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Investigation of barium iron oxides for CO_2 capture and chemical looping oxygen uncoupling

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ABSTRACT

The performance of two underexploited ternary oxides – $Ba_3Fe_2O_6$ and $Ba_5Fe_2O_8$ are investigated for carbon dioxide capture and chemical looping oxygen uncoupling. The ternary compound $Ba_3Fe_2O_6$ was found to have a structure characterised by space group $Pa\overline{3}$. Experimental results have shown that both $Ba_3Fe_2O_6$ and $Ba_5Fe_2O_8$ are capable of cyclically capturing CO_2 at temperatures above 800 °C. $Ba_5Fe_2O_8$ demonstrated superior CO_2 capture performance compared to $Ba_3Fe_2O_6$, with consistent gravimetric CO_2 uptake capacities of 4.35 wt% and 13.39 wt% at 900 °C and 1000 °C, respectively, over 20 cycles. In comparison, $Ba_3Fe_2O_6$ demonstrated high initial CO_2 uptake capacities which deteriorated cyclically, with 20 cycle average capacities of 7.73 wt% and 11.99 wt% at 900 °C and 1000 °C, respectively. $Ba_3Fe_2O_6$ also exhibits excellent recyclability and satisfactory chemical looping oxygen uncoupling (CLOU) activity over temperature swing cycles between 550 °C and 950 °C. In contrast, the strong affinity with CO_2 makes $Ba_5Fe_2O_8$ unsuitable for application in chemical looping oxygen uncoupling air separation, especially in the presence of substantial partial pressures of CO_2 .

1. Introduction

CO₂ constitutes the largest proportion of greenhouse gas emissions, with an estimated mean atmospheric lifetime of 300 years, while 25 % of CO₂ emissions last indefinitely [1]. Electricity and heat generation activities are responsible for approximately half of the total anthropogenic CO₂ emissions [2]. The combustion of fossil fuel emits many pollutants and poses a significant environmental risk due to activities associated with their procurement. Carbon dioxide (CO₂) emissions from industrial activities, particularly from the combustion of fossil fuels, have been identified as the primary cause of climate change [3]. In response, the development of effective technologies for capturing and storing CO2 has emerged as an urgent solution to climate change. One such carbon capture technique is chemical looping (CL), which utilises solid intermediates to separate a desired reaction into two or more constituent reactions, preventing direct contact between reactants and enabling easier separation of the products of combustion [4]. This versatile approach can be applied to a variety of processes, including fossil fuel combustion, water splitting, air separation, and the synthesis of high-value products [5-8]. In general, chemical looping offers high energy conversion efficiency, reduces exergy losses, and provides safer operation by preventing the use of gaseous oxygen compared to conventional gas-phase redox chemical processes [6,9,10]. In most CL processes, a solid metal oxide, nitride, or hydride is used as an intermediate to transfer oxygen, nitrogen, or hydrogen, respectively, between individual reactors running different constituent reactions. These solid materials, commonly known as carriers, play crucial roles in chemical looping processes. While the interests in nitrogen and hydrogen carriers emerged recently, most of the carriers studied to date have been oxygen carriers (OCs) [8]. In the context of carbon capture and storage, the relevant CL processes include chemical looping combustion (CLC), chemical looping oxygen uncoupling (CLOU), chemical looping reforming, and chemical looping CO₂ capture (also known as sorbent looping or CO₂ looping).

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1.1. CO₂ looping

As depicted in Fig. 1(a), CO_2 looping is a cyclic process involving the reversible reaction between CO_2 and a metal oxide to form metal carbonates. The metal carbonate is subsequently decomposed to release CO_2 and regenerate the metal oxide, which can be used for further CO_2 capture. The reversible reactions by a generic bivalent metal carbonate are shown in reactions (1) and (2).

$$MeO(s) + CO_2(g) \rightarrow MeCO_3(s)$$
(1)

$$MeCO_3 (s) \rightarrow MeO (s) + CO_2 (g)$$
(2)

Natural minerals, such as limestone and dolomite have been used for the CO_2 looping processes, because of advantages such as their high reactivity, abundant availability, and low cost [11]. The use of synthetic oxides, including those Ca-based oxides and non-Ca-based oxides [11–16], for CO_2 looping can also enable the CO_2 looping process to efficiently capture CO_2 from a wide range of industrial processes with various operating temperatures and pressures (e.g. sorbent-enhanced reforming [17,18]). For post-combustion capture, CO_2 looping can be integrated with various power generation systems, including coal-fired power plants, natural gas power plants, biomass power plants, and waste-to-energy plants [19].

