Introduction

Energy today is strongly related to the economic growth of countries around the world and plays an important role in the improvement of living standards, healthcare and educational services, digital information connections and in general an increased productivity.

Although it is difficult to predict the world’s energy consumption in a precise way in the future, it is a fact that this will significantly increase in the coming decades due to the constant growth of the human population and economies worldwide, which in turn demands the consumption of diverse natural resources to satisfy human basic needs.

Such is the case of fossil fuels, mainly used to obtain electricity, transport and artificial air conditioning of buildings. It is due to the decrease in reserves of these fuels and the environmental impacts that these generate that the scientific community around the world are searching for alternative energy sources and sustainable raw materials.1

In recent times, hydrogen (H2) has been of great interest, since it is considered as raw material for a wide variety of processes. For example, with nitrogen in the synthesis of ammonia, with CO and CO2 to produce methanol, in the manufacture of pharmaceuticals, production of hydrogen peroxide, in the electronics and petrochemical industries and to produce numerous chemical products in various processes.1,3

Moreover, hydrogen can be considered as a clean energy carrier, like electricity, which can be produced from various local resources such as renewable and nuclear energy. In the long-term, hydrogen is expected to reduce the dependence on fossil fuels and consequently the emission of greenhouse gases and other pollutants. Currently, hydrogen production accounts for around 2% of the primary energy demand.4 One of the main disadvantages of the use of hydrogen as an energy carrier deals with its relatively low volumetric energy density (8 MJ L−1), even though this presents a high gravimetric energy density (120 MJ L−1).
This low value in volumetric energy density makes very difficult to efficiently transport hydrogen because it needs to be either compressed or liquified, consequently employing large amounts of energy in the process. Therefore, it is necessary to develop better technologies, such as transportation and storage, to make it a more feasible energy vector.

The volumetric storage density (VSD) is a common way to measure the storage capacity of hydrogen in this field and it consists in the weight of hydrogen stored per unit volume of the storage media (g L⁻¹). Typically, the VSD for the liquid hydrogen is 70 g L⁻¹. However, a considerable amount of energy must be consumed to maintain the high pressure and low hydrogen liquefaction temperature.

Nowadays, various physical and chemical methods for hydrogen storage have been proposed. Such as high-pressure and cryogenic-liquid storage, adsorptive storage on high-surface-area adsorbents, chemical storage in metal hydrides and complex hydrides, storage in boranes, carbon materials, and metal organic frameworks. However, each storage method has its advantages and difficulties due to safety, size, weight, cost and efficiency requirements.

An innovative alternative to the actual hydrogen storage technologies has been proposed in recent years, which deals with the chemical looping process (CL). This CL offers many advantages over current technologies such as high efficiency, economic and ecological materials, and proven technical feasibility. This technique employs oxygen carrier materials with very high VSD values (≥ 200 g L⁻¹) that are exposed to a redox cycle. This CL process for hydrogen storage use carrier materials that once reduced by hydrogen convert to metals. These metals are then, in an oxidation step, reacted with steam to release hydrogen as a product of this reaction.

There are a large number of materials that have been evaluated under the CL concept for many thermochemical applications. However, there are a unique number of features that are required for an oxygen carrier to be successfully applied in the area of CL and specifically to CL hydrogen storage. First, the material must retain activity after a fair number of redox cycles, while avoiding deactivation by sintering (thermal stability), possess a high VSD and hydrogen storage capacity, avoid the use of toxic and harmful (to the environment) materials in its composition, to be constituted of abundant and non-expensive materials and finally, to present reasonable reduction and oxidation kinetics.

The use of redox reactions with metal oxides (MeO) under a CL reaction scheme for hydrogen storage is based on reactions (1) and (2). The use of redox reactions with metal oxides (MeO) under a CL reaction scheme for hydrogen storage is based on reactions (1) and (2).

\[ \text{Me}_x\text{O}_y + y\text{H}_2 \rightarrow x\text{Me} + y\text{H}_2\text{O} \]  \hspace{1cm} (1)

\[ x\text{Me} + y\text{H}_2\text{O} \rightarrow x\text{Me}_x\text{O}_y + y\text{H}_2 \]  \hspace{1cm} (2)

In order to be used in a CL process, MeO species must be thermally stable to withstand temperature gradients which are subjected during the process, to be able to store and release lattice oxygen at reaction conditions, and have good availability and affordable costs. The most common MeO reported is iron oxide in its different oxidation states (Fe₂O₄ and Fe₃O₄) and this process is commonly known as the steam-iron process. This process consists in the reduction of iron oxide with H₂ for storage purposes, and the subsequent release of H₂ when oxidizing Fe with steam. The theoretical maximum storage capacity of H₂ in this process is 4.8% w of Fe, i.e., based on the Fe as the only solid reactant. It is also reported a VSD of 374 g L⁻¹ for iron oxides, which is considered a high value. However, most of the oxygen carriers proposed in various research studies employ the impregnation and/or support on different materials of iron oxide to improve the thermal stability and to avoid the material sintering. This causes the generation of an additional inert load to the reactors, which in turn leads to the reduction of up to 3 times the storage capacity of H₂ based on the total solids in the reactive species (including Fe).

In other studies, related to the CL, it has been reported that some mixed metal oxides used as oxygen carriers, such as perovskites and ilmenites (ABO₃), are resistant to high temperatures. These investigations attribute the thermal resistance of the material to the cation A, while high valence B cations would contribute to the catalytic action. An example of these materials is ilmenite (FeTiO₃) that is reported by Schwebel et al. as an oxygen carrier for H₂ and CO as reducing gases. These materials were reported without the addition of extra inert species (supports) for their operation, a feature that can be of advantage towards the CL storage of H₂.

