Experimental Investigation of the Cycle Stability of different Iron Oxide Composites for a Redox Hydrogen Storage Process

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ABSTRACT – A promising process to store hydrogen is the thermochemical storage based on the repeated reduction and oxidation (redox) of iron oxide or iron. This storage process is an intermittent, two-phase reaction, which takes place under atmospheric pressure in a hydrogen atmosphere during reduction (charging) or in a steam atmosphere during oxidation (discharging). The investigations have been carried out at two constant temperatures, namely 700°C and 800°C. During the storage phase, only iron exists inside the storage reactor - a fact that makes the redox system much safer, compared to hydrogen storage under pressure in a tank.

This work aims at studying the effect of adding different supporting materials upon producing the iron oxide storage composite samples on their thermochemical cycle stability. Furthermore, the influence of the temperature during the initial sintering process and the redox cycles on the reaction behavior of the iron oxide composites is investigated.

It turned out that pure iron oxide pellets have lost about 65% of their redox potential after only three cycles. Applying 10 wt.% of calcium oxide has improved the cycle stability of the iron oxide pellets to over nine cycles. After nine cycles, a loss of redox performance by less than 5% was observed. This has been attributed to the densification of the sample’s outer surface, which is associated with slowing down the gas diffusion rate into/out of the investigated sample. In addition, reducing the temperature during the cycling (800°C to 700°C) and the sintering from 1100°C to 950°C has shown a positive effect on enhancing the cycle stability.

**Keywords:** Hydrogen Storage, Iron/Iron Oxide, Redox Reactions, Cycle Stability

1 INTRODUCTION

To achieve the climate targets of the Paris agreement [1], an increased use of renewable energy sources instead of fossil fuels for energy generation is necessary. However, renewable energies like solar and wind are subjected to strong daytime and weather related fluctuations. To maintain a secure energy supply mainly consisting of renewable energy and to avoid regulation of renewable energy plants, energy storage systems have gained an increased importance. Storage systems can provide a temporal balance between energy production and consumption. Accordingly, the power grid can be relieved and the regulation of renewable energy plants can be prevented. Regarding an energy supply mainly consisting of renewable sources, the demand of sufficient, affordable and safe storage capacities will increase [2,3].

Another key role in the energy transition will be the cross-sector energy supply [3]. Hydrogen, as a central energy carrier for renewable energies, has proven to be an ideal candidate for such sector coupling [4,5]. The excess electricity of renewables can be used to generate hydrogen by spitting water via electrolysis. In times with a high-energy demand, hydrogen can be converted into electricity and heat by a fuel cell or a combustion engine connected to a generator. Alternatively, the stored hydrogen can be utilized as an energy carrier for transport.

In previous research works, different technologies to improve the energy density and safety of hydrogen storages have been investigated and developed [4,7]. These technologies can be classified into two main categories: physical and chemical systems. Physical storages include compressed and liquefied hydrogen as well as adsorption on solid materials. Currently, the most common way is to store hydrogen in a tank after compression or liquefaction. However, about 9% to 12% of the stored energy is needed for the compression of hydrogen to 300 or 700 bar, respectively [8].

The volumetric energy density increases from 780 to 1300 Wh/dm³ upon increasing the storage pressure of hydrogen from 350 to 700 bar. Storing hydrogen in liquid form results in enhancing the storage density to 2300 Wh/dm³. [9] However, the necessary cooling down to -253°C at about 1 bar results in a loss between 20% to 30% of the lower heating value of the stored hydrogen [10]. Additionally, the amount of stored hydrogen decreases through evaporation losses and boil-off effects, which take pace over time [4,11].