Various types of synthetic oxides have also been investigated for CO_2 looping, including monometallic oxides, mixed metal oxides, and doped oxides. The most common type of sorbents used in CO_2 looping are CaO and MgO [20,21]. CaO is widely utilised in CO_2 capture systems because of the low-cost and the availability of limestone and other Ca-rich minerals. Similarly, MgO can be produced from natural minerals such as magnesite, through relatively simple and energy-efficient production processes. Therefore, CO_2 looping process is often referred to as calcium looping (CaL). It has also been shown that the cost of carbon capture through CO_2 looping is on par with or even cheaper than amine scrubbing technology [22].

In addition to CaO/MgO-based sorbents, solid oxide sorbents based on other alkali and alkaline earth metals are also subjects of study for CO₂ capture at elevated temperatures [23–25]. The chemical properties of these materials can be tailored to optimise their CO₂ capture performance. Recently, ternary oxide sorbents have attracted substantial attention as potential contenders for CO₂ looping [26], e.g. Ca₂SiO₄, Li₄SiO₄, Li₂ZrO₃, Na₂ZrO₃ and CaFeO_{2.5} [27–31]. These sorbents typically possess intricate crystal structures with diverse metal coordination environments, rendering them potentials for a broad spectrum of applications [29,32,33]. Additionally, ternary oxide sorbents have demonstrated superior stability over cycles of CO₂ capture and release compared to conventional oxide-based sorbents (e.g. CaO and MgO-based sorbents), owing to their robust resistance to sintering and their ability to retain their pore structure during the CO_2 looping process [12]. Furthermore, the inherent structural stability of the ternary oxide allows the A sites and B sites to be substitutionally doped without major structural changes, opening new avenues for synthetically optimising the sorbent performance. Therefore, the investigation of ternary oxide-based sorbents is important for developing effective CO_2 capture and storage processes with high CO_2 capture capacity, good cyclic stability, tuneable properties, and compatibility with a wide range of industrial applications. Further research is also needed to optimise their properties for improved CO_2 uptake and release kinetics, as well as long-term stability regenerability, before they can be commercially deployed for large-scale CO_2 capture.

1.2. Chemical looping oxygen uncoupling (CLOU)

Chemical looping oxygen uncoupling (CLOU) is a high temperature process exploiting principles similar to CLC, to produce oxygen-enriched process streams via the thermal decomposition of solid metal oxide carriers [5]. The uncoupling of oxygen is a two-step redox reaction:

$$Me_{x}O_{y-2}(s) + O_{2}(g) \rightarrow Me_{x}O_{y}(s)$$
(3)

$$Me_xO_y(s) \rightarrow Me_xO_{y-2}(s) + O_2(g)$$
 (4)

Similar to CLC, CLOU can be achieved using two reactors, namely the oxidation reactor and the reduction reactor, in which reactions (3) and (4) take place, respectively. The CLOU scheme is illustrated in Fig. 1(b).

As shown in Fig. 1(b), fully oxidised metal oxides enter the reduction reactor and undergo reaction (4) in the presence of a carrying gas (usually steam or CO_2). The mixture of carrying gas and gaseous O_2 exiting the reduction reactor can be used as an oxidant for various applications (including oxy-fuel combustion). The reduced particles are recycled to the oxidation reactor where reaction (3) takes place in the presence of hot air. The oxygen-depleted air is then either used to preheat the incoming air or, in the case of pressurised operation, expanded in a gas turbine to generate power. In ideal conditions, the net heat released over the two reactions is zero. In this context, the oxygen carrier particles also serve as heat carriers to transport heat from the exothermic reaction (3) to the endothermic reaction (4).

Amongst the oxides of 3d transition metals, the most suitable candidates for CLOU are MnO_2/Mn_2O_3 , Mn_2O_3/Mn_3O_4 , CoO/Co_3O_4 , and CuO/Cu_2O . Besides monometallic metal oxides, ternary oxide materials can be also used as oxygen carriers for CLOU applications [34–36]. The tunability of the site occupancies and oxygen vacancies makes it easy to tailor the oxide structures for specific chemical looping applications. For example, when high oxygen partial pressures are desired, perovskites



CO₂ looping using a metal carbonate

CLOU process

Fig. 1. Schematic illustration of (a) CO₂ looping using a metal carbonate (b) CLOU process.

such as SrFeO_{3- δ} can be used. The oxygen deficient SrFeO_{3- δ} was studied for hydrogen production and catalytic CH₄ combustion applications in TGA and fluidised beds [36]. The study showed reversible phase transitions between the perovskite and deep reduction products (viz. SrO and Fe). In addition, ternary oxides could potentially alleviating some of the problems and risks associated with conventional CLOU materials, such as loss of activity, deactivation due to thermal stress, attrition, agglomeration, and cost of OCs [37,38].