One strategy to improve the sintering resistance of the material has been to include refractory materials in combination with iron oxides such as Al₂O₃, MgAlO₄ and SiO₂. However, the introduction of an inert material has an important impact on the reduction of the VSD of the resulting oxygen carrier. In addition, many studies have focused in the increase of the oxygen-carrier reduction kinetics with hydrogen such as Otsuka et al. that added Rh and/or Ir to Fe/Fe₃O₄ resulting in a small VSD of 68 g L⁻¹ of this material. Furthermore, Takenaka et al. employed a co-doped Cu–Cr–FeOₓ sample showing a VSD of 65 g L⁻¹ at 980 K. In these studies, the addition of Rh, Ir, Pd, Cr can significantly improve the activity and kinetics of iron oxides. However, these not only added high-cost materials to the oxygen carriers, but also presented reduced VSD values compared to plain iron oxide (374 g L⁻¹) thus restraining their commercial utilization and causing researches to focus their investigations on earth-abundant materials. That is the case of oxygen carriers such as Co₀.₂₅Fe₂.₇₅O₄, Zn₀.₂₅Fe₂.₇₅O₄, Cu₀.₂₅Fe₂.₇₅O₄, Mn₀.₂₅Fe₂.₇₅O₄ and Ni₀.₂₅Fe₂.₇₅O₄ for CL hydrogen storage as reported by Xiao et al. Their results showed that Co₀.₂₅Fe₂.₇₅O₄ exhibited an acceptable VSD of 62.47 g L⁻¹ and average hydrogen production rate of 132 μmol g⁻¹ min⁻¹ with a fair performance after 10 redox cycles at 550 °C.

Recently, it has been reported that the oxygen diffusion process is one of the main rate-limiting steps towards the oxidation-reduction reactions within the oxygen carrier material under a CL reaction scheme. Li et al. studied the partial oxidation of methane employing a transient pulse injection experimental reactor. They found that the rate-limiting step was the bulk oxygen diffusion determined by the surface O coverage over the oxygen carrier, while the methane activation...
model oxygen carriers with the same time improve their oxygen-ion conductivity to enhance the sintering resistance of iron oxides and at the same time improve the redox stability over repetitive redox cycles. While, Xiao et al. achieved a high redox stability over repetitive redox cycles.

Moreover, tungstate materials such as NiWO$_4$ and CoWO$_4$ have been reported for several single crystal M$_2$WO$_4$ (M$_2$ = Mn, Co, Ni, Cu) transition metal tungstates. Furthermore, high temperature conductivity measurements have been performed for several single crystal M$^{2+}$WO$_4$ (M$^{2+}$ = Mn, Co, Ni, Cu) transition metal tungstates. Therefore, in principle metal tungstates such as CoWO$_4$ and NiWO$_4$ can be selected in this research to be synthesized, characterized (through XRD, SEM and BET), evaluated on its hydrogen storage capacity by thermogravimetric analysis (TGA), studied to establish its reduction and oxidation global kinetics and thermal stability by a multicycle redox test. All of these to determine its feasibility as a hydrogen storage oxygen carrier under a CL reaction scheme.

### Methodology

#### Synthesis

The NiWO$_4$ was synthesized by the chemical precipitation method using solutions at room temperature under constant stirring, as reported by Song et al. Briefly, 100 mL of 0.7 M precursors salts Na$_2$WO$_4$·2H$_2$O (Sigma Aldrich sodium tungstate dihydrate, white crystals, 99.9% w pure, melting point at 698 °C, and relative density of 4.18 g cm$^{-3}$) and Ni(NO$_3$)$_2$·6H$_2$O (Sigma Aldrich nickel(II) nitrate hexahydrate, emerald green crystals, 99.9% w pure, melting point at 56.7 °C, and relative density of 2.05 g cm$^{-3}$) solutions under strongly magnetic stirring at room temperature were mixed together drop by drop using a peristaltic pump. Once the precipitate was obtained, it was filtered and washed repeatedly with deionized water and then dried at 100 °C for 2 hours. The resulting solid was pulverized in an agate mortar. This powder was then calcined at 950 °C for 5 hours in a ceramic crucible and allowed to cool at room temperature in a desiccator. The powder was vialled for further characterization and tests.

#### Thermodynamic analysis

NiWO$_4$ reduction with hydrogen has been previously reported and its thermodynamics. However, regeneration...
conditions using H\textsubscript{2}O have not been attempted. Therefore, a thermodynamic sensitivity analysis using a Gibbs reactor in Aspen Plus software was carried out. This Gibbs reactor employs the Gibbs free energy minimization technique to obtain the equilibrium compositions for each gaseous and solid species within each reaction system. This technique is fully described elsewhere.\textsuperscript{21} The thermodynamic reaction system composed by the following gaseous species: H\textsubscript{2}(g) and H\textsubscript{2}O(g). The solid species included in the system were NiWO\textsubscript{4}, NiO, WO\textsubscript{2}, WO\textsubscript{3}, Ni and W. The feed stream to the reactor was fixed to 1 kmol h\textsuperscript{-1} of each Ni and W, while the amount of steam feed was varied from 3 to 14 kmol h\textsuperscript{-1}. In addition, the reactor temperature was varied from 600 to 950 °C. This thermodynamic analysis will provide experimental conditions where fully regeneration of NiWO\textsubscript{4} can be expected.

