Iron oxides as efficient sorbents for CO\textsubscript{2} capture

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**Abstract**
Carbon dioxide capture/release reactions using magnetite, Fe\textsubscript{3}O\textsubscript{4}, and hematite, Fe\textsubscript{2}O\textsubscript{3}, as sorbents were studied. Kinetics of mechanically activated chemical reactions between iron oxides and CO\textsubscript{2} was investigated as a function of CO\textsubscript{2} pressure and planetary ball mill process parameters. It was found that complete carbonation of iron oxides can be accomplished at room temperature and elevated CO\textsubscript{2} pressure (10–30 bar). Siderite calcination was studied in vacuum and argon atmospheres. FeCO\textsubscript{3} can be decomposed at 367°C yielding magnetite, carbon and/or iron. This mixture can reversibly re-absorb carbon dioxide in multiple carbonation–calcination cycles. These results suggest that siderite or iron oxides are prospective and efficient reversible sorbents for CO\textsubscript{2} capture.

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1. Introduction

The emission of carbon dioxide has been rising for the last several decades as a result of increasing use of fossil fuel by different sectors of industry. According to some recent studies, if the use of fossil fuels continues at this or increasing rate over next 50 years, the CO\textsubscript{2} concentration in the atmosphere could reach 580 ppm [1] causing significant global effects on living organisms and ecosystems. The Kyoto protocol, adopted in 1997, promotes the reduction of emissions of greenhouse gases notably CO\textsubscript{2}, which is the main cause of global warming [2]. This treaty has triggered off the development of new and improved technologies of carbon dioxide capture and sequestration.

Current technologies of carbon dioxide capture and sequestration are based on either physical or chemical methods. In physical methods, carbon dioxide is absorbed under different conditions of pressure and temperature by a solid or liquid and is released by the absorbents on decompression and/or heating. The chemical absorption methods are based on chemical reactions between absorbent and carbon dioxide [3–7]. Amine solutions are the most common absorbents. Several studies have shown that amine absorption is costly and energy intensive. Amine systems to capture 90% of the CO\textsubscript{2} from a flue gas of fossil-fuel power plant, could cost up to 30% of the electricity generated by the plant [8,9].

The adsorption and membranes are two other technologies developed, but like absorption due to the huge amount of
flue gas, they are not considered as suitable solutions [1,10,11] so far. Researchers have largely focused on improving membrane CO₂/N₂ selectivity, often by using facilitated transport mechanism membranes [12].

Another approach to CO₂ capture that is gaining relevance in the recent years focuses on metal oxides such as CaO, MgO and FeO, which can react with CO₂ yielding metal carbonates. Once the metal carbonates are formed they can be heated to release pure CO₂ gas and regenerate the oxides. Normally carbonation reactions are exothermic and carbonate decomposition reactions are endothermic which allow designing a cyclic process [13–15].

Iron oxides are attractive sorbents for iron mills because they can be carbonated and regenerated several times, and after that, be processed for steel production at the same iron making facility. The steel industry consumes large amount of fossil fuels which makes it one of the most energy-intensive industrial sub-sectors accounting for about 7% of total CO₂ emission [16]. The blast furnace gasses (BFG) are CO, N₂, H₂ and CO₂. The main source of carbon is a coke which generates CO₂ emissions between 17% and 25% of total flow of the gases. Nowadays, CO₂ is captured mainly from the BFG by chemical absorption methods using primary amines [17,18] or physical absorption methods like pressure swing absorption, and vacuum pressure swing absorption [14].

This work presents a study on the capture of CO₂ using iron oxides, synthesizing siderite. Normally, synthetic siderite can be produced either at high pressure and high temperature conditions from iron oxides and ferrous oxalate, or by means of chemical methods [19–23].