1.3. Ba-ferrite ternary oxides for chemical looping applications

Being the most earth-abundant and cheapest transition metal, ironcontaining ternary oxides have received widespread research interest by the chemical looping community, with most research focusing on Ca, Sr, and Ba ferrite systems [39–42]. A unique feature of ternary oxides systems is the presence of Fe⁴⁺, which is highly redox-active. In comparison, Fe generally takes +2 and +3 oxidation states in its native oxides, e.g., FeO, Fe₃O₄ and Fe₂O₃ [43]. A Mössbauer spectroscopy study examining the structure for the cubic perovskite SrFeO₃ confirmed the high valence of Fe. This compound was proven to be a metallic conductor due to the wide conduction bands and the strongly hybridised Fe-3d and O-2p orbitals [44]. Another alkaline earth metal ferrite, CaFeO₃ was also shown to exhibit similar conducting behaviour due to delocalisation of electrons [45,46].

Ternary oxides of Ba and Fe, e.g. $BaFeO_3$ has also been known to stabilise the Fe^{4+} [47]. They can be prepared by solid state reaction between magnetite and a barium precursor such as $BaCO_3$ and $Ba(NO_3)_2$ [47]. For chemical looping applications, $BaFe_2O_4$ has shown promising performance for chemical looping gasification of biochar [48]. Chen et al. [49] also used $BaFe_2O_4$ for chemical looping biomass gasification and found that an OC loading as low as 10 wt% (2 g fuel and 0.2 g OC) was able to produce higher hydrogen yield than other OC loadings tested (ranging from 0 to 50 wt%).

As shown in both the isopleth phase diagram (Fig. 2(a)) and the isothermal phase diagram (Fig. 2(b)) of the Ba-Fe-O system, moving from the Fe-rich side to the Ba-rich side on the left, the Ba-Fe-O system would form, in sequence, Fe_2O_3 , $BaFe_1_2O_{19}$, $BaFe_2O_4$, $BaFeO_3$, $Ba_2Fe_2O_5$, $Ba_3Fe_2O_6$ and $Ba_5Fe_2O_8$ and BaO. While mixed Fe-rich barium ferrites such as $BaFe_2O_4$ and $BaFe_{12}O_{19}$ (barium hexaferrite, commonly used as a permanent magnet) have been studied for chemical looping applications [50], the phases at the Ba-rich end of the Ba-Fe-O system, e. g., $Ba_3Fe_2O_6$ and $Ba_5Fe_2O_6$, remain largely unexplored. Montorsi and Brisi [51] found that $Ba_3Fe_2O_6$ was stable in air above 800 °C (as seen in Fig. 2(a)). Below 800 °C, $Ba_3Fe_2O_6$ would decompose into Fe⁴⁺-containing compounds, e.g., $BaFeO_3$. While the structure of

Ba₃Fe₂O₆ was reported to be similar to that of Sr₃Al₂O₆ (space group $Pa\overline{3}$), Ba₅Fe₂O₈ has only been identified by calorimetry measurements with unresolved crystal structure [52,53]. Compared to Fe₂O₃, BaFe₂O₄ and BaFeO₃, the reducibility of Ba₃Fe₂O₆ and Ba₅Fe₂O₈ appears low, as suggested by Fig. 2(b). On the other hand, given the high Ba/Fe ratio, Ba₃Fe₂O₆ and Ba₅Fe₂O₈ are expected to have high basicity for CO₂ capture via carbonation to form BaCO₃.

In this paper, the two Ba-rich barium ferrite phases, viz. $Ba_3Fe_2O_6$ and $Ba_5Fe_2O_8$, are studied as CO_2 sorbents and oxygen carriers for CO_2 capture and chemical looping oxygen uncoupling (CLOU) [54], respectively. This work represents an exploratory effort to experimentally investigate the two Ba-rich ferrite phases for their potential applications in chemical looping processes.

2. Experimental

2.1. Materials synthesis

The two ternary oxides, i.e., Ba₃Fe₂O₆ and Ba₅Fe₂O₈, were prepared using the modified Pechini method. In a typical synthesis to prepare 1 g of barium ferrite sample, calculated amount of barium nitrate (Ba (NO₃)₂) and iron nitrate nonahydrate (Fe(NO₃)₃·9H₂O) were dissolved in 150 ml Type II deionised (DI) water. The specific amounts of precursors used are shown in Table 1. The precursor solution was maintained at 50 °C on a heating plate with magnetic stirring at 300 rpm for 30 min. The chelating and crosslinking agents used were citric acid (C₆H₈O₇) and ethylene glycol (C₂H₆O₂), respectively. The specified amounts of these agents (see Table 1) were added to the solution. Then, the temperature of the solution was increased to 80 °C until a dark brown viscous gel formed. After gel formation, the stirring was stopped, and the gel was allowed to dry at 105 °C in a drying oven for at least 12 h. To remove the organic carbon, the dried gel was placed in a crucible and heated in a muffle furnace at 400 °C for 6 h. The organic components of this gel would devolatilise, decompose, and combust in the muffle furnace leaving behind a residue resembling a sponge like