In a chemical storage process, in most cases, hydrogen is chemically bound into a liquid or a solid carrier material. Using liquid organic hydrogen carriers (LOHC), unsaturated organic compounds such as N-ethylcarbazole,
naphthalene and toluene can be utilized. At a temperature of typically up to 200°C and a pressure of about 50 bar, the catalytic hydrogenation of the LOHC material takes place (charging phase). During the endothermic dehydrogenation of the hydrogen rich LOHC material, the absorbed hydrogen can be released with the use of a catalyst. For the discarding process, a temperature up to 310°C and a slight overpressure is needed. Positive aspects of LOHC systems are the flexibility upon transporting the liquid storage medium and the achievable storage capacity. A storage density of 5.8 wt.% is given upon using the carrier N-ethylcarbazole (H0-NEC). The necessary purification process of the released hydrogen after the dehydrogenation is still a challenge facing LOHC systems.

A further storage principle is to apply metal hydride (MeH2) systems, which are based on a reversible chemical reaction (Eq. 1). The process depends on the prevailing temperature, pressure and hydrogen concentration. During the exothermic charging process, the absorption of hydrogen (H2) into the existing metal lattice (Me) takes place. In the endothermic discharge (desorption) phase, the bonded hydrogen is released from the metal hydride.

\[ n \text{Me} + m \text{H}_2 \xrightarrow{\text{quench}} \text{Me}_n\text{H}_{2m} \]  

(1)

In particular, metal hydride storage systems prove to be promising due to the volumetric H2 storage capacity and safety. The obtainable storage density depends on the applied material and varies between 1.6-1.8 wt.% (AB5-type) and 13.5 wt.% for Lithium borohydride (LiBH4 + SiO2). Currently, one problem is the increasing system temperature due to the heat released during the absorption phase, which leads to a decreased storage capacity, unless the storage tank is additionally cooled.

Another chemical storage method is based on the repeated reduction and oxidation (redox) of metal oxide or metal, respectively. In this case, hydrogen is not chemically bound. Instead, it is used for the reduction of metal oxide (MeO) to pure metal (Me). In the same time, hydrogen is oxidized to steam (H2O). The discharge phase is the reverse process in a steam atmosphere (see Eq. 2).

\[ \text{Me}_n\text{O}_m + m \text{H}_2 \xrightarrow{\text{steam}} n \text{Me} + m \text{H}_2\text{O} \]  

(2)

A challenge of this process is the high temperature in a range of about 500°C to 1000°C, depending on the used material and the charging / discharging rate. However, unlike LOHC and metal hydride systems, the most heat demand is required during the charging phase, when excess energy is available. Additionally, the released heat during the exothermic discharging phase (oxidation) can be utilized as an output energy to enhance the overall system efficiency. Furthermore, the safety of the storage process represents a clear advantage. This is due to that only metal or metal oxide and no hydrogen is present during the storage phase. Furthermore, a high volumetric storage capacity can be reached. In an energy storage density of 2800 Wh/(dm³ bulk) for an iron based hydrogen systems was reported.

Currently, the kinetics of different solid metal materials such as Ce, Cu, Fe, Mn, Ni and W have been investigated during a repeated redox reaction. Research works in the field of solid oxide fuel cells (SOFC) and oxide ceramic batteries (ROB) as well as chemical looping (CL) processes have confirmed that the use of iron oxide as suitable carrier material is particularly advantageous. This is due to the thermodynamic behavior of the material, the cost-efficient production and the high availability.

\[ \text{Fe}_3\text{O}_4 + 4 \text{H}_2 \rightarrow \text{Fe}_3\text{O}_4 + 4 \text{H}_2 \]  

(3)

\[ \text{Fe}_3\text{O}_4 + 4 \text{CO} \rightarrow 3 \text{Fe} + 4 \text{CO}_2 \]  

(4)

\[ \text{Fe}_3\text{O}_4 + \text{CH}_4 \rightarrow 3 \text{Fe} + 2 \text{H}_2\text{O} + \text{CO}_2 \]  

(5)

\[ 3\text{Fe}_2\text{O}_3 + \text{H}_2 \rightarrow 2\text{Fe}_3\text{O}_4 + \text{H}_2\text{O} \]  

(6)