It was shown that iron oxides have active sites exposed at the surface which can react with the gaseous molecules of CO₂ [24]. Materials treated in grinding processes decrease in the particle size, forming a new fresh surfaces and creating more active sites which translates to increase of the surface area [25,26]. Iron can be a good reducing agent not only due to its thermodynamic properties but also because it is produced on a site where carbon dioxide capture takes place, steel mill [27]. Therefore, we propose the following initial carbonation reactions:

\[
\begin{align*}
\text{Fe}_2\text{O}_3(s) + \text{Fe}(s) + 4\text{CO}_2(g) & \rightarrow 4\text{FeCO}_3(s) \quad (1) \\
\text{Fe}_2\text{O}_3(s) + \text{Fe}(s) + 3\text{CO}_2(g) & \rightarrow 3\text{FeCO}_3(s) \quad (2)
\end{align*}
\]

Reactions (1) and (2) allow producing iron(II) carbonate; reaction (1) was previously studied using a thermo-pressure system with additions of water to improve the reaction kinetics [14]. In nature, carbonate minerals are present of different rocks mainly represented by the quaternary system FeCO₃–MgCO₃–CaCO₃–MnCO₃. Natural samples of iron carbonate show different amounts of substitutions of Mg, Ca, Mn for Fe in the lattice, which suggests that pure siderite seldom occurs [28]. Natural siderite forms complete series of solid solutions with magnesium and manganese carbonates, while a wide miscibility gap reported between iron and calcium carbonates. Solubility of Ca²⁺ in siderite does not exceed 10 mol.% at 550 °C, while calcite (CaCO₃) can contain up to 20 mol.% of Fe²⁺ at the same temperature. Siderite is a commercial mineral and has many applications such as source of iron in steel industry, raw material in cement industry, hydrogen production, refining of ferrosilicon, shale oil production, etc. [29,30].

The thermal decomposition of siderite is very important issue mainly in processing of oil shales or in the combustion of coals [31,32]. Mechanism of the thermal decomposition of siderite is complicated and depends both on its composition and experimental conditions. Formation of solid solutions between FeCO₃ and other metal carbonates increases decomposition temperature of natural siderite samples compared to synthetic one. Temperature, atmosphere, microstructure and heating rate are those experimental parameters which determine phase composition of siderite decarbonation products. In oxygen-rich atmosphere the only end product is hematite, while magnetite, wustite and pure iron could form in vacuum or in an inert atmosphere [28].

There is a remarkable difference in the thermal decomposition behavior of natural and synthetic siderite. For example, Gotor et al., [28] reports that the decomposition temperature of synthetic siderite is approximately 200 K below of the decomposition temperature of the natural sample.

The thermodynamic stability fields of siderite, wustite and magnetite have been studied as a function of P₀₂ and P₀₃ at different temperatures [19,33]. It was found that the pressure range of siderite stability is smaller at higher temperatures, on the other hand, the stability pressure of magnetite and wustite is extended at higher temperatures. Existence of siderite at high temperatures is possible only at high P₀₂ and P₀₃; at those conditions the molar ratio of FeO/Fe₂O₃ increases with temperature [33].

FeO formed by mechanistic chemical reaction between Fe₂O₃ and Fe can be decomposed into Fe₂O₄ and Fe by an endothermic reaction at 250 °C according to DSC study [34]. Results reported in Ref. [29] confirm that siderite decomposition proceeds through formation of FeO and CO₂ and after that FeO undergoes transformation to Fe and Fe₂O₃. Moreover, several studies have reported carbon (C), oxygen (O), carbon monoxide (CO) [19,30,35–37] as by-products of siderite decomposition at different experimental conditions.

In order to obtain a reducing agent after calcination, siderite decomposition was studied in inert atmosphere, hence, the magnetite regeneration reactions are shown below:

\[
\begin{align*}
6\text{FeCO}_3(s) & \rightarrow 2\text{Fe}_3\text{O}_4(s) + \text{C}(s) + 5\text{CO}_2(g) \quad (3) \\
4\text{FeCO}_3(s) & \rightarrow \text{Fe}_3\text{O}_4(s) + \text{Fe}(s) + 4\text{CO}_2(g) \quad (4)
\end{align*}
\]

Products of siderite decomposition, Fe₃O₄, Fe and/or C are decarbonated sorbent which can re-absorb carbon dioxide, e.g.:

\[
2\text{Fe}_3\text{O}_4(s) + \text{C}(s) + 5\text{CO}_2(g) \rightarrow 6\text{FeCO}_3(s) \quad (5)
\]

Hence, initial material (Fe₃O₄/Fe₂O₃ + Fe) can be used for multiple cycles according to the combination of (1)–(5) reactions, as can be seen in Fig. 1. Carbon has the advantage that it is cheaper than iron.
2. **Experimental**