**Hydrogen storage performance evaluation**

The redox behavior of the NiWO\textsubscript{4} powder was followed by a TA-Instruments TGA model Q500, following the weight change (% w) signal with respect to time. All redox experiments were carried out at atmospheric pressure, the total gas flowrate through the sample was 100 mL min\textsuperscript{-1} and the amount of NiWO\textsubscript{4} was of ca. 20 mg placed in a platinum-made sample holder.

A first redox cycle experiment was performed at a constant temperature of 800 °C using 5% v H\textsubscript{2}/Ar and 2.2% v H\textsubscript{2}O/Ar for reduction and reoxidation, respectively, aiming to evaluate the initial NiWO\textsubscript{4} hydrogen storage by comparing the degree of reduction and oxidation in a complete redox cycle. This test then was compared with results of other oxygen carriers previously reported in the literature.

**Characterization**

Characterization of the calcined sample was examined to study its crystalline structure on a Panalytical X’Pert PRO powder diffractometer using Cu-K\textsubscript{α} radiation generated at 40 kV and 30 mA. BET specific surface area was examined by N\textsubscript{2} adsorption–desorption isotherms with a NOVA 4200e surface area and pore size analyzer apparatus, morphology and microanalysis composition by scanning electron microscopy (SEM), and energy dispersive X-ray spectroscopy (EDS) were examined on a JEOL JSM-7401F field emission scanning electron microscope.

The XRD, SEM and EDS analyses were performed on the fresh synthesized sample, after the first thermogravimetric analysis (TGA) reduction (R), after one TGA redox cycle (RO), after a second TGA reduction (ROR) and after 17 TGA redox cycles. This was, in order to characterize the material during its exposure to different steps of reduction and oxidation cycles, as well as to evaluate the stability of NiWO\textsubscript{4} to determine the chemical species present in each stage and establish their reaction paths.

**Global kinetic study**

In order to evaluate the reduction and oxidation kinetics, different temperatures and gas concentrations were used. Table 1 shows 18 experimental conditions that were employed for the global kinetic study.

Prior to each TGA test, the sample was heated in an inert atmosphere (Ar) to reach the targeted experimental temperature (730, 800 or 870 °C) and then followed by an isothermal step under a reducing (H\textsubscript{2}/Ar) or oxidizing (H\textsubscript{2}O/Ar) atmosphere. Before switching from reduction to oxidation atmospheres and vice versa, the remaining reactive gases were removed by an argon flush flow for 5 min.

For the reduction step, different H\textsubscript{2}/Ar flowrates (2, 3.4 and 5% v) were used. The duration of this step was determined in order to achieve a complete reduction of NiWO\textsubscript{4} to Ni and W, i.e., 20.9% of theoretical weight loss.

For the reoxidation of the sample, a mixture of steam and argon was supplied by a water temperature-controlled saturator using an Ar flow, with concentrations of 0.8, 1.5 and 2.2% v H\textsubscript{2}O/Ar. The duration of this step was established in order to complete the full reoxidation of Ni and W back to NiWO\textsubscript{4} (until no mass change could be detected).

Based on TGA results according to experimental conditions of Table 1, global kinetic parameters such as initial reaction rates, reaction constants and orders, as well as activation energies for the reduction and oxidation steps were calculated.

**Hydrogen storage multicycle stability test**

It is important to point out the importance of the thermal stability of the materials used to storage hydrogen in CL operation, because the materials employed in this process are expected to be exposed to multiple (redox cycles) absorption and release of hydrogen. Therefore, a multi-cycle redox test was performed with the purpose of testing the thermal stability of NiWO\textsubscript{4}. This was evaluated by exposing the sample to seventeen consecutive redox cycles and comparing the weight change progress of each cycle as function of time at a constant temperature of 800 °C using 5% v H\textsubscript{2}/Ar and 2.2% v H\textsubscript{2}O/Ar as reducing and oxidizing gases, respectively.

**Results and discussion**

**Thermodynamic analysis**

Fig. 1 shows the sensitivity analysis of NiWO\textsubscript{4} formation as function of reactor temperature and H\textsubscript{2}O molar feed in combination with 1 kmol h\textsuperscript{-1} of W and Ni each to a Gibbs reactor using Aspen Plus process simulator. In this figure, it is possible to observe the conditions for full regeneration to NiWO\textsubscript{4}, which according to the results is a window with
temperatures from 600 to 870 °C and H₂O molar flow rates from 6.5 to 13.5 kmol h⁻¹, respectively.

**Hydrogen storage performance evaluation**

Fig. 2 shows a synthesized NiWO₄ sample evaluated for a hydrogen storage performance at different stages of reduction and oxidation using TGA at 800 °C. In this figure theoretical weight changes of NiWO₄ and reduced Ni + W species are represented by horizontal lines corresponding to 100% w (orange) and 79.1% w (green), respectively. Fig. 2(a) starts with the fresh synthesized NiWO₄ sample corresponding to a 100% w, while Fig. 2(b) shows the first reduction of this material, which achieved a complete reduction using 5% v H₂ (R-sample), since this reached the theoretical Ni + W% w. After this, the sample was oxidized using 2.2% v H₂O/Ar. Fig. 2(c) shows the sample weight change at an intermediate oxidation stage (TGA, 93% w) between the fully reduced and oxidized steps. Furthermore, Fig. 2(d) illustrates the sample reaching the total % w corresponding to the complete theoretical oxidation to NiWO₄, using the same oxidation conditions that started at point 2(b) (RO-sample). In addition, Fig. 2(e) indicates that the reduced sample reached again the fully reduced species (Ni + W), corresponding to the second cycle performed at the same reducing conditions previously employed (ROR-sample). Finally, Fig. 2(f) identifies the final oxidation step after 17 consecutive redox cycles, reaching again the % w corresponding to the fully NiWO₄ oxidized species.