2 IRON REDOX PROCESS

The use of iron oxide goes back to the early 20th century, where iron oxide was chosen as a solid carrier material for the production of hydrogen. Instead of hydrogen, carbon monoxide (CO) was utilized for the reduction of iron oxide. Based on this method, one of the first large-scale process for the production of high-purity hydrogen was developed, the so-called “steam-iron” process. The temperature range was typically about 550°C to 900°C. Due to the oxidation of iron in a steam atmosphere, hydrogen was produced (see Eq. 3). The reduction of iron oxide takes place in a carbon monoxide (CO) or methane (CH4) atmosphere (see Eq. 4-5), which leads to the emission of carbon dioxide (CO2).

\[ 3\text{Fe} + 4\text{H}_2 \rightarrow \text{Fe}_3\text{O}_4 + 4\text{H}_2 \]  

(3)

\[ \text{Fe}_3\text{O}_4 + 4\text{CO} \rightarrow 3\text{Fe} + 4\text{CO}_2 \]  

(4)

\[ \text{Fe}_3\text{O}_4 + \text{CH}_4 \rightarrow 3\text{Fe} + 2\text{H}_2\text{O} + \text{CO}_2 \]  

(5)

2.1 Iron redox storage process

The iron redox storage process takes places at atmospheric pressure conditions. When hematite (Fe2O3) is used as the initial oxide, it is converted to magnetite in an upstream step (see Eq. 6-4). The stability range of hematite in relation to the prevailing temperatures (600°C to 800°C) and the oxygen partial pressure (<10⁻¹⁴ bar) prevents the formation of hematite again (cf. Fig. 1).
Theoretical energy density for products magnetite, wüstite or iron and finally the des-tice, the formation and growth of nuclei of the reaction
reductions progresses, the reducing gas di ffuses through the pellets. The mass transfer between the flowing gas and Bogdandy, 1967) fusion processes in the solid reaction partners. (Engell

Fe₂O₄ + H₂ ⇌ 3FeO + H₂O

During the charging phase, magnetite (Fe₂O₄) is first reduced to wuestite. In the second step, further reduction of wuestite to pure iron (Fe) takes place. Fig. 2 illustrates the reduction process, in the case of a porous structure. Starting from the gas phase (flow boundary layer), the gas transport takes places through the macro pores to the grain boundary (A). Through micro pores, the transport continues into the interior of the solid particles (B). Subsequently, at the gas/solid interface, oxygen is released from the crystal lattice and reacts with H₂ to steam (H₂O) (C). This leads to the migration of iron cations (Fe⁴⁺⁺) (solid-state diffusion, D). The transport of the generated steam takes place in a reverse order via micro and macro pores (E,F). [48,49]

Figure 1 Equilibrium oxygen partial pressures of H₂O/H₂ mixtures as a function of temperature (Dashed lines) and dissociation pressures of the different iron oxide reactions (Solid lines); Calculated on the basis of [47].

Regarding the equilibrium condition (Eq. 9), the amount of storage or released oxygen (mass change during the redox process) is proportional to the amount of hydrogen or steam. Accordingly, the difference between the sample mass in the oxidized and reduced state represents a measure for the storage capacity.

O₂ + 2H₂ ⇌ 2H₂O

2.2 Aging process

The repeated redox reactions of pure iron oxide samples results in strong aging effects. In particular, the occurrence of aging effects leads to the agglomeration of iron particles and the increasing densification of the sample structure. The formation of a dense outer surface reduces the gas transport into the interior of the sample. This leads to a decrease in the reaction rate and can cause an incomplete conversion to iron or magnetite. [50,52]

N. Menzler et al. [34] have observed aging effects, which took place due to the repeated redox reaction when pure iron/iron oxide sample are used. After 5.5 redox cycles (reduced state), it came out to an agglomerated structure and the formation of an outer dense iron layer. The investigations were carried out at a process temperature of 800°C and a reaction atmosphere of argon (Ar) with additionally ether hydrogen or steam (reduction: Ar with 2% H₂; oxidation: Ar with 2% H₂ and 7% H₂O).