Mechanochemical reactions between magnetite or hematite and carbon dioxide were performed at room temperature and elevated CO$_2$ pressure (10–30 bar). Planetary ball mill Retsch PM100 was operated at 200–400 rpm. Vessel for the ball milling was a stainless-steel jar of 50 mL volume capable for holding up to 100 bar gas pressure. High purity CO$_2$ gas (Airgas, 99.999%) was loaded into the reactor at different pressures together with 3.0 g of magnetite (Alfa Aesar, nanopowder, 97%) or hematite (99.945%) and iron (Good Fellow, 99% purity, <60 μm) mixture in a molar ratio 1:1. The powder to balls (stainless steel) weight ratio was 2:27. Reactor was flushed several times with CO$_2$ gas to ensure a pure CO$_2$ atmosphere inside the reactor. Mechanochemical reaction was run for different periods of time from 30 min up to 36 h. Each 1 h milling interval was followed by half an hour cooling interval to avoid overheating of the sample.

Thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) were conducted in a temperature range of 25–1000 °C using TA Instruments SDT Q600 instrument. The measurements were performed in air and Ar atmospheres with a heating rate of 10 °C/min. Powder X-ray diffraction patterns were collected using Bruker GADDS/D8 diffractometer equipped with Apex Smart CCD Detector and molybdenum rotating anode. Collected 2D diffraction patterns were integrated using Fit2D software [38]. Quantitative phase analysis of the samples was performed using Rietveld method and GSAS package [39,40]. The CO$_2$ sorption capacity was calculated using the results generated by Rietveld refinement of XRD patterns. Scanning electron microscope (JEOL JSM-6330F) was used to study the morphology of powder particles. Surface area of the powders was measured using Brunauer–Emmett–Teller (BET) method and Microeritics Tristar II 3020 instrument. Raman spectroscopy characterization was used to identify carbon in siderite decomposition products. A continuous wave (CW) argon ion (Ar$^+$) laser (model 177G02, Spectra Physics) of 514.4 nm in wavelength was used as a source of monochromatic radiation. Backscattered Raman spectra were collected by a high-throughput holographic imaging spectograph (model HoloSpec f/1.8i, Kaiser Optical Systems) with volume transmission gratings, a holographic notch filter, and thermoelectrically cooled charge-coupled device (CCD) detector (Andor Technology). The Raman system has spectral resolution of 4 cm$^{-1}$. The spectra were usually collected with 10 min exposure.

3. **Results**

3.1. **Thermodynamic simulation of Fe$_3$O$_4$–Fe–CO$_2$ and Fe$_3$O$_4$–Fe–CO$_2$ systems**

FACTSAGE software and the databases therein, FACT – F’A’C’T 5.0, SGPS – SGTE and SGSL [41] were used to verify the thermodynamic feasibility of carbonation process for the systems Fe$_3$O$_4$–Fe–CO$_2$ and Fe$_3$O$_4$–Fe–CO$_2$. Fig. 2 shows the calculated equilibrium temperature as a function of pressure (phase diagram) in both systems. These results indicate that the siderite formation is favored either by high pressures at a constant temperature or by low temperatures at a constant pressure. Both systems have similar behavior. The Fe$_3$O$_4$–Fe–CO$_2$ system shows slightly lower equilibrium temperatures at the same pressure. This calculation indicates that carbonation is thermodynamically favorable during ball milling process at room temperature and decarbonation is possible at relatively low temperatures (about 180 °C at 1 atm pressure).

3.2. **Siderite formation through ball milling**

Fig. 3 illustrates kinetics of iron(II) carbonate formation in mechanochemical reaction between magnetite and iron at 30 bar CO$_2$ pressure and 400 rpm planetary ball mill revolution speed. Weak diffraction peaks of FeCO$_3$ are detected after 30 min milling interval and their intensities increase with milling time. Diffraction peaks of elemental iron vanish after 2 h of milling while Fe$_3$O$_4$ phase persists for 15 h of the reaction. Disappearance of Fe diffraction peaks but not those of magnetite in XRD patterns could be an indication of elemental iron amorphization on early stage of ball milling. The 36 h sample reveals the presence of siderite as a major phase and small amount of magnetite. Theoretical CO$_2$ absorption capacity of Fe$_3$O$_4$–Fe mixture in 1:1 molar ratio is 0.6126 g CO$_2$/g sorbent. Rietveld refinement of XRD pattern of 36 h sample yields siderite content 99.76% which is equivalent to 0.6101 g CO$_2$/g sorbent capacity. CO$_2$ absorption capacity which was calculated from TG experiment (see Section 3.3) is 0.5213 g CO$_2$/g sorbent. The difference is associated with small oxidation that occurs in TG when the regenerated oxides come into contact with CO$_2$ when it is released.