Table	1
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Amounts of precursors used for the synthesis	of 1 g	s of barium	ferrites
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Precursor	$Ba_3Fe_2O_6$ (amount in g)	$Ba_5Fe_2O_8$ (amount in g)
Ba(NO ₃) ₂	1.27	1.41
Fe(NO ₃) ₃ ·9H ₂ O	1.30	0.87
Citric acid	3.88	3.63
Ethylene glycol	1.88	1.76



Fig. 2. (a) A quasi-binary section of BaO-Fe₂O₃ in air 800–1200 °C and (b) Phase diagram of the Ba-Fe-O system showing oxygen potential versus Ba/(Ba+Fe) ratio at 827 °C [53].

texture. This residue was ball milled in a Focucy planetary ball mill (F-P4000) for 2 h at ~400 rpm and sieved into size fraction of 100–200 μ m. The sieved particles were subsequently calcined in air in a muffle furnace at 1000 °C for 15 h with a heating rate of 5 °C/min. The calcined samples were sieved again to the size fraction of 100–200 μ m before testing. The oxide samples are named Ba_x, where x = 3 refers to Ba₃Fe₂O₆ and x = 5 refers to Ba₅Fe₂O₈.

2.2. Thermogravimetric analysis

The barium ferrites' ability to (a) reversibly take up CO_2 over CO_2 looping cycles and (b) release and take up gaseous oxygen over reduction-oxidation cycles was investigated by thermogravimetric analysis (TGA, Mettler Toledo TGA/DSC2 LF 1100). The TGA equipment consisted of an electrically heated high temperature furnace and a microbalance with an accuracy of 0.1 µg incorporated into the furnace to measure the mass of the sample. Calibrated rotameters were utilised to regulate the flow rates of both the purge gas and protective gas.

2.3. CO₂ uptake

In a typical CO₂ looping experiment, approximately 50 mg powdered sample was loaded into a 150 μ L alumina pan, which was inserted into the reaction chamber of the TGA. During each experiment, the sample was firstly heated from ambient temperature to the reaction temperature, which ranged between 800 and 1000 °C, while purged by pure N₂ gas flowing at a rate of 50 ml/min. At the reaction temperature, the sample was alternatingly exposed to CO₂ and N₂ gases, each flowing at 50 ml/min for 30 min, for 20 cycles. Therefore, each CO₂ uptakeregeneration cycle was 60 min. Based on the recorded weight changes, the CO₂ capture efficiency of the sorbents is estimated according to:

$$\eta_{\text{CO2,up}} = \frac{\text{observed mass change per cycle, averaged over 20 cycles}}{\text{stoichiometric CO}_2 \text{ uptake capacity}} \times 100\%$$
(5)

where the stoichiometric CO_2 uptake capacity, in mass basis (i.e. mass of CO_2 taken up/mass of the fully calcined sample), is calculated based on the following reactions:

$$Ba_3Fe_2O_6 + 2CO_2 \rightleftharpoons 2BaCO_3 + BaFe_2O_4 \tag{6}$$

$$Ba_{5}Fe_{2}O_{8} + 4CO_{2} \rightleftharpoons 4BaCO_{3} + BaFe_{2}O_{4}$$
(7)

Separately, the experimentally measured CO_2 uptake capacity, also referred to as the gravimetric CO_2 uptake capacity in $mg_{CO2}/g_{sorbent}$, is defined by the ratio of weight gain by the sample during carbonation relative to its minimum weight measured during calcination, within a CO_2 capture-regeneration cycle.

2.4. Chemical looping oxygen uncoupling

In a typical CLOU experiment, approximately 50 mg of powdered sample was put in a 150 μ L alumina pan, which was subsequently inserted into the reaction chamber of the TGA, where the sample was exposed to alternating oxidising (50 ml/min of air at 550 °C) and reducing (50 ml/min of N₂ at 950 °C) environments. Once the sample temperature reached the set points of 550 °C and 950 °C, the sample was immediately heated up or cooled down at a rate of 10 °C /min, respectively, without any isothermal dwelling. Based on the measured weight changes, the O₂ release efficiency of the sorbents is estimated according to:

$$\eta_{O2,rel} = \frac{\text{observed mass change per cycle, averaged over 20 cycles}}{\text{stoichiometric }O_2 \text{ release capacity}} \times 100\%$$

where, the stoichiometric O_2 release capacity is calculated based on the following reactions:

$$2BaFeO_3 + BaO \rightleftharpoons Ba_3Fe_2O_6 + 0.5O_2 \tag{9}$$

$$2BaFeO_3 + 3BaO \rightleftharpoons Ba_5Fe_2O_8 + 0.5O_2 \tag{10}$$

2.5. Material characterisation

2.5.1. X-ray diffraction

Powdered XRD spectra at room temperature and pressure were collected using a Bruker D8 Advance diffractometer with filtered Cu K α radiation ($\lambda = 1.5418$ Å) at 40 kV and 40 mA under ambient conditions. The XRD patterns were collected in a 2 θ range from 10 to 90° with a step size of 0.02° and a collection time of 1 s/step.

