles are generally needed to reach high capacitance values in a pseudocapacitive material. The computed Fukui functions also illustrate that, interestingly, the content of which is linearly correlated to the Fe(d) band integration around the Fermi level, is also correlated to the capacity obtained in our different samples. This demonstrates that the pseudocapacitance experimentally measured is directly related to the concentration of native point defects in the $\square_y Fe_{(1-3y)}^{II} Fe_{2y}^{III} WO_4$ system. This finding can therefore explain the discrepancies in the capacitance values displayed by iron tungstate depending on its synthesis route. ^[10a]

3. Conclusion

In this study, *operando* X-ray absorption spectroscopy was used to show how the capacitive-like electrochemical signature of FeWO₄-based electrode can be unambiguously assigned to a pseudocapacitive charge storage. The experiments were performed upon operating the electrode at a realistic timescale (one-minute charge/discharge cycles), thus enabling to monitor the variations of the iron and tungsten oxidation states occurring in the FWO material, unveiling its charge storage behavior upon cycling. Our results lead to the conclusions that i) iron tungstate is a pseudocapacitive material since the charge storage mechanism involves a linear variation of the Fe oxidation state over the applied potential window, ii) the capacity measured with the use of electrochemical characterization fits well with the redox capacity related to Fe³⁺/Fe²⁺ redox couple, iii) these findings are in good agreement with the charge storage mechanism recently predicted and presently confirmed either by *operando* XAS measurements and Fukui function calculations, iv) W⁶⁺ acts as a spectator cation, probably helping to maintain the crystallographic structure upon cycling the electrode.

This *operando* XAS approach can be therefore advantageously used to investigate the electrochemical behavior of other pseudocapacitive multicationic oxides or transition metal nitrides for which the charge storage mechanism is still speculative. This fundamental knowledge is crucial to facilitate the research and the design of new pseudocapacitive materials and architectures with improved performance.

4. Experimental Section

Reagents: All reagents used in this work were purchased commercially from Sigma-Aldrich or Alfa Aesar, were at least 99% purity and were used as received.

Materials synthesis: Iron tungstate (FWO) was synthesized *via* a microwave-assisted coprecipitation route. First, stoichiometric 2 M aqueous solutions of Fe^{2+} and W^{6+} precursors were obtained by separately dissolving iron (II) chloride and sodium tungstate in distilled water. Then, both solutions were slowly mixed together under vigorous stirring before being subjected to microwave heating. The synthesis was carried out at a constant temperature of 100°C with a 200 W power for 1 h in a SAIREM MiniFlow 200SS microwave reactor. The obtained brown powder was centrifugated and resuspended in water and ethanol several times, before being dried overnight at 35°C under primary vacuum.

X-ray powder diffraction: X-ray diffraction patterns of FeWO_4 were collected in Bragg-Brentano configuration with a PANalytical X'Pert Pro diffractometer using $\text{Cu K}\alpha$ radiation, equipped with an X'Celerator detector operating at 40 kV – 40 mA in the range $2\theta_{\text{Cu K}\alpha} = 10 - 90^\circ$ with a 2θ step size of 0.017°.

Electron microscopy: The size and morphology of the synthesized FWO nanoparticles were characterized using a Hitachi H9000-NAR transmission electron microscope. Prior to examination, the samples were ultrasonically dispersed in ethanol. A drop of the suspension was then deposited on a copper grid previously covered with a thin holey carbon film.

Electrodes preparation: All electrodes were prepared using FWO, carbon black (100% compressed; Alfa Aesar; > 99.9 %) and polytetrafluoroethylene (60 wt % dispersion in H_2O , Sigma Aldrich) with weight ratios of 60:30:10, as described elsewhere.^[23] Ethanol was added

to the components and the obtained mixture was stirred on a hot plate to slowly and partially evaporate the solvent. The resulting homogeneous paste was cold-laminated into thick films (120-150 μm) that were subsequently dried at 60°C in air. Finally, disk-shaped electrodes (10 mm diameter) were cut out from the films and pressed at 900 MPa onto stainless steel grid current collectors. Rectangle-shaped holes (1x5 mm) were drilled in all current collectors beforehand for XAS measurements (see **Figure S2B**). All the electrodes used in this study displayed a mass loading of $10.0\pm 0.3 \text{ mg}_{\text{FeWO}_4}\cdot\text{cm}^{-2}$, which is a standard value for real-life commercial devices.

Electrochemical measurements: The prepared electrodes were electrochemically characterized using a VMP3 galvanostat-potentiostat from Biologic run under EC-Lab software. All the experiments were conducted in a three-electrode configuration using a platinum wire and an Ag/AgCl (3M NaCl) assembly as the counter and reference electrode, respectively. FWO composite electrodes were tested in cyclic voltammetry between -0.6 and 0 V vs. Ag/AgCl in an aqueous 5 M LiNO₃ electrolyte at room temperature.

Electrochemical cell for operando X-ray Absorption Spectroscopy: The operando measurements were performed in a three-electrode cell assembly that has been specifically designed and manufactured in our lab for this study. The FWO working electrode was placed in the electrolyte between two thin (10 μm) Kapton[®] windows. Both reference and counter electrodes were placed on the sides of these windows so that the thickness of electrolyte is limited to 1.5 mm, which enables quick data acquisitions at the investigated energy ranges. More detailed information on the electrochemical cell assembly can be found in **Figure S2**.

X-ray Absorption Spectroscopy (XAS): The X-ray absorption spectroscopy experiments were performed at the ROCK beamline of SOLEIL synchrotron radiation facility (France).^[24] The energy of the synchrotron radiation emitted by a bending magnet source was selected using a Si(111) channel-cut quick-XAS monochromator. The spectra were collected first at Fe K-edge (7112 eV) and then at W L₃₋₁-edges (from 10200 to 12200 eV) using a quick-XAS monochromator oscillating with a frequency of 2 Hz with an amplitude of 2.4°.