According to Ref. [28] synthetic siderite crystallizes in hexagonal space group R-3c with lattice parameters $a = b = 4.6919(8)$ Å and $c = 15.407(2)$ Å. Calculated lattice...
parameters of siderite in this study are \(a = b = 4.679(3) \text{ Å}\) and \(c = 15.281(7) \text{ Å}\) which agree with reported values.

Broadening of diffraction peaks of siderite during milling indicates about crystallite size reduction. Average crystallite size of FeCO\(_3\) in 36 h sample estimated using Scherrer's formula is 82 Å.

The energy spent in ball milling process can be calculated. According to Burgio et al. [42], the transferred energy to a mass, \(E_{\text{tm}}\), in a planetary ball milling can be obtained as a function of filling factor in the jar, \(\phi\), ball mass, \(m_b\), ball diameter, \(d_b\), number of balls, \(N_b\), reaction time, \(t\), angular rotation speeds of plate and jar, \(W_p\) and \(W_v\), radius of plate and jar, \(R_p\) and \(R_v\), and sample mass, \(P_W\), as follows:

\[
E_{\text{tm}} = \phi N_b m_b t (W_p - W_v) \left[ \frac{W_p^2 (R_v - (d_b/2))}{W_p + W_v W_p R_p} \right] \left( \frac{(R_v - d_b/2)}{2\pi P_W} \right)
\]

Fig. 2 – Equilibrium temperature of siderate decomposition as a function of pressure for the systems \(\text{Fe}_3\text{O}_4 + \text{Fe} + 4\text{CO}_2 \rightarrow 4\text{FeCO}_3\) and \(\text{Fe}_2\text{O}_3 + \text{Fe} + 3\text{CO}_2 \rightarrow 3\text{FeCO}_3\).

Fig. 3 – Evolution of X-ray diffraction patterns \((\lambda = 0.71073 \text{ Å})\) of \(\text{Fe}_3\text{O}_4 + \text{Fe}\) mixture at 30 bar \(\text{CO}_2\) pressure with a time of ball milling (400 rpm). Plot at the bottom shows XRD patterns of Fe (JCPDS #006-0696; black), \(\text{Fe}_3\text{O}_4\) (JCPDS #001-1111; blue) and \(\text{FeCO}_3\) (JCPDS #029-0696; red). Red dotted lines show positions of the diffraction peaks of siderite.
Fig. 4 – Evolution of XRD patterns (λ = 0.71073 Å) of Fe₂O₃ + Fe mixture during ball milling (400 rpm) at 30 bar CO₂ pressure. Plot at the bottom shows XRD patterns of Fe (JCPDS #006-0696; black), Fe₂O₃ (JCPDS #089-2810; green), Fe₃O₄ (JCPDS #001-1111; blue) and FeCO₃ (JCPDS #029-0696; red). Red dotted lines show positions of the diffraction peaks of siderite. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of the article.)

Fig. 5 – Siderite yield as a function of reaction time at 400 rpm for 30, 20 and 10 bars of CO₂ pressure in the Fe₃O₄–Fe–CO₂ and Fe₂O₃–Fe–CO₂ systems.

Here, to obtain siderite, at conditions of 30 bar, 400 rpm and 36 h, the spent of energy is 168.746 W h/g. Taking into account that siderite formation kinetics is faster while metallic iron is present, the material could be initially carbonated for 3 h, spending 14.062 W h/g, thus achieving a CO₂ capture capacity of 8.397 mmol CO₂/g sorbent, which translates to 60.234% of conversion. Sorbent performance in subsequent carbonation cycles improves due to reduction of particle size (see Section 3.4). In industrial processes, the demand for energy can be supplied by renewable energy sources.

Fig. 4 shows evolution of XRD patterns in the Fe₂O₃–Fe–CO₂ system during ball milling at 30 bar CO₂ pressure, 400 rpm and 0.5–3 h of reaction time. Formation of siderite in this system proceeds through Fe₃O₄ as an intermediate phase. It was shown [34] that hematite quickly converts to magnetite in mechanical ball milling process.