The *in situ* XRD measurements were carried out with a PANalytical Empyrean X-ray powder equipped with a X'Celerator Scientific ultrafast line detector and Bragg-Brentano HD incident beam optics using Cu K α radiation (45 kV and 40 mA). The instrument is fit with an Anton Paar XRK 900 reactor chamber with a Macor sample holder in reflection geometry and gas flow through the sample with uniform temperature control. A 2 θ range of 20°–80° with a scanning speed of 0.022° s⁻¹ and a step size of 0.016° was employed. Phase identification was performed with the help of Match! V3.0 software.

In situ XRD measurements were performed to investigate the phase transitions of the barium ferrites at high temperature chemical looping cycles. To confirm the successful formation of the barium ferrite phases, the fresh samples were measured by XRD while being heated in air from 500 to 900 °C. Similarly, to explore the phase transitions of Ba₃Fe₂O₆ during CO₂ uptake, XRD measurements were conducted when the sample was firstly heated in air at 550 °C, then exposed to a mixture of 20 % CO₂ in N₂ for 20 min. The CO₂ mole fraction of 20 % was chosen because it is the upper limit that the *in situ* XRD reaction chamber could tolerate. The temperature was subsequently raised to 900 °C under the same gas mixture and sustained in 900 °C for a further 60 min, followed by a further XRD measurement. Finally, the inlet gas was switched to N₂ for an additional 60 min at 900 °C to simulate the sorbent regeneration, before the last XRD pattern was measured.

2.5.2. Scanning electron microscopy

The microstructure and morphological features of the ternary oxide samples were analysed using field emission scanning electron microscopy (FESEM). The measurements were carried out on a JEOL JSM-7200F electron microscope. The imaging was done at an accelerating voltage of 5 kV under low vacuum and room temperature conditions. Prior to imaging, the samples were mounted onto cylindrical aluminium sample holders using sticky carbon tape. The samples were then sputter coated with a thin layer of platinum using a sputter coater to increase the conductivity and enhance the contrast of the samples. A JEOL JEC-30000FC sputter coater was used for this purpose. The sputter coating was carried out at a current of 20 mA for 60 seconds. The magnification of the SEM images ranged from x100 to x20,000 depending on the morphological features of interest.

The SEM-EDX (energy-dispersive X-ray) analysis was performed with the above-mentioned FESEM unit equipped with an Oxford Instruments X-Max N EDX detector. The SEM was operated at an acceleration voltage of 20 kV. The SEM-EDX data was acquired using the Aztec software package. The software was used to control the SEM and EDX detector, and to acquire SEM images and EDX spectra from the sample. EDX spectra were collected from multiple areas of interest, and elemental maps were generated to visualise the spatial distribution of the elements.

(8)

3. Results and discussions

3.1. Structural characterisation

The energy-dispersive X-ray spectroscopy (EDX) analysis of the Ba₃ and Ba₅ samples, as shown in Fig. 3, confirms the elemental composition of the samples, i.e. the relative contents of Ba, Fe, and O are in accordance with the target stoichiometries (i.e. 66.5 wt% and 74.1 wt% barium for Ba₃ and Ba₅, respectively). The Pt signals in the EDX spectra arises from the Pt sputter coating during sample preparation for SEM. Therefore, the synthesis was deemed successful in producing samples with the correct stoichiometry, of Ba₃Fe₂O₆ and Ba₅Fe₂O₈ at high temperatures.

We examine the phase composition of the synthesised barium ferrite samples by means of *in situ* XRD. As seen in Fig. 4(a), the XRD spectra of Ba₃ at different temperatures show that the sample exhibited two distinct phases, namely BaFeO₃ and BaCO₃, in the temperature range of 500-800 °C. Above 800 °C, a unique set of peaks attributed to Ba₃Fe₂O₆ were observed, indicating that Ba₃Fe₂O₆ is only stable above temperatures of 800 °C. The unit cell of the Ba₃Fe₂O₆ lattice is shown in Fig. 4(b); it is isostructural to Sr₃Al₂O₆, belonging to the *Pa*3 space group [53].