In 0.5 second, two spectra were acquired (for example at Fe K-edge), one with increasing Bragg angles followed by one with decreasing Bragg angles. We have then considered only the upward Bragg angles and we have averaged out the XAS spectra over periods of 10 seconds. It is important to mention that we acquire a full XAS spectrum, *i.e.* the near edge and the extended fine structure regions. The XAS acquisitions were carried out in transmission mode using three ionization chambers mounted in series for simultaneous measurements on the electrode and a reference. The *in situ* electrochemical cell was placed between the first and the second ionization chambers. The energy calibration was established with simultaneous absorption measurements on a Fe or a W metal foil placed between the second and the third ionization chamber. Standard procedure was used to calibrate the energy and normalize the spectra using the Demeter package^[25] and a Python routine developed for PCA analysis.^[26] Commercial powders of (NH₄)₂Fe(SO₄)₂ · 6H₂O, Fe₃O₄ and Fe₂O₃ were used as Fe²⁺, Fe^{2.67+} and Fe³⁺ references, respectively.

Before running *operando* XAS measurements, the electrodes were placed in the *in situ* cell and soaked in the electrolyte for several minutes at open-circuit potential (OCP) (reference electrode Ag/AgCl; counter electrode: platinum wire). In such conditions, XAS measurements were performed at both Fe K-edge and W L₃-edge in order to determine the valence of both iron and tungsten in the pristine material. The corresponding spectra are presented in **Figure S3 and S4**.

The XAS data related to the first 10 cycles at the Fe K- and W L_{1,3}-edges were analyzed using a chemometric approach based on a combination of Principal Component Analysis (PCA) and Multivariate Curve Resolution Alternating Least-Squares (MCR-ALS).^[27] Principal Component Analysis (PCA) was applied in order to evaluate the number of orthogonal components needed to describe the whole evolution of the XAS spectra for each edge. We found that only the Fe K-edge XAS spectra were affected by the applied potential. Then, Multivariate Curve Resolution alternating Least-Squares (MCR-ALS) was used to reconstruct the 2 principal components (PCs) at Fe K-edge. The extended X-ray absorption fine structure (EXAFS) oscillations extracted at Fe K-edge for the PCS and at W L₃-edge on the pristine material were modeled using the Artemis software.^[25] As input model to refine the EXAFS oscillation, we have used the structure reported by Almeida and coworkers.^[28] The details about the EXAFS analysis are provided in the SI.

Supporting information

Supporting Information is available from the Wiley Online Library or from the author.

Acknowledgements

This work was supported by a public grant overseen by ANR IVEDS (ANR-15-CE05-0011-01), the Université de Nantes and the French “Réseau sur le stockage électrochimique de l'énergie” (RS2E) through the project ANR-10-LABX-76-01. This study was also supported by the Global Innovation Re-search Organization in TUAT. The authors would like to thank P. Léone (IMN) and M. Sougrati (ICGM) for running the Mössbauer spectroscopy experiments and F. Guillou (IMN) for his valuable help with the design and the fabrication of the electrochemical cell. We also thank S. Belin for her help during the experiment 20151183 on the ROCK beamline (financed by the French National Research Agency (ANR) as a part

of the "Investissements d'Avenir" program, reference: ANR-10-EQPX-45) of SOLEIL synchrotron.

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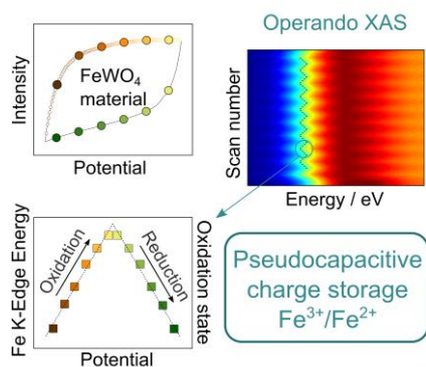
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Unveiling Pseudocapacitive Charge Storage Behavior in FeWO₄ Electrode Material by Operando X-ray Absorption Spectroscopy

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TOC :



Operando X-ray Absorption Spectroscopy (XAS) has been used to elucidate the charge storage mechanism of FeWO₄ **when used as a negative electrode material in an aqueous electrochemical capacitor**. Both Fe K-edge and W L₃-edge measurements point out the involvement of the Fe³⁺/Fe²⁺ redox couple, while W⁶⁺ acts as a spectator cation, **demonstrating the pseudocapacitive charge storage mechanism of FeWO₄**.

Supporting Information

Unveiling Pseudocapacitive Charge Storage Behavior in FeWO_4 Electrode Material by Operando X-ray Absorption Spectroscopy

*Nicolas Goubard-Bretesché, Olivier Crosnier, Camille Douard, Antonella Iadecola, Richard Retoux, Christophe Payen, Marie-Liesse Doublet, Kazuaki Kisu, Etsuro Iwama, Katsuhiko Naoi, Frédéric Favier, and Thierry Brousse**