Formation of siderite has strong dependence on CO₂ pressure. Fig. 5 depicts siderite yield at different CO₂ pressures as a function of reaction time for two systems. Iron(II) carbonate yield increases with CO₂ pressure and reaction time.
Porous solid materials have interconnected pathways where gas molecules collide between them or with the pore walls. If the CO$_2$ gas pressure is high, the molecules can remain within each available pore of magnetite, hematite or iron forming FeCO$_3$. On the other hand, if the pressure is low, collisions between the CO$_2$ molecule and pore walls are dominant and the free path is restricted with the geometry of void space [43,44]. Initially, the CO$_2$ capture kinetics is faster in hematite system and at longer times, the siderite yield trends to stabilize, because of inverse reaction occurs at those conditions.

In mechanical ball milling, the reaction kinetics is strongly affected by revolution speed. Fig. 6 depicts CO$_2$ capture capacity of material for revolution speeds of 200 and 400 rpm as a function of reaction time for the two studied systems. According to these results, if the number of revolutions is lower, the siderite formation decreases; this behavior is observed in both systems. It suggests that, increase in revolution speed transfer higher kinetic energy to the materials promoting formation of crystal defects such as stacking faults, dislocations, vacancies, and increased number of grain boundaries [45-48].
The presence of these defects generates more active sites in magnetite, hematite and iron which facilitates their reaction with CO₂. In a planetary ball mill, increasing the speed of rotation will increase the speed with which the balls move. Above a critical speed, the balls will be pinned to the inner walls of the jar and do not fall down to exert any impact force. Therefore, the maximum speed should be just below this critical value so that the balls fall down from the maximum height to produce the maximum collision energy [48]. It is clear that for the studied conditions, 200 and 400 rpm are below such a critical speed. Here faster speeds were not used because of industrial processes limitations.

3.3. TG/DSC study of siderite decomposition

Siderite decomposition reactions were studied on a sample of pure siderite obtained at 30 bar CO₂ pressure, 400 rpm and 36 h in the system Fe₂O₃–Fe–CO₂. The calcination temperature (absorbent regeneration) of siderite was experimentally identified using thermogravimmetrical analysis.

According to the literature a two-step mechanism of decomposition of FeCO₃ is established. Formation of non-stoichiometric wüstite, FeO, in the first step of decomposition is followed by its transformation to other products depending on the experimental conditions, oxygen partial pressure, in particular:

$$\text{FeCO}_3 \rightarrow \text{FeO} + \text{CO}_2 \quad 37.99\% \text{weight loss} \quad (7)$$

Second-step reactions are listed below in order of increasing oxygen partial pressure. Wüstite is not thermodynamically stable below 563 °C and undergoes in a vacuum or inert atmosphere a disproportionation reaction, e.g. [34]:

$$4\text{FeO} \rightarrow 2\text{Fe}_2\text{O}_3 + \text{Fe} \quad \text{noweightchange} \quad (8)$$

Overall reaction of reactions (6) and (7):

$$4\text{FeCO}_3 \rightarrow 2\text{Fe}_2\text{O}_3 + \text{Fe} + 4\text{CO}_2 \quad 37.99\% \text{weight loss} \quad (9)$$

Wüstite also can be oxidized by carbon dioxide produced in the reaction (7) according reactions (10) and (12). Decomposition reactions of siderite (11) and (13) can also generate carbon dioxide:

$$3\text{FeO} + \text{CO}_2 \rightarrow 2\text{Fe}_2\text{O}_3 + \text{CO} \quad 7.42\% \text{weight gain} \quad (10)$$

$$3\text{FeCO}_3 \rightarrow 2\text{Fe}_2\text{O}_3 + \text{CO} + 2\text{CO}_2 \quad 33.39\% \text{weight loss} \quad (11)$$

$$2\text{FeO} + \text{CO}_2 \rightarrow 2\text{Fe}_2\text{O}_3 + \text{CO} \quad 11.14\% \text{weight gain} \quad (12)$$

$$2\text{FeCO}_3 \rightarrow 2\text{Fe}_2\text{O}_3 + \text{CO}_2 \quad 31.08\% \text{weight loss} \quad (13)$$