**Characterization**

**XRD.** Fig. 3(a) shows the XRD pattern of the synthesized material. This diffractogram corresponds to the crystallographic phase of nickel tungstate (NiWO₄) in agreement with the ICSD collection code 015852. Furthermore, Fig. 3(b) presents the diffractogram obtained from reduced NiWO₄ under 5% v H₂/Ar atmosphere in a TGA (Fig. 2(b)). Here it is clearly observed that Ni and W are the only metallic species being present, which indicate a complete reduction of nickel tungstate. These results are consistent with studies reported elsewhere. On the other hand, Fig. 3(c) shows the diffractogram of the intermediate TGA weight change (93% w) between the reduced sample and oxidation using 2.2% v H₂O/Ar corresponding to Fig. 2(c). In this diffractogram it is evident the presence of WO₂ as an intermediate specie along with Ni and NiWO₄ in the process towards the total reoxidation to NiWO₄. At this stage, it is important to notice that metallic Ni is not thermodynamically possible to be oxidized with steam towards NiO. Furthermore, Fig. 3(d) shows the XRD pattern of the sample after being exposed to seventeen redox cycles in a TGA (Fig. 2(f)). According to this diffractogram (Fig. 3(d)), the only crystallographic phase obtained agreed with the nickel tungstate
structure, ICSD collection code 015852, which corresponds to the same crystallographic phase found at the beginning of the redox cycles (fresh sample).

Based on results from XRD diffractograms Fig. 3(c) and (d) it is proposed that the reaction path that explains these results are as follows:

\[
\begin{align*}
W + 2H_2O &\rightarrow WO_2 + 2H_2 \\
Ni + WO_2 + 2H_2O &\rightarrow NiO \cdot WO_3 + 2H_2
\end{align*}
\] (5) (6)

First, W is oxidized to WO\(_2\) according to reaction (5) and secondly, Ni and WO\(_3\) are oxidized by steam according to the gas–solid reaction (6). It is important to note that previously formed tungsten dioxide (WO\(_3\)) reacts along with Ni through a gas–solid reaction with H\(_2\)O, this last providing the required oxygen for the formation of NiWO\(_4\). Furthermore, one way to explain these results can be based on the fact that presumably the inclusion of Ni in the crystal lattice of tungsten dioxide (WO\(_3\)) inhibits the formation of the WO\(_3\) phase during its oxidation with water vapor. This simultaneously favors the formation of nickel tungstate (NiO\(^\cdot\)WO\(_3\) or NiWO\(_4\)) phase when is further oxidized, which is consequently the final phase found according to the X-ray diffraction results reported in Fig. 3(d). The latter is because it is thermodynamically impossible to oxidize Ni with steam as indicated in the thermodynamic analysis section. Therefore, the only source of oxygen capable of oxidizing W + Ni must be through a reaction that involved the formation of WO\(_2\) as an intermediate specie, which is consistent with results from XRD characterization leading to the overall regeneration reaction (4).

**BET surface area.** Moreover, BET analysis of the fresh NiWO\(_4\) sample indicates a surface area of 4.25 m\(^2\) g\(^{-1}\), which agrees with results found by Quintana et al.\(^{24}\) who reported a range of BET surface areas between 16 to 3 m\(^2\) g\(^{-1}\) produced by the precipitation method and calcined at 400 and 800 °C, respectively. Due to the small amount of sample employed during the TGA tests (20 mg) it was not possible to perform BET analysis of other samples produced during the TGA test.

**SEM and EDS analysis.** Fig. 4 presents SEM images of the NiWO\(_4\) sample at different steps described in the TGA test presented in Fig. 2. In Fig. 4(a) it can be observed that fresh NiWO\(_4\) sample is composed of non-porous (rock-like) large particles of polygonal morphologies with sizes varying between 1 and 5 μm and similar to those reported by Quintana et al.\(^{24}\). Fig. 4(b1 and b2) shows the same sample after the first reduction according to TGA Fig. 2(b) (R-sample). In this image it can be observed how the initial particle changed its surface morphology from a non-porous rock-like regular shape to a porous coral-like morphology formed by a network of particle agglomerates of about 50 nm size (see ESI†) with cavities between 50 to 100 nm size. These morphologies have been observed previously by Tatsumi et al.\(^{35}\) for NiO–NiFe\(_2\)O\(_4\) and Ahmed et al.\(^{36}\) for NiWO\(_4\) calcined in a temperature range from 750 to 900 °C. According to these authors and Kang and Jeong,\(^{33}\) the reduction process is as follows:

\[
\begin{align*}
\text{NiWO}_4 + H_2 &\rightarrow Ni + WO_3 + H_2O \\
\text{WO}_3 + H_2 &\rightarrow WO_2 + H_2O \\
\text{WO}_2 + H_2 &\rightarrow W + H_2O
\end{align*}
\] (7) (8) (9)

However, no attempt has been made to explain the formation of the resulting solid porous morphology after the reduction process of NiWO\(_4\) with H\(_2\).