Supporting figures

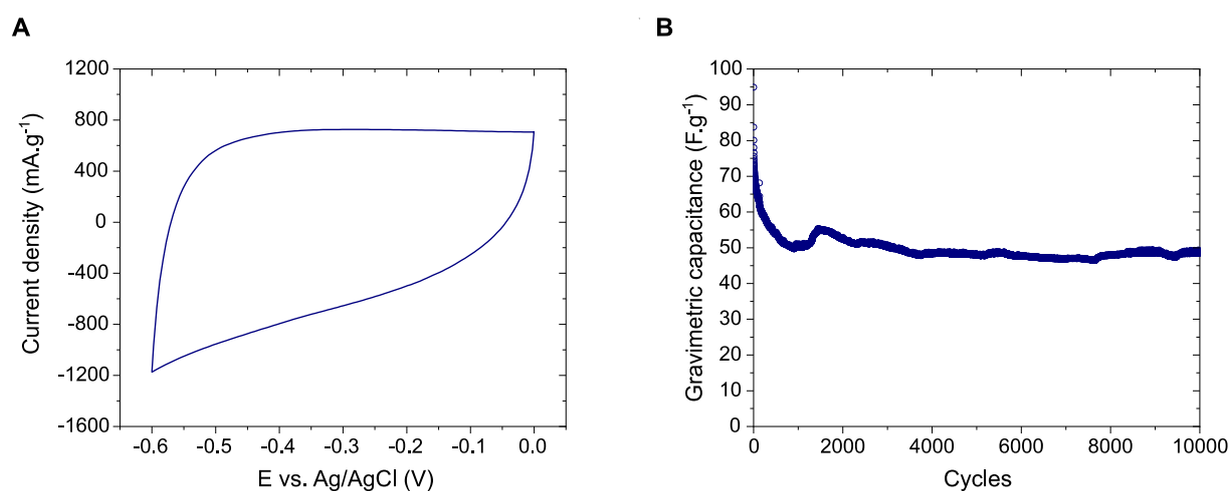


Figure S1: A) Typical voltammogram and B) Long term cycling behavior of the FeWO_4 material over 10 000 cycles. Electrolyte: 5M LiNO_3 ; Scan rate: 20 mV.s^{-1} . The decay observed on the first 1000 cycles maybe due to a rearrangement at the surface of the particles during cycling.

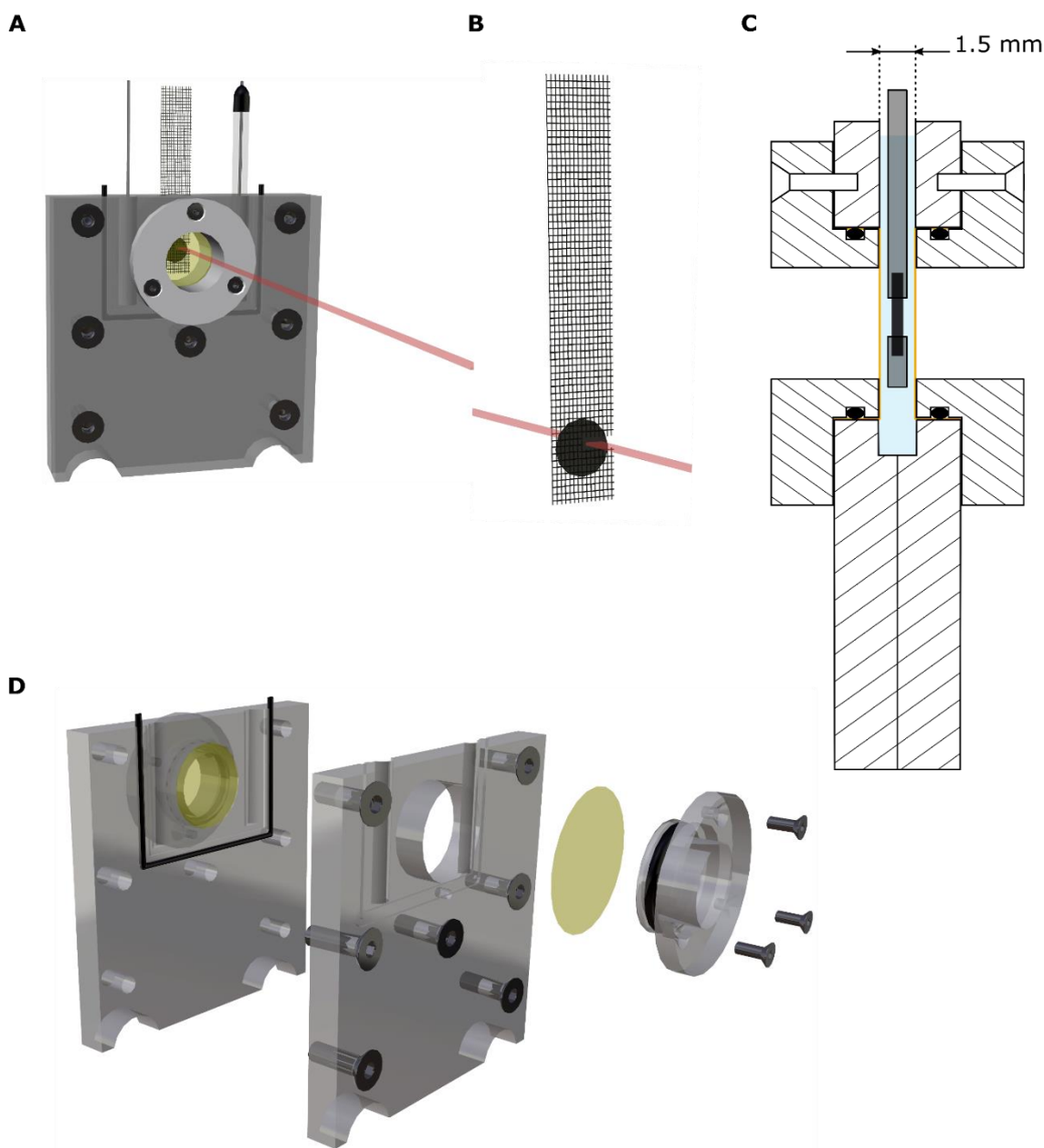


Figure S2: 3D Representation of the electrochemical cell used for the *operando* experiments. The FeWO_4 working electrodes were immersed in a 5M LiNO_3 aqueous electrolyte confined between two polyimide (Kapton®) $10\mu\text{m}$ -thick films (represented in yellow), separated by a distance of 1.5 mm from each other. The reference and counter electrodes were put on both sides of the Kapton films to avoid increasing the thickness of electrolyte. A) General overview with the platinum counter electrode, the working electrode and the Ag/AgCl reference electrode; B) Typical FeWO_4 electrode, with a rectangle-shaped hole of $5\text{ mm} \times 1\text{ mm}$ cut in the current collector (stainless steel grid) to avoid any absorption of the incident beam (represented in red); C) Cross-sectional schematic view of the inside of the *in situ* cell; D) Exploded view.