In the presence of oxygen, oxidation of FeCO₃ takes place very rapidly, yielding hematite or magnetite (in moderately oxidative atmosphere):

$$3\text{FeCO}_3 + 1/2\text{O}_2 \rightarrow 2\text{Fe}_2\text{O}_3 + 3\text{CO}_2 \quad 33.39\% \text{weight loss} \quad (14)$$

$$4\text{FeCO}_3 + \text{O}_2 \rightarrow 2\text{Fe}_2\text{O}_3 + 4\text{CO}_2 \quad 31.08\% \text{weight loss} \quad (15)$$

Moreover, it is worth to mention one more FeCO₃ decomposition pathway which was experimentally observed at low oxygen partial pressure in the presence of the graphite buffer [19]:

$$3\text{FeCO}_3 \rightarrow \text{Fe}_3\text{O}_4 + 3\text{C} + 5/2\text{O}_2 \quad 31.66\% \text{weight loss} \quad (16)$$

Fig. 7(a) shows the TG/DSC plot for the siderite decomposition in argon atmosphere. Endothermic peak at 367 °C can be assigned to the first step of siderite decomposition. According to TG plot the weight loss in the first step of siderite decomposition is 34.27% which is lower than predicted by reaction (7) 37.99%. The difference is due to the presence of small amount of unreacted Fe and Fe₂O₄. This amount can be calculated from TG curve of siderite decomposition in air atmosphere (Fig. 7(b)). Decomposition of FeCO₃ in air atmosphere is accompanied by iron oxidation to Fe₂O₃ according to reaction (15). Theoretical weight loss in this reaction is 31.08%. Fig. 7(b) shows TG curve of FeCO₃ decomposition in air atmosphere. Experimental weight loss is 30.68% and it is less than theoretical one. Siderite which was used in this TG experiment was produced by ball milling of equimolar mixture of Fe₂O₄ and Fe at 30 bar CO₂ pressure for 36 h. Difference between theoretical and measured weight loss in reaction (15) must be related to the presence in the siderite sample of unreacted equimolar mixture of Fe₂O₄ and Fe. Oxidation of both Fe and Fe₂O₄ are accompanied by the weight gain and the difference between theoretical and measured weight change in reaction (15) can be used to calculate amount of unreacted iron and magnetite. Calculations show that siderite contains 0.95 wt.% of equimolar mixture of Fe and Fe₂O₄.

If we take into account 0.95 wt.% of Fe + Fe₂O₄ as impurity in the siderite, the first decomposition step in argon atmosphere should be accompanied by 37.63% weight loss which is still larger than observed experimentally for siderite decomposition in Ar atmosphere loss of 34.27%. This evidences that simultaneously to the reaction of FeO formation from siderite and its disproportionation to Fe and Fe₂O₄, oxidation of FeO by CO₂ must occur according either to reaction (10) or (12). Reaction (16) also should be considered as a possible pathway of FeCO₃ decomposition. Fe and Fe₂O₄ are stable below 550 °C and react with each other above 550 °C yielding FeO. Thus, endothermic peak observed on DSC curve in Ar atmosphere at 566 °C can be assigned to FeO re-formation from Fe₂O₄ and Fe [33,34,49]. Small weight loss of 1.95% which is accompanied by observed endothermic peak at 668 °C could be related to Fe₂O₃ reduction by carbon [50]. Most of the reports on thermal decomposition of siderite refer to the natural mineral samples which are essentially solid solutions with Ca, Mg, and Mn carbonates. As it was mentioned earlier these impurities affect siderite decomposition temperature and kinetics. One of the rare reports on decomposition of synthetic FeCO₃ is by Gotor et al. [28]. They found that natural siderite with (Fe₀.₇Mg₀.₃)CO₃ composition decomposition obeys n-order kinetic model and the synthetic one could be described by an Avrami-Erofeev law. Also, difference in the final decomposition products was observed. Natural siderite sample yields wustite phase of the composition Fe₀.₇Mg₀.₃O and small amount of magnetite while products of synthetic siderite sample decomposition are magnetite and iron. Thermal decomposition of natural
siderite in air atmosphere is reported in the temperature range 465–550 °C (exothermic peak at 525 °C) [51], 478–580 °C (exothermic peak at 539 °C) [33]. In the inert atmosphere decomposition temperature shifts to slightly higher temperature 438–540 °C (endothermic peak at 500 °C) [33]. Kinetics of thermal decomposition of siderite in air atmosphere obeys three-dimensional diffusion model [33].