For Ba₅, two distinct phases of BaFeO₃ and BaCO₃ are also seen the range of 500-800 °C, as shown in Fig. 4(c). Similarly, phase change was observed above 800 °C, when the mixture of BaFeO₃ and BaCO₃ was replaced with a set of unindexed peaks. Given the unsolved crystal structure of Ba₅Fe₂O₈ [52], it is not entirely certain whether Ba₅Fe₂O₈ has truly formed, despite the calorimetric observation by Montorsi and Brisi [51]. Nevertheless, the phase transition at 800 °C is in line with the isopleth Ba-Fe-O phase diagram shown in Fig. 2(a). Based on the *in situ* XRD results, the phase transitions observed can be summarised by Eqs. (11) & (12), and denoted by the black arrows shown in the magnified triangular phase diagram of the Ba-Fe-O system, in Fig. 4(e). Notably, the phase transitions involve transformation of Fe⁴⁺ containing phases

(e.g. SrFeO₃) to Fe³⁺ containing phases (e.g. $Ba_3Fe_2O_6$ and $Ba_5Fe_2O_8$) as the temperature increases beyond 800 °C.

$$2BaFeO_3 + BaCO_3 \rightleftharpoons Ba_3Fe_2O_6 + CO_2 + 0.5O_2$$

$$(11)$$

$$2BaFeO_3 + 3BaCO_3 \Rightarrow Ba_5Fe_2O_8 + 3CO_2 + 0.5O_2$$
 (12)

3.2. CO₂ capture experiments

3.2.1. TGA performance of CO₂ uptake at 1000 °C

Fig. 5(a) shows the cyclic performance of both Ba3 and Ba5 at 1000 °C over 20 cycles in alternating atmosphere of CO₂ (simulating carbonation) and N_2 (simulating calcination). $Ba_3Fe_2O_6$ exhibited a notable decay in CO₂ uptake capacity, which may be attributed to the apparent deterioration in carbonation kinetics. Specifically, the carbonation of Ba₃Fe₂O₆ was rapid during the first cycle, reaching a uptake of 142 mg CO₂/g sorbent within 600 s. However, by the 20th cycle, a relative weight gain of only 96.1 mg CO_2/g sorbent was reached within 600 s. On the other hand, the apparent kinetics of regeneration (i. e., decomposition of BaCO₃) of Ba₃ appears consistent and fast over the 20 cycles. The average gravimetric CO₂ uptake capacity of Ba₃Fe₂O₆ over the 20 CO₂ capture cycles (as shown in Fig. 5(a) & 5(b)) was 127.88 mg CO_2/g sorbent. It should be noted that, given the limitation of the TGA instrument, which only allowed us to perform multiples of 5cycle programmes with small pauses in-between, the samples were subjected to a longer calcination period every five cycle, resulting in a lower calcined mass and subsequently higher apparent gravimetric CO2 uptake capacity for every 5 cycles.

 $Ba_5Fe_2O_8$ exhibits slower carbonation during the initial 10 cycles (see Fig. 5(a)). Interestingly, the carbonation kinetics gradually accelerated over cycles, resulting in increased cyclic CO₂ uptake, as shown in Fig. 5(b). The cycle-averaged gravimetric uptake capacity of Ba₅ is



Fig. 3. SEM images of (a-b) fresh samples of Ba₃ and Ba₅, respectively and (c, d) EDX map spectrum of Ba₃ and Ba₅, respectively.



Fig. 4. (a) *In situ* XRD diffraction spectra of Ba_3 in air from 500 to 900 °C (b) Unit cell of $Ba_3Fe_2O_6$ (c) *in situ* XRD diffraction spectra of Ba_5 in air from 500 to 900 °C (d) Isothermal section of the Ba-Fe-O system for temperatures between 700 and 1000 °C. (e) a zoomed-in view of the triangular phase diagram [53].

147.5 mg CO₂/g sorbent. The decomposition kinetics of Ba₅ remained consistently fast, suggesting that the irreversible changes in the ternary oxides that are responsible for the changes in carbonation kinetics do not affect the decomposition kinetics. In addition, the fact that the apparent change in carbonation kinetics could be captured over CO₂ looping cycles suggests that the TGA measurements were not dominated by mass transfer effects.

Based on the measured maximum gravimetric CO_2 uptake capacities of 265 mg CO_2/g sorbent wt% and 271 mg CO_2/g sorbent for Ba₃ and Ba₅, respectively, the following reaction stoichiometries are proposed:

$$Ba_3Fe_2O_6 + 2CO_2 \rightleftharpoons 2BaCO_3 + BaFe_2O_4$$
(13)

 $Ba_{5}Fe_{2}O_{8} + 4CO_{2} \rightleftharpoons 4BaCO_{3} + BaFe_{2}O_{4}$ (14)

Reactions (13) and (14) corresponds to gravimetric CO₂ uptake

capacities of 284 mg CO₂/g sorbent and 379.80 mg CO₂/g sorbent, respectively. Using the cycle-averaged CO₂ uptake capacities of 239.80 mg CO₂/g sorbent and 267.80 mg CO₂/g sorbent, the CO₂ capture efficiencies, $\eta_{CO2,up}$ of Ba₃ and Ba₅ are calculated to be 84.43 % and 70.51 %, respectively.