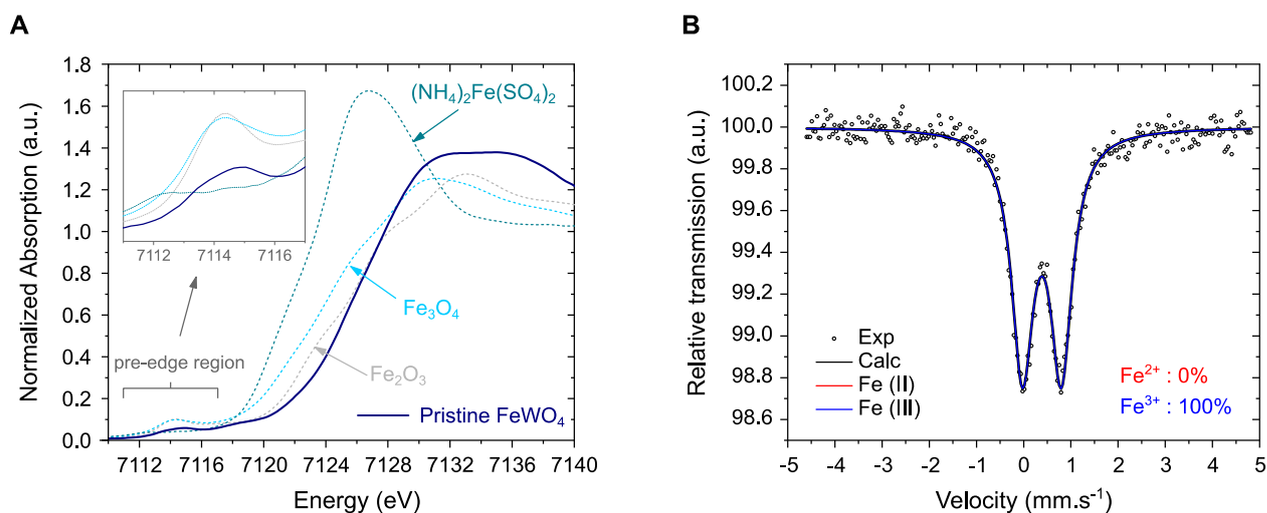


Figure S3: A) XANES spectra of FeWO_4 (dark blue), and the iron-containing references used in the study. Inset: zoom in the pre-edge region; B) Mössbauer spectrum of the FeWO_4 sample, along with its calculated Fe^{2+} and Fe^{3+} contents. The pristine material only contains Fe^{3+} .

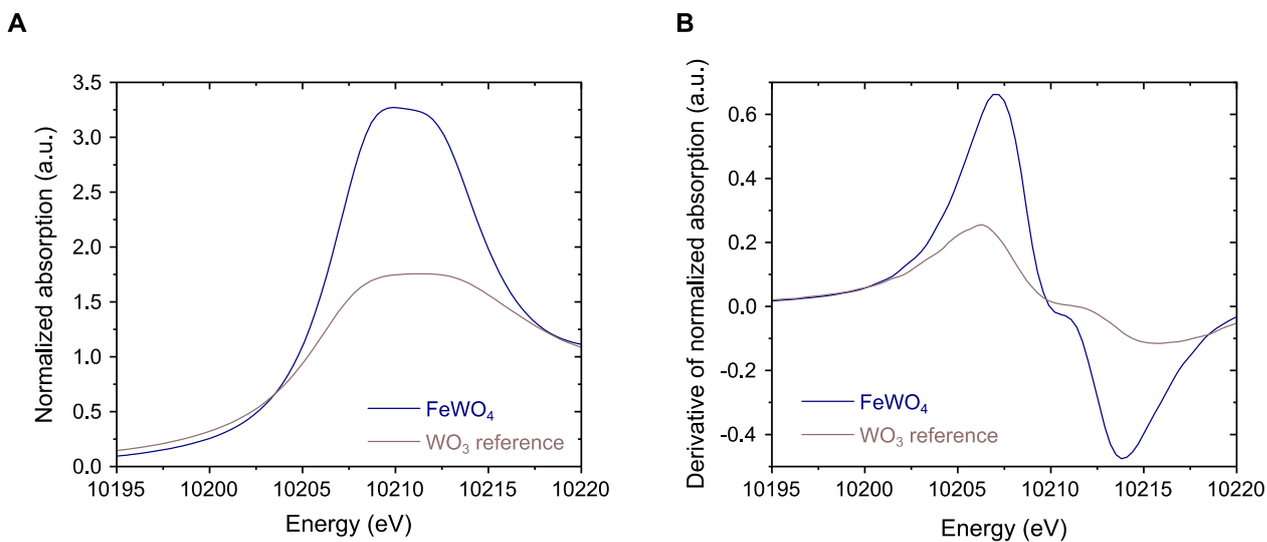


Figure S4: A) W L_3 -edge XANES spectra of the FeWO_4 electrode (dark blue), compared to the WO_3 reference sample (W^{6+} , brown); B) Corresponding derivative spectra showing the W^{6+} oxidation state of tungsten in the pristine FeWO_4