Fig. 5 shows a representation of a proposed reduction process to explain the obtained morphology. First, it starts with a solid non-porous NiWO\(_4\) particle, which according to reaction (7) leads to the formation of Ni particles surrounding and enclosing WO\(_3\) particles (1st stage). These WO\(_3\) species enclosed by Ni particles can be explained in terms of results from a research conducted by Inomata et al.\(^{27}\) who showed that the solid-state diffusion between W and Ni in a similar range of temperatures (as used in the present research) is mainly controlled by boundary and volume diffusion. According to Fig. 5, Ni formed in the first stage prevents shrinkage of the resulting material, whilst W formed species begin to diffuse through Ni and the formation of inner particles of W species surrounded by Ni particles take place. They called this phenomenon a diffusion induced recrystallization (DIF) that predominantly produce W species into Ni by volume diffusion, which is enhanced by the high diffusivity of W in Ni (\(D = 9.2 \times 10^{-18} \text{ m}^2 \text{ s}^{-1} \) at \(T = 800^\circ\) C) where initial pores are formed due to the reduction of NiO to Ni. However, WO\(_3\) is still not reduced and the size of the original NiWO\(_4\) particles can be maintained during this first reduction reaction. In the second reduction step (2nd stage), WO\(_2\) is reduced to WO\(_2\), with the consequent release of gaseous water according to reaction (8). This gas generation causes pores to form because of its escape from the inner to the outer solid structure of the material. Finally, WO\(_2\) is further reduced by H\(_2\) (3rd stage) according to reaction (9), thus generating additional porosity due to the release of H\(_2\)O according to this reaction and forming species Ni + W as overall reduction products. This generating a coral-like morphology and a network of porous structure, which will be crucial for explaining the fast steam-oxidation kinetics observed in the kinetic section of this paper.

Fig. 4(c) shows an SEM image of the sample after one complete redox cycle (RO-sample) using 5% v H\(_2\)/Ar and 2.2% v H\(_2\)O/Ar during reduction and oxidation steps, respectively. Here it can be observed a different morphology compared to the fresh sample of Fig. 4(a), in this image, clusters of particle agglomerates are observed and this can be explained in terms of the high temperature exposure of the sample (800 °C). However, particle sizes were reduced from 1–5 μm from the fresh sample to 0.2–1 μm, this caused by the oxidation of the porous network of particles, thus producing smaller particles forming clusters of agglomerates.

On the other hand, Fig. 4(d) shows the SEM image of the sample after a second reduction (ROR-sample, spot d in TGA test) where it can be observed that a similar structure to Fig. 4(b1 and b2) was produced, but containing larger porous
within the particles. This behavior can be explained because, unlike in the first reduction, here the reduction starts with smaller initial particles. This is consistent from this cycle onwards, which is confirmed by the morphology of the oxidized sample after 17 redox cycles shown in Fig. 4(e).

In summary, after the first cycle, the most significant change in the sample was its morphology that suffered important changes as shown in Fig. 4(b1 and b2) (first reduction SEM images) and Fig. 4(c) (first cycle oxidation). The morphology of the oxidized sample after the first cycle is clearly not the same as the fresh NiWO₄ sample (Fig. 4(a)). However, the morphology of the oxidized sample at the beginning of the second cycle (Fig. 4(c)) and after the reduction during the second cycle (Fig. 4(d)) is very alike to the morphology of the sample after
seventeen cycles (Fig. 4(e)). The reason why a comparison in terms of characterization results was made between the second and after seventeen cycles obeys to the fact that after the second cycle the material is consistent in terms of morphology and this is a key feature in the behavior of the material under cyclic operation. This behavior indicates that after the first cycle the material begins to adapt to the reaction cycling conditions and reaches its reaction and thermal stability, since after this second cycle and beyond (17 consecutive cycles) no significant difference in terms of reactivity, composition and morphology was observed. This was reflected in the corresponding characterization (XRD, SEM and EDS analyses) results obtained after seventeen consecutive cycles, which established the NiWO₄ stability.

Table 2 shows results from EDS analysis of the fresh sample (a), first reduced sample (b), first oxidized sample (d), second reduced sample (e) and the oxidized sample after 17 consecutive redox cycles (f). EDS images also are reported in Fig. S3 of ESL† Elemental composition values from Table 2 of the fresh sample are consistent with the theoretical composition of NiWO₄. On the other hand, the (b) sample, which theoretically should be composed only by Ni and W presents traces of oxygen (1.96% w). This is possibly due to the oxidation that takes place between the sample and the environmental air during vialing and handling this sample to different analyses.

Furthermore, column (d) shows the elemental composition of the sample after being oxidized under steam and completing the first redox cycle. This composition is very similar to the theoretical values of NiWO₄. This result confirms that NiWO₄ was fully regenerated after the oxidation reaction and agrees well with results from the TGA test and XRD analysis.

Likewise, column (e) confirms the elemental composition reached in a second reduction with hydrogen, which again is very close to the theoretical Ni + W composition. While column (f) shows that the sample returns to a composition close to the theoretical NiWO₄ values after completing 17 redox cycles, thus showing the high thermal stability of this material. From this table it can be seen that there is a good agreement between the proposed reaction steps and results from EDS, TGA and XRD analyses previously described.

**Global kinetic study.** From the TGA signal response in Fig. 2, the slope obtained \((\Delta w/\Delta t)\) is proportional to the initial reaction rate \((-r_A)\). Initial reaction rates were evaluated for each one of the 17 tests that were previously described in Table 1.