The aging effects have been attributed to the high mobility and sintering tendency of iron cations during the redox process at a temperature range between 600°C and 800°C. Due to these effects, the available storage capacity decreases strongly with the number of cycles using pure iron/iron oxide samples. [33]

Despite the research carried out so far, the low cycle stability and, consequently, the short lifetime represents the biggest challenge of thermochemical hydrogen storage based on the redox process. In order to fulfill the requirements of an effective and stable storage material, a significant reduction of such aging effects seems quite necessary. This work aims, therefore at giving a hand towards enhancing the cycle stability by introducing different supporting materials upon producing the composite samples. Several redox cycles have been measured simultaneously to detect any aging effects. Moreover, microscopic images of the changed structure of the tested composites after cycling have been analyzed in order to explore the reason behind any obtained aging effect.

3 MATERIALS AND METHODS

Different supporting materials were added to an iron oxide powder with a particle size distribution of max. 60μm and a purity of 99.1% Fe₂O₃ (Linox GmBH).
3.1 Selection of supporting materials

A possible mitigation of aging effects, especially agglomeration of iron, has been achieved by adding suitable support materials. This could be proved in research works, which are dealing with the investigation of redox behavior of iron oxide materials for application in Solid Oxide Fuel Cell (SOFC), Rechargeable Oxide Battery (ROB) [32,34] or Chemical Looping (CL) processes [35-37].

In [35], different iron oxide samples with a supporting material amount of 15% and a diameter of about 4.5 mm were investigated at a temperature range of 700°C to 900°C. The comparatively best results were obtained in particular with CaO-rich samples. By comparing the mass change during oxidation and a temperature of 700°C, the reaction behavior of the CaO-rich sample (10%CaO, 5%Al2O3) was about four times faster than that of the sample without CaO (10% SiO2, 5%Al2O3).

In a research work [35] focused on the investigation of iron oxide based oxygen storage materials for a SOFC, numerous support materials were tested with respect to their cycling stability over 20 redox cycles. The most promising results were obtained by adding CaO. In contrast, the use of other materials like TiO2, CeO2, Cr2O3, Mn2O3, MgO and CuO led to a strongly decreasing reaction behavior or even to a mixed crystals formation, which did not allow a repeated reduction of the sample. [35] In further work [35], the addition of pure SiO2 and Al2O3 resulted in a negative influence on the redox reaction behavior. As a reason, the formation of inert mixed crystals (SiFeO3 and FeAl2O4) was reported. By adding a support material consisting of a combination of SiO2 and Al2O3 with CaO, an improvement in the cycling stability of the sample was achieved. This was attributed to a possible stabilizing effect of CaO through the formation of a dispersed structure, which mitigate the sintering effects of iron.

In this research work, CaO based supporting materials were chosen (cf. Tab. 1). These are a CaO powder with a purity of 96.8% and a CaO-rich Portland cement CEM1 42.5R with a clinker content of at least 95%. In addition, a fire cement with an composition of about 78% Al2O3 and 21% CaO was selected. To keep the loss of active storage material as low as possible, the supporting materials were added with an amount of 10 wt. %.

Table 1 Applied supporting materials including the introduced abbreviations, chemical composition and manufacturer.

| Name | Material | Composition |
|------|----------|-------------|
| Z    | CEM1 42.5R | ≥95% cement clinker, Heidelberg Cement |
| ZF   | Secar 80  | ≥78.1% Al2O3, ≤21.4% CaO, Kerneos |
| CaO  | nekafin0  | 96.8% CaO, Kalkfabrik Netstal |
| ZF   | ZF        |                          |
| CaO  | CaO       |                          |
| CaO-02 | CaO        |                          |
| CaO-03 | CaO        |                          |
| Fe2O3 | -         |                          |

3.2 Manufacturing process and operation conditions

The manually produced samples have been sintered in an air atmosphere for a period of six hours. To investigate the influence of the temperature during the sintering process, some samples have been sintered at 950°C, while the others have been sintered at 1100°C. The initial mass of the investigated samples, after sintering, were about 1.4 g (see Tab. 2).