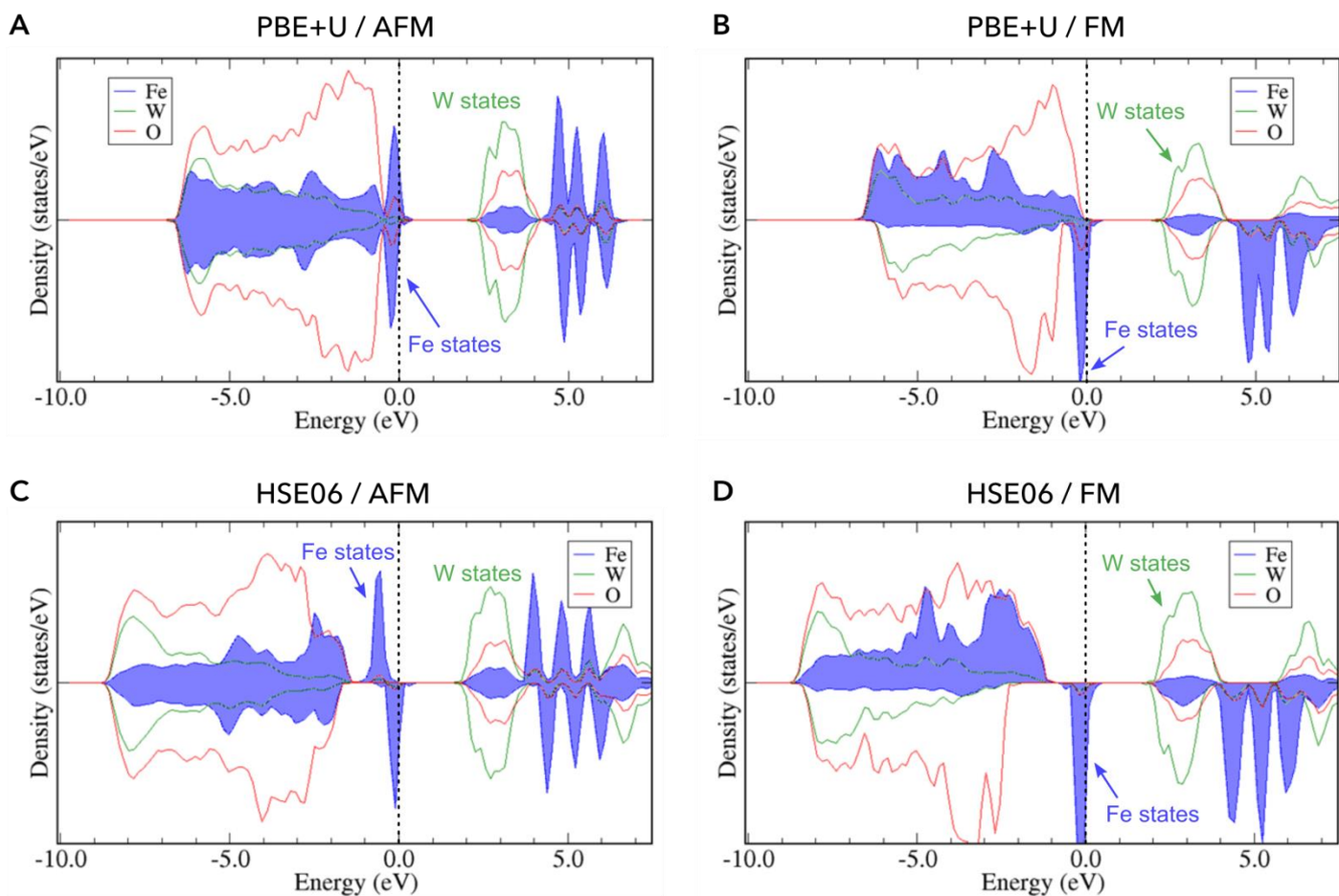


Figure S5: Atom-projected Density of states for the stoichiometric FeWO_4 material, computed within the DFT framework using various XC functionals: PBE+U (A, B) and HSE06 (C, D), and different spin configurations: Antiferromagnetic AFM (A, C) and Ferromagnetic FM (B, D). All calculations invariably show a narrow Fe-band below the Fermi level consistent with the polaronic effect already reported for this system and a wider W-band above the Fermi level. Note that a small fraction of holes is already visible in the stoichiometric FeWO_4 material, in full agreement with the occurrence of polarons and Fe-deficiency in that system.