According to what is widely known, this reaction can be studied by a usual model of a gas–solid reaction rate according to the following expression:²⁸

\[
-r_A = kC_A^n
\]  

where \(k\) is the reaction rate constant, \(n\) is the reaction order with respect to reactant gas and \(C_A\) is the \(H_2\) or \(H_2O\) concentrations in mol fraction for the reduction or oxidation steps, respectively.

![Fig. 6 Reaction order plots for the first and second reduction steps of NiWO₄ respect to H₂ concentration.](image)

Table 2  EDS results of the fresh sample (a), R-sample (b), RO-sample (d), ROR-sample (e) and the sample after 17 consecutive redox cycles (f)

| Element  | (a) | (b) | (d) | (e) | (f) | Theoretical oxidized sample | Theoretical reduced sample |
|----------|-----|-----|-----|-----|-----|-----------------------------|---------------------------|
| O        | 22.49 | 1.96 | 25.22 | 3.19 | 23.00 | 25.21 | 0.00 |
| W        | 59.05 | 70.34 | 55.73 | 70.00 | 57.10 | 56.49 | 75.80 |
| Ni       | 18.46 | 27.70 | 19.05 | 26.81 | 19.90 | 18.30 | 24.20 |
Reduction kinetics. The global reaction order for the NiWO₄ reduction with respect to the hydrogen concentration at experimental conditions previously described is shown in Fig. 6. The logarithm of the reaction rate vs. the logarithm of the hydrogen concentration is plotted in this figure. A linear equation results from the analysis at each temperature condition, where the ordinate axis intersect represents the logarithm of the reaction rate constant ($k$) and the slope ($n$) the reaction order at a specific temperature.

By a careful observation of the TGA response from Fig. 2 during the NiWO₄ reduction, a change of slope is evident, which is related to different reactions that take place in this process. According to and Kang and Jeong, Ahmed and Sidharta the reduction of NiWO₄ kinetics is reported to occur in two steps. The initial step is associated to the reduction of NiO to Ni as indicated in reaction (7) while W species remain in its oxidation state as WO₃. During the second step of the kinetics, WO₃ is reduced to WO₂ and this is further reduced to W according to reactions (8) and (9). Therefore, a kinetic evaluation of these two steps was made in order to determine global kinetic parameters (reaction order, reaction constants, and activation energies).

According to results from Fig. 6, for the first reduction step (step 1), the reaction order varied between 1.05 and 1.11. Therefore, it can be concluded that the global reduction reaction of NiWO₄ to Ni + WO₂ is first order with respect to the hydrogen concentration. Furthermore, the second step reduction from Ni + WO₂ to Ni + W presents a reaction order that varied from 0.97 to 1.02, which represents also a global first order reaction with respect to the H₂ concentration. By using the $k$ values generated in Fig. 6, Arrhenius plots for each reduction step can be calculated and these are presented in Fig. 7. From these plots, it can be observed that activation energies for the reduction of NiWO₄ during step 1 and 2 are 22.11 and 43.41 kJ mol⁻¹, respectively. These values agree well with those found by Ahmed et al., who reported activation energies of 18 and 38 kJ mol⁻¹ for these two steps, respectively. It is important to indicate that most of the previous kinetic studies performed of the reduction of NiWO₄ do not report the influence of the H₂ concentration as the present study does. Li et al. recently reported several ferrite-spinel materials for hydrogen storage using hydrogen and steam during reduction and oxidation, respectively. They found hydrogen reduction global activation energies in a temperature range from 450 to 850 °C for Fe₃O₄ ($E_A = 98.63$ kJ mol⁻¹), Zn₀.₂₅Fe₂.₇₅O₄ ($E_A = 86.67$ kJ mol⁻¹), Mn₀.₂₅Fe₂.₇₅O₄ ($E_A = 76.31$ kJ mol⁻¹), Cu₀.₂₅Fe₂.₇₅O₄ ($E_A = 52.05$ kJ mol⁻¹), Ni₀.₂₅Fe₂.₇₅O₄ ($E_A = 45.36$ kJ mol⁻¹) and Co₀.₂₅Fe₂.₇₅O₄ ($E_A = 35.36$ kJ mol⁻¹). By comparing these values with the global activation energies for steps 1 and 2 of NiWO₄ reduction with hydrogen, it can be seen that lower activation energies are needed for nickel tungstate than those recently reported in the literature for mixed oxides under similar conditions, which confirms the potential of NiWO₄ in hydrogen storage applications.

Oxidation kinetics. According to results from Fig. 2, it can be seen that the oxidation of the reduced metallic species (Ni + W) occurs in a fast single straight line that covers almost the entire oxidation process (ca. 99% w). The remaining 1% w can be attributed to diffusional limitations of H₂O molecules to reach...
a small portion of the unreacted reduced species. Therefore, the kinetic analysis of the oxidation step was made by taking into account the main oxidation reaction process (reaction (4)).

Fig. 8, presents results of a plot of ln[H2O] vs. ln(−rA) for the three oxidation temperatures of 730, 800 and 870 °C, where the slope of each line represents the reaction order and the intercept with the y axis the reaction constant at each temperature. According to results from this figure, the reaction order varied from 0.99 to 1.08, which represents a global reaction order of one with respect to the H2O concentration, whereas the reaction constants were 0.053, 0.074 and 0.102 s−1 for 730, 800 and 870 °C, respectively.