Table 2 List of parameters of produced storage samples with a support material content of 10 wt. %

| Name | Support material | Initial weight (g) | Sinter temp. (°C) | Process temp. (°C) |
|------|------------------|--------------------|------------------|-------------------|
| Z    | Z                | 1.49               | 950              | 800               |
| ZF   | ZF               | 1.36               | 950              | 800               |
| CaO  | CaO              | 1.36               | 950              | 800               |
| CaO-02 | CaO           | 1.38               | 1100             | 800               |
| CaO-03 | CaO           | 1.32               | 950              | 700               |
| Fe2O3 | -                | 1.41               | 950              | 700               |

In Fig. 3 the experimental setup is schematically illustrated. The reaction kinetics of the samples were investigated using a simultaneous thermogravimetric analyzer (STA 449 F3 Jupiter, Netzsch). Regarding the weight measuring, the accuracy of the STA device is ±0.1 µg, while the accuracy for the sample temperature adjustment is about ±0.3°C. To adjust different gas concentrations (H2, N2), mass flow controllers (MFCs) were applied. The accuracy of one MCF is ±0.5% regarding the gas flow in a range of 0 to 250 ml/min. For oxidation, the amount of steam was generated via an evaporator (DV2MK, ADROP) with an accuracy of ±1% of the final value and a resolution of 0.1 g/h. To evacuate the experimental setup after each experiment as a safety precaution, a vacuum pump is used.

Figure 3 Schematic illustration of the setup for the experimental investigation of the reaction kinetics of storage samples [24].

The redox reactions have been carried out at 700°C or 800°C. The volume flow rate of the reacting gas amounts...
4 RESULTS AND DISCUSSION

Regarding the prevailing process conditions, magnetite is the highest stable oxide. In the first redox cycle, the conversion of hematite to magnetite takes place. This leads to a reduction of the total sample mass (Tab. 2) of about 3%, according to Equation [6]. With respect to the process conditions, hematite cannot be generated due to its unstable behavior (see Fig. 1). Thus, the relative mass of 100% represents the complete oxidation to Fe$_2$O$_3$.

4.1 Different supporting Materials

Fig. 4 depicts the reaction kinetics of different samples with the supporting materials (Z, ZF and CaO, cf. Tab. 2) during oxidation (discharging phase) at 800°C. Up to a relative mass of about 90% of the first oxidation cycle (C1), an approximately similar reaction behavior can be observed (Fig. 4a). Regarding the complete conversion to Fe$_2$O$_3$, a mass change of about 24 wt.% was measured. The ZF-sample reaches its maximum in 19 min. In comparison, 6 and 8 minutes more are needed for the oxidation of the CaO- and Z-sample, respectively.

Fig. 4b shows the reaction kinetics during the oxidation of the third and seventh redox cycles, respectively. Despite offering almost the same kinetics, the ZF-sample was mechanically damaged after three cycles. Consequently, no further investigations could be carried out. Concerning both the Z- and CaO-sample, a significant reduction in the reaction rates become obvious in the seventh cycles. After a reaction time of 100 min, a mass change of only 8 wt.% could be achieved using Z as a supporting material. In contrast, the CaO-sample could nearly reach its maximum (97%). In total, the loss of storage capacity is about 67% (Z-sample) and less than 3% (CaO-sample).

Figure 4 Mass change over the reaction time of the prepared samples with different supporting materials (Z: portland cement; ZF: fire cement; CaO: pure calcium oxide) during the oxidation at 800°C. a) first cycle (C1); b) third (C3) or seventh cycle (C7)

The reason for the low mechanical stability of the ZF-sample could be the condensation of vacancies caused by the structural change of the crystal lattice during the redox reaction [45-53]. In Fig. 5, the formation of a cavity between the interior and the rim can be seen, which causes low adhesion of the outer layer.