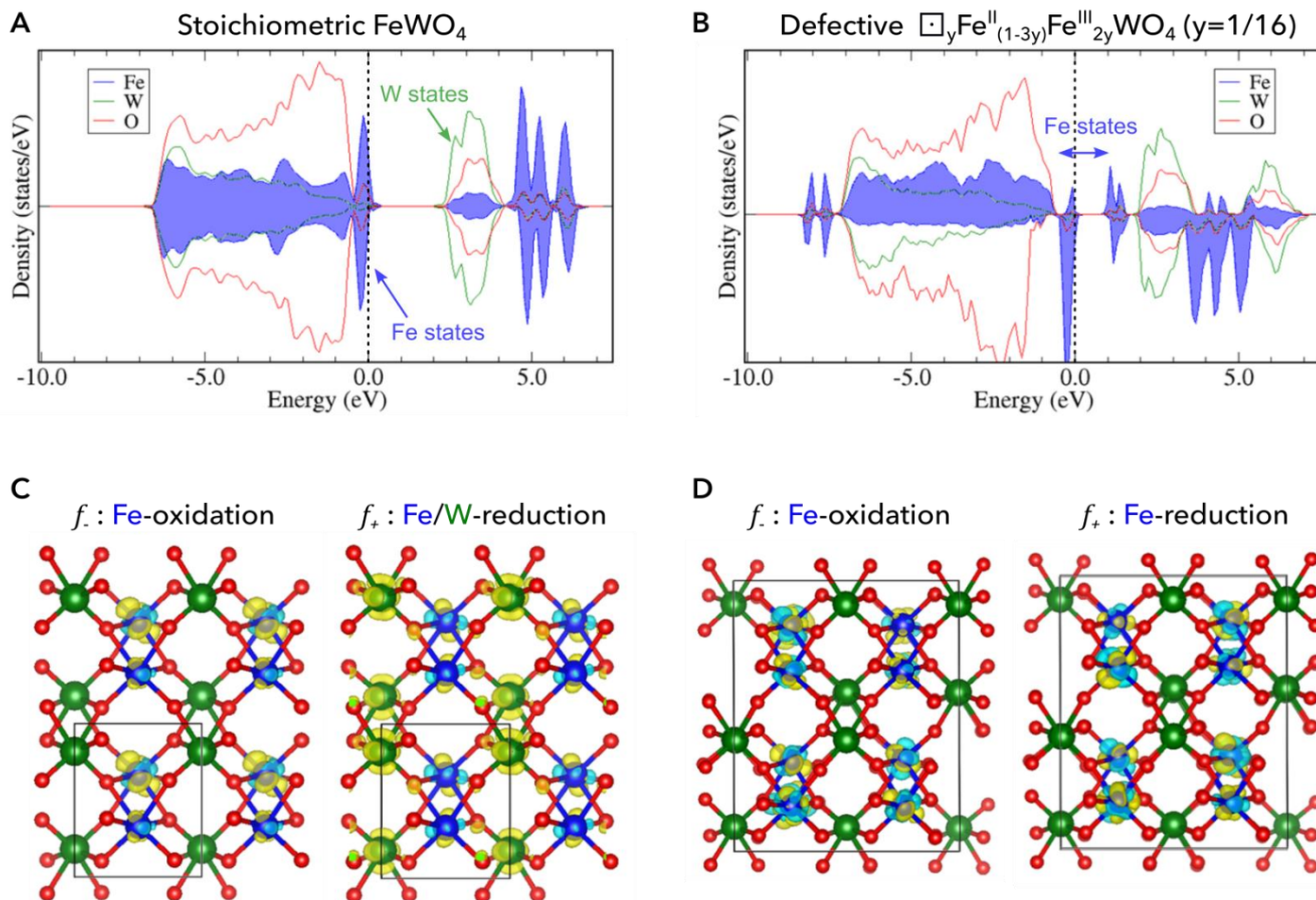


Figure S6: Atom-projected Density of states for A) stoichiometric FeWO_4 and B) Fe-defective $\square_y \text{Fe}_{(1-3y)}^{\text{II}} \text{Fe}_{2y}^{\text{III}} \text{WO}_4$ bulk, as well as Fukui functions f_+ and f_- for C) stoichiometric FeWO_4 and D) Fe-defective $\square_y \text{Fe}_{(1-3y)}^{\text{II}} \text{Fe}_{2y}^{\text{III}} \text{WO}_4$ bulk, as computed from first-principles DFT calculations in the AFM spin configuration. Iron, tungsten and oxygen are represented in blue, green and red, respectively both in the pDOS plots and in the crystal structures. The Fermi level is indicated by the vertical dotted line in the pDOS. The yellow and blue volumes of the Fukui functions represent the accumulation and depletion of charge around each atom. For the stoichiometric FeWO_4 , the pDOS shows a narrow Fe-band below the Fermi level and a wider W-band above the Fermi level. Accordingly, the Fukui functions f_- (hole addition) and f_+ (electron addition) show that Fe and W are the main centers of the oxidation (anodic scan) and reduction (cathodic scan), respectively. When Fe-vacancies are introduced in the FeWO_4 supercell (16 formula units), the splitting of the Fe-states around the Fermi level leads to a different electrochemical response of the system with Fe being the redox center of both the anodic and cathodic scans, even at very low y -content (here 1/16).

Note that the increase of y is directly correlated to the increase of the Fe-band DOS above the Fermi level and therefore with the increase of the capacitance of the cathodic scan.

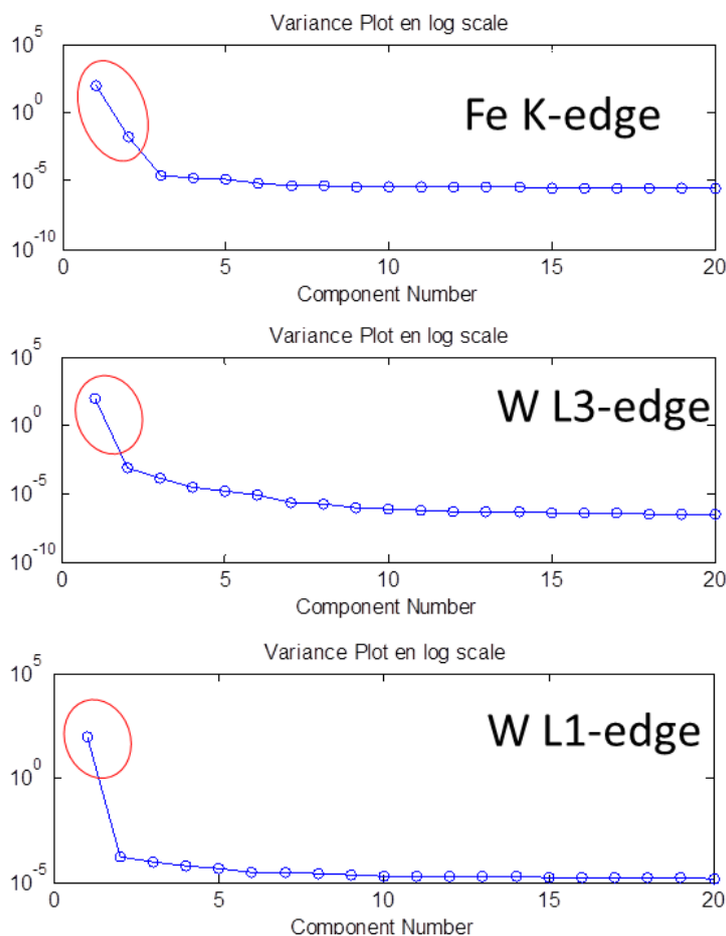


Figure S7: Variance plot in logarithmic scale on the XAS dataset at Fe K- and W $L_{1,3}$ -edges related to the first 10 cycles. It is clear that 2 principal components are needed to describe the whole evolution of the XAS data at Fe K-edge while no evolution is found for the W L-edges XAS data.