Furthermore, Fig. 9 presents the Arrhenius plot for the oxidation step. In the case of steam oxidation reaction, the global activation energy obtained was 53.4 kJ mol−1. Here, it is important to indicate that according to XRD results of the oxidation process presented previously, the mechanism identified for the oxidation of Ni + W back to NiWO4 starts with W being oxidized by steam to form WO2 and H2 (reaction (5)) and in a second step Ni, WO2 and H2O simultaneously react to form NiWO4 + H2 (reaction (6)). This means that during the oxidation process WO3 was not formed as an intermediate specie. This behavior is consistent with previous studies performed by Wendel et al. and Smolik et al. that described the oxidation of W with H2O as follows:

\[
\begin{align*}
W + 2H_2O &\rightarrow WO_2(s) + 2H_2(g) \\
WO_2(s) + 2H_2O &\rightarrow WO_3H_2O(g) + H_2(g)
\end{align*}
\] (11)

They have found that the formation of gaseous specie WO3H2O leads to the sublimation of W an ultimately to the loss of W under steam atmospheres in a temperature range between 200 to 1100 °C. Furthermore, Sabourin and Yetter performed the oxidation of W with a mixture of steam and H2 in order to inhibit the formation of W volatile species an consequently the loss of tungsten.

The facts that all W was reacted with Ni and H2O to form NiWO4 and that no loss of W was found during the TGA tests corroborates that WO3 is not an intermediate during the oxidation step and that the formation on NiWO4 prevents the volatilization of W in the present research. Furthermore, activation energy for the oxidation of W with steam has been reported to be 129 and 139 kJ mol−1 according to Smolik et al., which is 2.6 times greater than the activation energy reported in Fig. 9, this is presumably attributed to the presence of Ni during the oxidation process and to the unique coral-like morphology of W + Ni that presents an interparticle porous network as seen in SEM Fig. 4(b2). This particular morphology presumably eases the diffusion of reactant gaseous H2O, thus increasing Ni + W oxidation kinetics. A further modeling kinetic study would be needed in order to determine the intrinsic activation energy and possible mass transport and/or diffusional limitations during the oxidation step, however, this study is out of the scope of the present investigation.

Finally, according to results obtained in this research, the global kinetic expressions for the reduction [steps 1 and 2] and oxidation reactions are respectively as follows:

\[
\begin{align*}
-r_1[NiWO_4] &\rightarrow 2.98e^{-\frac{22.6}{RT}} \text{H}_2 \\
-r_2[NiWO_4] &\rightarrow 7.83e^{-\frac{43.4}{RT}} \text{H}_2 \\
-r_r[Ni + W] &\rightarrow 6.51e^{-\frac{33.4}{RT}} \text{H}_2O
\end{align*}
\] (13)

**Hydrogen storage multicycle stability test.** A multicycle thermal stability test was carried out by TGA and this is presented in Fig. 10. Weight loss by reduction and weight gain by oxidation was in average approximately equal to 20.7% w compared with the theoretical 20.9% w, which corresponds to the weight change from NiWO4 to Ni + W species. Results obtained clearly show that the material presents a significant
thermal stability during the seventeen-hydrogen storage-release cycles, where no apparent deactivation is observed by comparing the % w difference between the first and last cycles. Furthermore, XRD and EDS results from Fig. 3(d) and Table 2, respectively, confirm that the sample composition obtained after seventeen consecutive cycles is the same as the initial NiWO$_4$, which demonstrates the high stability of the sample under the exposed conditions.

By comparing the present results to other materials reported in the literature under similar multicycle tests it can be assessed the potential of NiWO$_4$ for hydrogen storage through a CL operating scheme. Lorente et al.\(^4\) studied the effect of added metals Al, Cr and Ce to Fe$_3$O$_4$ in the steam-iron process. They reported an improvement in the thermal stability and oxidation activity of the solids during seven consecutive redox cycles. However, their materials showed a reduction in hydrogen storage capacity between 7 to 16% w according to the type of additive used, while no key information related to the reduction and oxidation kinetics was reported. In another research using a ternary ferrite-spinel (Cu$_{0.5}$Co$_{0.5}$Fe$_2$O$_4$) reported by Zeng et al.,\(^5\) they found that the hydrogen storage performance was maintained with only negligible deactivation being observed after twenty consecutive redox cycles. This behavior compares well with the stability exhibited by other similar hydrogen storage materials containing rare earths. However, their material presents only a theoretical volumetric hydrogen storage density (VSD) of 65.58 g L$^{-1}$ compared to a high VSD of 496 g L$^{-1}$ of NiWO$_4$. Furthermore, Zeng et al.\(^5\) studied a CoFeAlO$_2$ spinel material, which was exposed to twenty CO–H$_2$O redox cycles at 800 °C. They found a significant deactivation of this material. Nevertheless, their reduction lasted for 90 minutes ($t_{\text{red}} = 0.16%$ w min$^{-1}$) and the oxidation was completed after 60 minutes ($t_{\text{oxi}} = 0.25%$ w min$^{-1}$), which corresponds to the kinetics of this material. However, they did not evaluate the global kinetics of this material based on an activation energy. In another recent research, Fe$_2$Mn$_2$O$_4$ was exposed to a redox cycle at 775 °C using 5% v H$_2$/Ar and 2% v H$_2$O/Ar during the reduction and oxidation, respectively.\(^3\) However, the redox kinetics of this material was not studied, but the oxidation lasted for 73 minutes with a $t_{\text{oxi}} = 0.28%$ w min$^{-1}$. Huang et al.\(^3\) studied the NiFe$_2$O$_4$ oxygen carrier under a water-splitting CL reaction scheme. They subjected this material to CO–H$_2$O redox cycles and found that the reduction of NiFe$_2$O$_4$ was 19.61% w at 790 °C, while its reduction time lasted 60 min ($t_{\text{red}} = 0.36%$ w min$^{-1}$); however, oxidation with steam needed 130 min ($t_{\text{oxi}} = 0.15%$ w min$^{-1}$) to be achieved. Comparing TGA results from NiWO$_4$ as a hydrogen storage material with above results reported in the literature, it is possible to observe (Fig. 10, a redox TGA weight change of 20.7% w at 800 °C) that the reduction time lasted 30 minutes ($t_{\text{red}} = 0.69%$ w min$^{-1}$) while oxidation under steam took 20 minutes ($t_{\text{oxi}} = 1.03%$ w min$^{-1}$). Therefore, it can be concluded from the above comparison that NiWO$_4$ exhibits a superior behavior in terms of reducibility, hydrogen storage capacity and reduction–oxidation kinetics. Furthermore, Li et al.\(^5\) recently reported multicycle stability tests for ferrite-spinel materials under hydrogen reduction and steam oxidation during 10 redox cycles at 550 °C. They found a percentage reduction (%) in hydrogen storage capacity after 10 cycles for Fe$_3$O$_4$ (33%), Zn$_{0.25}$Fe$_{2.75}$O$_4$ (24%), Mn$_{0.25}$Fe$_{2.75}$O$_4$ (15%), Cu$_{0.25}$Fe$_{2.75}$O$_4$ (13%), Ni$_{0.25}$Fe$_{2.75}$O$_4$ (11%) and Co$_{0.25}$Fe$_{2.75}$O$_4$ (2%). Whereas the percentage reduction for NiWO$_4$ after 17 redox cycles (Fig. 10) at a higher temperature of 800 °C was negligible. This significant result confirms the high multicycle redox thermal stability of NiWO$_4$ for hydrogen storage applications.