Figure 5 Microscopic image of an iron oxide sample with 10 wt.% ZF after the three redox cycles.

Regarding the Z- and CaO-sample, the change of the reaction kinetics with the number of redox cycles indicates that sintering effects occurred. Fig. 5 shows the microscopic structure of the outer layers and the sample surface after seven redox cycles. In both cases, a denser outer layer was formed. The densification of the sample structure leads to an impeded gas exchange (H$_2$, H$_2$O) into/out of the interior of the sample, which results in a slowdown of the reduction and oxidation process, respectively [34-52]. In case of the Z-sample, the sharp decreases of the reaction rate and the storage capacity implies the formation of an strongly densified outer layer, which can be seen in Fig. 5a. Furthermore, comparing the sample structure, a higher porosity can be observed by the CaO-sample. This correlates with the observed higher cycle stability.
Based on the obtained results, ZF and Z are not suitable to be applied as a support material in iron redox storage samples. However, compared to Z and ZF, CaO as supporting materiel showed promising results. This might be attributed to the mixed crystal formation of CaO and iron oxide, which leads to a higher stability, despite the significant increase in the required process time [36, 44]. Due to the positive results, further investigations have been carried out only on iron oxide samples with 10 wt. % CaO.

### 4.2 Influence of the temperature during sintering

In order to explore the effect of the temperature during the sintering process on the cycle stability, CaO-samples were sintered at 950°C (CaO-sample, S950) and 1100°C (CaO-01-sample, S1100), respectively. The redox cycles have been carried out at 800°C.

Fig. 7 shows the reaction kinetics of the samples during the first and second oxidation cycle. Compared to the CaO-sample, the first oxidation of the CaO-01-sample is already four times slower ($\Delta t_{\text{C1-S950}} = 25$ min; $\Delta t_{\text{C1-S1100}} = 100$ min). In the second cycle, a further decrease of the reaction rate can be observed. After 37 min, a mass change of only about 9 wt. % was reached, while in the first cycle, about 18 wt. % could be archived. Unlike this, the reaction behavior of the CaO-sample remained constant during the first two cycles.

The observed strong decrease in reaction rate of the sample sintered at 1100°C suggests that the increased temperature during the sintering process leads to a densification of the sample structure. This results in a reduced gas transport through the internal structure and a decreased conversion of iron to iron oxide, leading to a reduced cycle stability and storage capacity [33, 34, 54].

### 4.3 Influence of the process temperature

The influence of the temperature during the redox cycles on the reaction kinetics has been studied. To this aim, CaO-samples were sintered at 950°C and investigated at a constant process temperature of 700°C (CaO-03-sample; ro700) and 800°C (CaO-sample; ro800), respectively. In Fig. 8, the mass changes in the fifth and eighth oxidation cycle are depicted.

Fig. 8 shows the reaction kinetics of the samples during the first and second oxidation cycle. Compared to the CaO-sample, the first oxidation of the CaO-01-sample is already four times slower ($\Delta t_{\text{C1-S950}} = 25$ min; $\Delta t_{\text{C1-S1100}} = 100$ min). In the second cycle, a further decrease of the reaction rate can be observed. After 37 min, a mass change of only about 9 wt. % was reached, while in the first cycle, about 18 wt. % could be archived. Unlike this, the reaction behavior of the CaO-sample remained constant during the first two cycles.

The observed strong decrease in reaction rate of the sample sintered at 1100°C suggests that the increased process temperature causes a strong decrease in the kinetics. At a relative mass of about 80%, a clear decline of the reaction rate can be observed. As a
result, 300 minutes are needed to achieve a mass change of 23 wt.%. In comparison, at 700°C, almost the same mass change was obtained in 34 min. This indicates that a higher process temperature of 800°C during the cycling does stimulate the occurrence of effects.