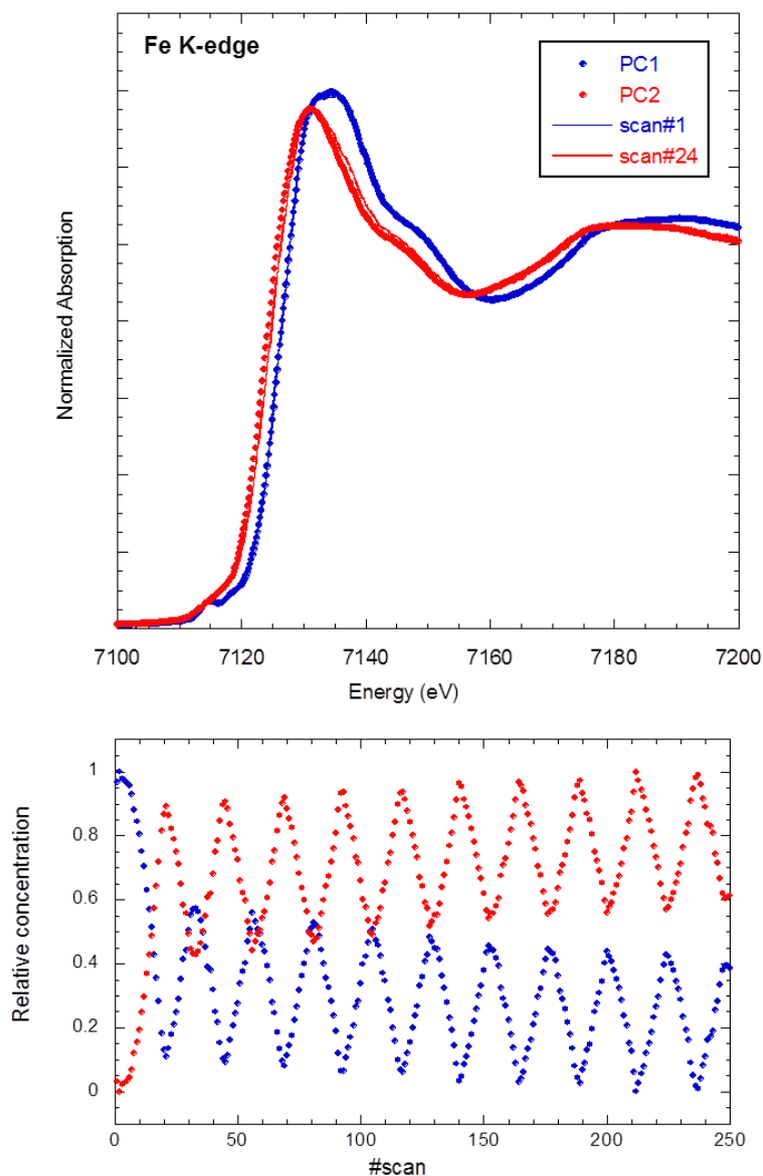


Figure S8: Fe K-edge XANES spectra of the reconstructed components (PC1 and PC2) along with the corresponding operando scan (#1 and #24); the relative concentration of the PC1 and PC2 species during the first 10 cycles, confirming the robustness of the charge storage mechanism in FeWO_4 . PC1 corresponds to the reconstructed component of the pristine sample ($\text{FeWO}_4\text{-ox}$, in **Figure 6** of the manuscript) and PC2 to the reduced state (at -0.6 V vs. Ag/AgCl, *i.e.* $\text{FeWO}_4\text{-red}$ in **Figure 6**).

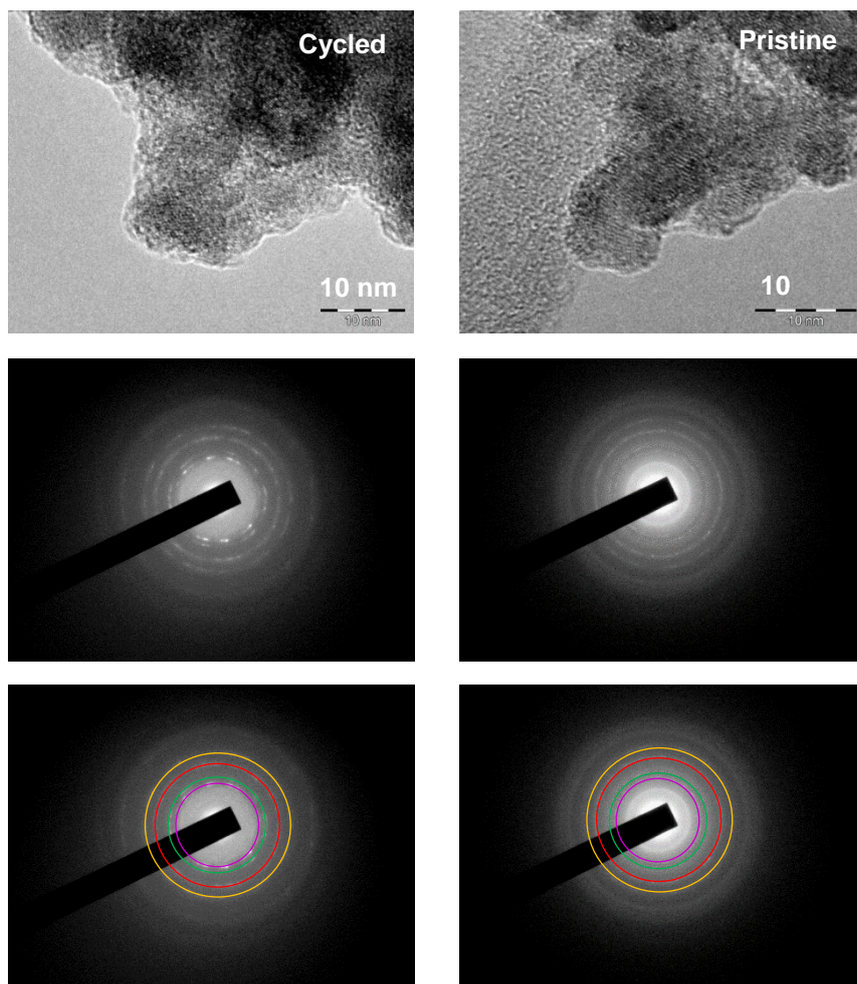


Figure S9: HRTEM images of uncycled (pristine) and cycled (10000 charge/discharge cycles) of iron tungstate particles. Electron diffraction only shows a slight amorphization of the particle surface after 10000 cycles.