According to the thermogravimetric analysis, the siderite decomposition and therefore the release of carbon dioxide occurs in the temperature range 300–400 °C. Total weight loss in this range of temperature is 34.27%, translating the CO2 capture capacity of sorbent at that conditions to 0.5213 g CO2/g sorbent or 11.84 mmol CO2/g sorbent; this value is relatively close to the theoretical maximum which is 13.91 mmol CO2/g sorbent, demonstrating the high feasibility to capture CO2 by mechanical ball milling method. In Fig. 8 the X-ray diffraction patterns of the products of siderite decomposition in Ar atmosphere and vacuum are shown. It was found that siderite starts decomposing above 300 °C in Ar atmosphere and above 200 °C in vacuum. Magnetite is detected as decomposition product.
of siderite in both conditions, and metallic iron is identified at vacuum. In addition to XRD decomposition product were also characterized by Raman spectroscopy to test the presence of elemental carbon which could form in FeCO$_3$ thermolysis reaction (16).

Raman spectroscopy is a very useful technique to molecular morphology characterization of carbon materials. It is because of its high sensitivity to highly symmetric covalent bonds with little or no natural dipole moment. The carbon–carbon bonds that make up these materials fit this criterion perfectly and as a result Raman spectroscopy is highly sensitive to these materials and able to provide a wealth of information about their structure. Every band in the Raman spectrum corresponds directly to a specific vibrational frequency of a bond within the molecule [52,53]. The 1582 cm$^{-1}$ band of graphite is known as the G band [52,53]. Raman vibration at 1370 cm$^{-1}$ is disorder-induced mode of graphite (D mode) [53]. Fig. 9 shows Raman spectra of the products of siderite decomposition at 200°C and 300°C in vacuum. Characteristic peaks of graphite at 1582 cm$^{-1}$ and 1370 cm$^{-1}$ observed in this spectra confirm presence of graphite in decomposition product.

Metallic iron is one of the products of siderite decomposition in vacuum. This fact agrees with the earlier discussed mechanism and with Refs. [28–30] which report FeO as intermediate product of siderite thermolysis.

3.4. Carbonation–calcination cycles

Products of siderite decomposition were subsequently cycled in CO$_2$ absorption/release reactions. It was found that the capture capacity of the sorbent in the second carbonation cycle is significantly reduced due to the low content of reducing agent (Fe and C) in the mixture. Reducing agent deficiency was compensated by adding to the sorbent carbon in the form of graphite. XRD patterns of the products which are formed in the second carbonation cycle with different amounts of graphite are shown in Fig. 10. The carbonation was performed at 400 rpm, 30 bar CO$_2$ pressure for 20 h in planetary ball mill. Addition of 1 mol.% of graphite to the mixture leads to the reduction of Fe$_3$O$_4$ to FeO. Very little yield of siderite was detected for 1 mol.% graphite loading. Increase in graphite loading above 3 mol.% facilitates carbonation reaction and as it is evident from Fig. 10. Pure siderite can be obtained in the second carbonation cycle with graphite content 5.00 mol.%. Samples of FeCO$_3$ produced in the second and subsequent carbonation cycles were decomposed at 300°C in vacuum.

Siderite with a yield close to 95% was formed during third and fourth carbonation–calcination cycles, without addition of any extra reducing agent, which suggests that carbonation at this point is favored by two conditions, first, porosity features of material exposed to grinding treatment and second, the reducing agent yielded in the calcination. Fig. 11(a) shows XRD pattern of the products formed after the fourth cycle of carbonation, and Fig. 11(b) shows the products after calcination at 300°C in vacuum. The products of fourth CO$_2$ desorption cycle were identified as Fe$_3$O$_4$, Fe, and graphite which suggests that the material can be used in posterior CO$_2$ absorption/desorption cycles.