3.2.2. Effect of operating temperature on CO_2 looping performance

To examine the impact of operating temperature on the CO_2 capture performance of $Ba_3Fe_2O_6$ and $Ba_5Fe_2O_8$, 20 isothermal CO_2 swings were performed at 800 and 900, in addition to 1000 °C. All experimental results are shown in Fig. 6.

For Ba₃ (Fig. 6(a)), the weight change curve measured at 900 °C is similar to that of 1000 °C. At 900 °C, the rate of carbonation appears more rapid, but also undergoes obvious cyclic deactivation, probably owing to sintering. Overall, the CO₂ capture capacities at 900 °C are in



Fig. 5. (a) Relative mass change and (b) gravimetric CO₂ capture capacity of the two samples over 20 carbonation-calcination cycles in TGA at 1000 °C in alternating CO₂ and N₂ environment.



Fig. 6. Relative mass change over carbonation-calcination cycles, facilitated by alternating CO₂ and N₂ environment isothermally at 800, 900 and 1000 °C in a TGA for (a) Ba₃Fe₂O₆ and (b) Ba₅Fe₂O₈.

the range of 70–80 mg CO₂/g sorbent ($\eta_{CO2,up} = 48.19 - 55.07$ %). At 800 °C, Ba₃ showed much lower CO₂ uptake capacity, in the range of 12–16 mg CO₂/g sorbent. In fact, Ba₃ appears to stay in a largely carbonated state at 800 °C with rather sluggish calcination. This observation is in line with the *in situ* XRD results shown in Fig. 4(a), which suggests that Ba₃ consists of a mixture of BaCO₃ and BaFeO₃ at 800 °C. The residual activity registered at 800 °C is attributed to the reversible carbonation of a small fraction of BaO or BaFeO₃. In general, Fig. Fig. 6(a) shows that calcination is kinetically hindered at lower temperatures, i.e., CO₂ looping employing Ba₃ should be preferably carried out at >800 °C.

For Ba₅Fe₂O₈ (Fig. 6(b)), contrasting the trend observed at 1000 °C, the CO₂ looping performance at 900 °C showed cyclic deactivation, with CO₂ uptake capacity falling from 49.76 mg CO₂/g sorbent ($\eta_{CO2,up} = 25.79$ %) in the first cycle the 1st cycle from to 35.61 mg CO₂/g sorbent ($\eta_{CO2,up} = 18.59$ %) in the 20th cycle. This decay appears to be associated with the increasingly sluggish carbonation kinetics, probably as a result of sintering and the significant resistance to mass transfer through

the product layer. Similar to Ba₃, the performance of Ba₅ at 800 °C is limited owing to high stability of the BaCO₃, which were difficult to decompose. The highest weight changes (i.e., gravimetric CO₂ uptake capacities) observed at 800 °C and 900 °C were 1.96 % and 7.87 %, respectively. Interestingly, the sintering-induced deactivation that is apparent at 900 °C seems less obvious in experiments at 1000 °C, although the samples are expected to sinter more at higher temperature. This disparity may be attributed to the presence of other rate-enhancing factors at higher temperatures such as faster ionic diffusion through the carbonate product layer, which could alleviate the rate-inhibition effects induced by sintering. Therefore, both Ba-rich barium ferrite phases are suitable reversible CO₂ sorbents for high temperature CO₂ capture cycles (i.e., >900 °C).

3.2.3. Mechanism of carbonation and calcination

Between the two ternary oxides investigated, the crystal structure of $Ba_3Fe_2O_6$ has been resolved. Regardless, *in situ* XRD analysis was performed on both the ternary oxides to further investigate the phase

transitions occurring during high-temperature CO_2 looping cycles. The resulting diffraction patterns of Ba_3 and Ba_5 ternary oxides are shown in Fig. 7(a) and (b).

From Fig. 7(a), it can be seen that after 20 min carbonation in 20 % CO₂ at 550 °C, Ba₃ primarily consisted of BaFeO₃, in agreement with the TGA results. Upon increasing the temperature to 900 °C, while keeping the gas environment as 20 % CO₂, the BaCO₃ phase transitioned from an orthorhombic structure (Pmcn) to a trigonal one (Fm-3m), resulting in a change in the diffraction pattern. This phase transition of BaCO₃ typically occurs at 811 °C, according to Antao and Hassan [55]. The reference XRD patterns along with the crystal structures of the two encountered BaCO₃ phases are shown in Fig. 7(c) and (d). BaFe₂O₄ is also detected upon high temperature carbonation, in agreement with Eq. (13). After carbonation at 900 °C for 60 min, the atmosphere was switched to N₂ for 60 min at 900 °C to facilitate calcination. During calcination, Ba₃Fe₂O₆ was regenerated. The changes observed via the *in situ* XRD measurements validate that the cyclic carbonation and calcination of Ba₃ indeed follows Eq. (13).