Supporting Tables

Table S1: Summary of the characteristics of the FeWO₄ sample used as the electrode material

Particle size (TEM Images)	~ 8x15 nm
Specific surface area S _{BET} (BET method, N ₂ adsorption)	73 m ² .g ⁻¹
Mean diameter from S _{BET} (Based on spherical particles)	11 nm
Measured density (He pycnometry)	5.12(3) g.cm ⁻³
Fe mean oxidation state (Mössbauer spectroscopy)	3.00

Table S2: ICP-OES analysis of Fe and W in the iron tungstate sample solution.

Element	Emission wavelength (nm)	Replicate 1 (ppm)	Replicate 2 (ppm)	Replicate 3 (ppm)
Fe	234.350	31.10	31.08	31.06
Fe	238.204	30.82	30.80	30.79
Fe	259.940	30.73	30.62	30.69
W	207.912	148.03	147.53	147.71
W	209.475	148.53	148.83	148.78
W	220.449	149.75	150.01	150.61

Fe Concentration (average): 30.85 ppm

W Concentration (average): 148.86 ppm

Calculated molar ratio Fe/W in the sample: 0.68

Table S3: Summary of the EXAFS analysis parameters. The variation of the Fe-O and Fe-Fe bondlengths between the pristine sample and the reduced electrode confirms the Fe³⁺/Fe²⁺ redox activity, while no bondlength modification is detected at W L₃-edge.

Pristine (PC1) Fe K-edge	Reduced (PC2) Fe K-edge	Pristine W L ₃ -edge
4 x O at 1.96 Å	6 x O at 2.03 Å	4 x O at 1.80 Å
2 x O at 2.01 Å	2 x Fe at 3.07 Å	2 x O at 2.10 Å
2 x Fe at 3.04 Å	4 x O at 3.37 Å	2 x W at 3.21 Å
4 x O at 3.34 Å		8 x O at 3.30 Å
		8 x Fe at 3.56 Å
σ^2 (Fe-O) = 0.009 Å ²	σ^2 (Fe-O) = 0.013 Å ²	σ^2 (W-O) = 0.007 Å ²
σ^2 (Fe-Fe) = 0.014 Å ²	σ^2 (Fe-Fe) = 0.016 Å ²	σ^2 (W-W) = 0.007 Å ²
S_0^2 = 0.8	S_0^2 = 0.8	S_0^2 = 1
ΔE_0 = -4.0 eV	ΔE_0 = -2.3 eV	ΔE_0 = 2.5 eV
R-factor = 0.0027	R-factor = 0.0042	R-factor = 0.0055

Supporting Experimental Section

Specific Surface Area measurements: Specific surface areas were obtained by Brunauer–Emmett–Teller (BET) analyses performed with a Quantachrome Nova 4200e operated with nitrogen gas.

Density measurements: Density measurements were carried out by pycnometry under He pressure using an AccuPyc 1330 system from Micromeritics Instrument Corporation.

Mössbauer Spectroscopy: The ⁵⁷Fe Mössbauer spectrum was collected at ambient temperature in transmission geometry on a constant acceleration spectrometer using a ⁵⁷Co(Rh) γ -ray source. Velocity and isomer shift calibrations were performed using α -Fe as a standard.

Quadrupole splitting data were analyzed as discrete 0.1 mm.s^{-1} step distributions in the 0 to 6 mm.s^{-1} range and fitted with Gaussian distribution curves.

DFT Calculations: Spin-polarized density functional theory (DFT) calculations were performed using the plane-wave density functional theory VASP (Vienna ab initio simulation package) code^[S1] within the generalized gradient approximation of Perdew–Burke–Ernzerhof (PBE)^[S2] or the range-separated (HSE06)^[S3] hybrid functionals for exchange and correlation potentials. The rotationally invariant Dudarev method (DFT + U)^[S4] was used to correct the self-interaction error of conventional DFT for correlated d-electrons with $U = 5.3 \text{ eV}$ and 2 eV for the Fe and W, respectively.

Fukui functions are used as a probe of the highest-occupied and lowest-unoccupied states of a system. It represents the variation of the electron density of a given system when a fraction of electron or hole is added to the system. It is a typical electrochemical descriptor as it allows identifying the redox center of the cathodic and anodic reactions.^[S5]

EXAFS Analysis: The k^2 -weighted EXAFS oscillation were extracted at Fe K-edge and at W L_3 -edge using a Gaussian window in the k range $[2.4, 12] \text{ \AA}^{-1}$ and $[3.6, 13.4] \text{ \AA}^{-1}$, respectively. The Back-Fourier transforms were performed in the R range $[1, 3] \text{ \AA}$ at Fe K-edge and $[1.05, 3.8] \text{ \AA}$ at W L_3 -edge using the Hanning window. The phase shifts and structural parameters were obtained by Demeter package, using the crystallographic structure reported by Almeida and coworkers as starting model.^[S6] The coordination number, the reduction factor S^2_0 and ΔE_0 were kept fixed during the fitting procedure at both edges. Only the radial distances R_i and the mean square related displacements (MSRD) σ^2_i were allowed to vary. The uncertainty on the reported values of distances was $\pm 0.02 \text{ \AA}$ and on the MSRDs was $\pm 0.002 \text{ \AA}^2$.

Elemental analysis (ICP): The composition of the synthesized FeWO₄ sample was determined by Inductively Coupled Plasma Spectroscopy (ICP) analysis, using an Agilent 5100 ICP-OES spectrometer. The sample solution was prepared by dissolving a few mg of the iron tungstate powder in concentrated hydrochloric acid (35%). The obtained yellow solution was then diluted by adding nanopure water, so that the final HCl and FeWO₄ sample concentrations were 5% and ca. 200 ppm. As tungsten has a tendency to reprecipitate with time in such a medium (by hydrolysis, as WO₃), the ICP analysis was performed immediately after the preparation of the solution. Each measurement consisted in the analysis of three different wavelengths per element (Fe and W) and the measurements were replicated three times. The obtained concentration values are presented in **Table S3**.

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