Table 1 shows pore volume and surface area of initial mixture of magnetite and iron, after 2 h of milling and after four cycles of carbonation–calcination. The surface area increases as a result of ball milling. Noticeably, the surface area after 4 cycles of carbonation and calcination is more than 15 times

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**Fig. 8 – X-ray diffraction patterns ($\lambda = 0.71073$ Å) of the products of siderite decomposition in Ar atmosphere and vacuum.** Plot at the bottom shows XRD patterns of Fe (JCPDS #006-0696; black), Fe$_3$O$_4$ (JCPDS #001-1111; blue) and FeCO$_3$ (JCPDS #029-0696; red). Red dotted lines show positions of the strongest diffraction peaks of siderite. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of the article.)
Fig. 9 – Raman patterns of siderite decomposition products at 200 and 300 °C in vacuum.

Fig. 10 – XRD patterns (λ = 0.71073 Å) of the products of the second carbonation cycles with different amounts of extra-graphite. Plot at the bottom shows XRD patterns of FeO (JCPDS #085-0625; black), and FeCO₃ (JCPDS #029-0696; red). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of the article.)
Fig. 11 – XRD patterns (λ = 0.71073 Å) of siderite formation products (a) after four cycles of carbonation and (b) products identified after calcination of the fourth cycle. Plot at the bottom shows XRD patterns of Fe_3O_4 (JCPDS #001-1111; blue), Fe (JCPDS #006-0696; black) and FeCO_3 (JCPDS #029-0696; red). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of the article.)

### Table 1 – Pore volume and surface area for initial mixture of magnetite and iron, after 2 h of milling and after four carbonation–calcination cycles.

<table>
<thead>
<tr>
<th></th>
<th>Surface area (m^2/g)</th>
<th>Pore volume (cm^3/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial mixture</td>
<td>4.66</td>
<td>0.013</td>
</tr>
<tr>
<td>2 h milled</td>
<td>16.36</td>
<td>0.045</td>
</tr>
<tr>
<td>After four cycles</td>
<td>73.45</td>
<td>0.120</td>
</tr>
</tbody>
</table>

higher than surface area of initial Fe_3O_4 + Fe mixture. Pore volume also increases with ball milling time.

Additionally, SEM reveals the appearance of material in the three states above mentioned. Fig. 12(a) shows that in the material not treated is possible to identify independently the particles of iron and magnetite; the iron particles are bigger and have spherical shape. The particle size of Fe_3O_4 is approximately 300 nm, while for iron the particle size rises 2.5 μm. In the appearance of the milled material for 2 h predominates the agglomeration as is shown in Fig. 12(b); despite of this agglomeration, material is more homogeneous and has larger surface area and pore volume; the particle size of milled material is approximately 380 nm. Bigger particle size and agglomeration in the milled material suggests that the tendency to cold welding predominates over fracture to conditions of 400 rpm revolution speed [45]. The carbonated and calcined material shows high discrepancy in the appearance of the particles, developing a broad range between particles size (0.1–30 μm) as can be seen in Fig. 12(c), indicating that during the process, cold welding predominates over fracture.

Fig. 12 – SEM images of (a) as received Fe_3O_4 and Fe (b) 2 h ball milled (c) after four carbonation–calcination cycles at magnification of 5000x, 5000x and 3000x, respectively.
Formation of biggest particles at 400 rpm and long reaction times (more than 20 h per cycle) is due to three factors: (1) the lattice strain falls with the refinement, hence the rate of cold welding is more than the rate of fracturing; (2) strong agglomerating force due to high energy per impact; (3) the ductile nature of iron based components [45].

4. Conclusion

This study presents a novel method to capture CO2 by mixtures of raw materials used in steel industry (Fe2O3 + Fe and Fe2O3 + Fe). Increases in siderite formation (CO2 capture capacity) during planetary ball milling conditions were achieved by increases of pressure, revolution speed of ball milling, and reaction time. Predominantly pure siderite was produced at 30 bar CO2 pressure, 400 rpm and 36 h of reaction time from Fe2O3 and Fe, which yields a CO2 capture capacity of 0.6101 g CO2/g sorbent. CO2 capture capacity in Fe2O3 and Fe system was higher compared with Fe2O3 and Fe system at the same conditions of pressure, temperature and reaction time. It was found that hematite quickly converts to magnetite during ball milling. Experimentally observed FeCO3 decomposition temperature was above 300 °C. Magnetite, carbon and iron were identified as decomposition products. The sorbent material was used in four carbonation–calcination cycles showing good stability of CO2 absorption capacity.

Conflicts of interest

The authors declare no conflicts of interest.

REFERENCES

Mechanical behaviour of particles between...