### 4.4 Pure iron vs. CaO samples

Fig. 9 presents the reaction kinetics of storage samples with only pure iron oxide (Fig. 9a) and 90% iron oxide and 10% CaO as supporting material (Fig. 9b). Compared to the inertial sample weight, the mass loss in the first redox cycle is about 3.5 wt.% (pure iron oxide sample) and 3 wt.% (CaO-03-sample), respectively. This is due to the conversion of hematite to magnetite.

In case of pure iron oxide sample, a mass change of 27 wt.% can be measured in the first redox cycle. However, already in the second reduction cycle a strong decrease of the reaction rate can be observed (cf. $\Delta t_{\text{red}1}=95$ min, $\Delta t_{\text{red}2}=864$ min). Despite the nine times longer reaction time, a mass change of about 23 wt.% can be achieved during the second reduction and 21 wt.% during the second oxidation. In the subsequent third oxidation process, the mass change is further decreased to 9.5 wt.%. This represents already a loss of storage capacity of about 65%. These aging effects emerge mainly from the agglomeration of iron particles and the increasing densification of the sample structure, which includes the formation of a dense outer layer. [50][52]

Regarding the CaO-03-sample, the mass change in the first redox cycle is about 24 wt.%. The lower storage capacity (compared to 27 wt.% of pure iron oxide) is due to the amount of the support material. Over the first three cycles, an almost identical reaction behavior can be observed (cf. Fig. 9b). The reaction time is about 85 min for the reduction and about 20 min for the oxidation.

Through the repeated redox reaction, a comparatively small decrease in the kinetics sets in. In the ninth cycle, a capacity loss of less than 5% is observed. Thereby, the reaction time has slowed down ($\Delta t_{\text{red}9}=95$ min, $\Delta t_{\text{ox}9}=150$ min). This indicates that the addition of 10 wt.% CaO has clearly improved the cycle stability but could not completely prevent the occurrence of aging effects.

### 5 CONCLUSION

In order to enhance the cycle stability of iron oxide samples for a hydrogen redox storage process, three various supporting materials (Portland cement (Z), fire cement (ZF) and calcium oxide (CaO)) were selected. Composite samples incorporating 10 wt.% of each supporting material with 90% of Fe$_2$O$_3$ have been prepared and cycled in a Thermo-Gravimetric Analyzer upon simultaneously measuring the reduction and oxidation kinetics in an atmosphere of H$_2$/N$_2$ during reduction and H$_2$O/N$_2$ during oxidation at different process temperatures. In addition, the influence of the sintering temperature on the cycle sta-
bility of the composites has been investigated. The structure of the investigated composite samples has been analyzed by a microscope to explore the reasons behind the low mechanical stability or the reduced reaction kinetics. The obtained results can be concluded as follows:

1. The cycle stability could be enhanced and the reaction kinetics could be maintained over more cycles by adding supporting materials, compared to the pure iron oxide samples.

2. CaO as a support material showed the best improvement concerning both cycle stability and reaction kinetics.

3. Reducing the sintering temperature from 1100°C to 950°C showed a significant improvement on both cycle stability and reaction kinetics of the investigated samples.

4. Reducing the process temperature from 800°C to 700°C resulted in another significant improvement of the composite stability of the investigated iron oxide composites based on CaO.

5. The reaction slowing down effect associated with the repeated redox cycles of the investigated samples could be attributed to the formation of a dense outer layer, which enhances the diffusion resistance of the reacting gases into/out of the samples.

6. Pure iron oxide samples lost about 23% and 65% of their redox storage capacity during the 2nd and 3rd redox cycles compared to the 1st cycle. The needed cycle time is increased by a factor of 9 between the 1st and 2nd redox cycles.

7. Adding 10 wt.% of CaO as a supporting material to iron oxide has significantly improved the redox cycle stability but could not completely prevent the occurrence of aging effects. Accordingly more developments and investigations are needed to further enhance the cycle stability of the iron oxide composites.

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