**Conclusions**

NiWO$_4$ with a volumetric storage density (VSD) of 496 g L$^{-1}$ was studied to evaluate its feasibility as a hydrogen storage material under a CL reaction scheme using hydrogen and steam during its reduction and oxidation steps, respectively. The material was synthesized by precipitation and calcined at 950 °C for 5 hours in air. Characterization consisted in XRD, BET surface area, SEM and EDS analysis. A thermodynamic analysis was performed to explore oxidation conditions of NiWO$_4$ reduced species (Ni + W). NiWO$_4$ hydrogen storage evaluation was performed by TGA using 5% v H$_2$/Ar and 2.2% v H$_2$O/Ar at 800 °C. Global kinetics for the reduction step was studied in a temperature range from 730 to 870 °C using 2 to 5% v of H$_2$/Ar. While oxidation kinetics was examined from 730 to 870 °C using 0.8 to 2.2% v H$_2$O/Ar. Finally, a hydrogen storage multicycle stability test was performed by exposing NiWO$_4$ to twenty consecutive redox cycles. XRD results indicate that NiWO$_4$ was the only crystalline phase obtained of the synthesized material. Fully reduced material consisted of only W and Ni species, while reoxidation led back to NiWO$_4$ specie. Furthermore, XRD results also helped to propose a reaction path for the oxidation of Ni + W species to form NiWO$_4$. BET surface area of the synthesized material was 4.25 m$^2$ g$^{-1}$. SEM results showed that fresh NiWO$_4$ was composed of non-porous (rock-like) large particles of polygonal morphologies with sizes varying between 1 and 5 μm. After the reduction step, the reduced species changed to a porous coral-like morphology formed by a network of particle agglomerates of about 50 nm size with cavities between 50 to 100 nm size. A reduction process was proposed based on SEM images to explain the drastic morphology changes observed during the reduction step in terms of a solid-state diffusion phenomenon between W and Ni, which presumably prevents the shrinkage of the resulting material leading to a porous material. EDS analysis results confirmed the compositions of the reduced (Ni + W) and fully oxidized NiWO$_4$ species, respectively. Global kinetic studies indicate a first order reaction for the first and second steps during reduction of the material ($n = 1$), while a reaction order of $n = 1$ for the oxidation, with activation energies of 22.1, 48.4 and 53.4 kJ mol$^{-1}$ for the two reduction and oxidation steps, respectively. NiWO$_4$ multicycle stability test shown no loss of VSD and fast reduction and oxidation kinetics under the studied conditions after seventeen consecutive redox cycles, which confirms the potential of this material with respect to current oxygen carriers reported in the literature for hydrogen storage applications.

Finally, the proof of concept of NiWO$_4$ as an oxygen carrier employed for hydrogen storage under a CL operating reaction
scheme completed in the present research (synthesis, characterization, TGA evaluation and reaction kinetics), will set the foundations for future studies in this field. These studies should focus on the use of a greater amount of sample along with another experimental configuration such as fixed or fluidized bed reactors. Particle size and attrition resistance (mechanical strength) during a cycling reaction scheme play important roles to evaluate the suitability of this material under CL operation as well as the concentration of the product gases generated in each step and ultimately, a process simulation with another experimental configuration such as fixed or fluidized bed reactors. These studies will be the matter of upcoming endeavors of our research group.

**Author contributions**

P. E. González-Vargas: writing – original draft, writing – review & editing and investigation; J. M. Salinas-Gutiérrez: formal analysis; M. J. Meléndez-Zaragoza: formal analysis; J. C. Pantoja-Espinoza: data curation and supervision; V. Collins-Martínez: investigation, methodology and supervision; A. López-Ortíz: writing – original draft, writing – review & editing, investigation, supervision and project administration.

**Conflicts of interest**

There are no conflicts to declare.

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