Unlike Ba₃, the low temperature carbonation of Ba₅ C in 20 % CO₂ at 550 ° for 20 min resulted in BaCO₃ and BaFe₂O₄ (Fig. 7 (b)), corroborating the proposed carbonation mechanism in Eq. (14). Similar to the case of Ba₃, BaCO₃ formed from Ba₅ underwent a phase transition from an orthorhombic structure (Pmcn) to a trigonal one (Fm-3m). Calcination of the Ba₅ sample at 900 °C in N₂ for 60 min resulted in the extensive formation of Ba₃Fe₂O₆. Despite the similarity in forming Ba₃Fe₂O₆ upon calcination, the two barium ferrite phases behaved differently over CO₂ looping cycles, as shown in Figs. 5 and 6. The nature, the origin and the consequence of this difference ought to be further investigated in future studies. In addition, the fate of the excess Ba in the calcined Ba₅ is also a subject for future investigations.

sorbents and oxygen carriers are almost always affected by the morphologies (including porosity and surface area) of the looping materials. Accordingly, the morphological changes of the barium ferrite samples over CO_2 capture cycles were examined by SEM and BET analysis.

Fig. 8 shows the SEM images of Ba₃ and Ba₅ before and after 20 redox cycles of CO₂ capture. Fig. 8(a) and (b) show that the freshly prepared samples are porous, consisting of mesoporous aggregates of grains of size 2 –4 μ m and 3–5 μ m for Ba₃ and Ba₅, respectively. These porous structures give rise to moderately high BET surface areas, as shown in Table 2, which in turn result in faster carbonation kinetics during the first few CO₂ looping cycles. After cycling, the porous structure was replaced by a dense one, while the specific surface area of Ba₃ dropped drastically from 8.8 m²/g to 0.87 m²/g; both are obvious signs of sintering, accommodated by the significantly slowed carbonation kinetics, as shown in Fig. 5(a). Interestingly, the calcination kinetics were not affected by the reduction in BET surface area. This observation resembles the disconnection between severe sintering and stable CO₂ uptake performance reported by an earlier studying using Ba₄Sr₂O₉ as the CO₂ sorbent [12].

For Ba₅, the fresh sample also displayed a moderately high surface area of $12 \text{ m}^2/\text{g}$. After 20 cycles of CO₂ capture at 900 °C, the surface of Ba₅ appears to have lost most of its porosity, showing signs of severe sintering (Fig. 8(d)). Similarly, the specific surface area of the cycled Ba₅ dropped to 1.93 m²/g after CO₂ cycling (Table 2). Therefore, sintering is responsible for the decaying performance of Ba₅ during CO₂ looping cycles at 900 °C, as shown in Fig. 6. Nevertheless, the severe sintering does not seem to affect the CO₂ uptake by Ba₅ at 1000 °C.

3.3. Chemical looping oxygen uncoupling

3.2.4. Morphological changes over CO₂ looping cycles

In chemical looping systems, the activity and performance of the CO₂

3.3.1. Oxygen release and uptake performance in TGA

The chemical looping oxygen uncoupling (CLOU) performance of the barium ferrite samples was also investigated by TGA. Each CLOU cycle



Fig. 7. In situ XRD pattern of **(a)** Ba₃Fe₂O₆ and **(b)** Ba₅Fe₂O₈, after being exposed to 20 min in 20 % CO₂ at 550 °C, followed by 60 min in 20 % CO₂ at 900 °C, and subsequently 60 min in N₂ at 900 °C. Reference XRD spectra and crystal lattice of BaCO₃ in (c) orthorhombic form and (d) trigonal form.



Fig. 8. SEM images of (a) and (b) - fresh samples, (c) and (d) after 20 cycles of CO₂ capture at 900 °C.

Table 2 BET surface area and pore size of the fresh and spent samples post \mbox{CO}_2 looping in TGA.

Sample	Surface area (m ² /g)	Pore size (nm)
Fresh Ba ₃ Fe ₂ O ₆	8.8	8.30
Fresh Ba ₅ Fe ₂ O ₈	12	10.96
Spent CO ₂ looping Ba ₃ Fe ₂ O ₆	0.87	20.85
Spent CO ₂ looping Ba ₅ Fe ₂ O ₈	1.93	24.50

consists of (1) air oxidation while heating up from 550 to 950 °C at 10 °C/min followed by (2) N₂ decomposition while cooling down from 950 to 550 °C at 10 °C/min, as shown in Fig. 9. The kinetics of both reduction and oxidation of Ba₃ were fast and consistent over 10 cycles. The observed mean mass change of Ba₃ is 1.25 wt% over the 10 redox cycles. Based on stoichiometry, the following oxygen release reactions are proposed:

$$2BaFeO_3 + BaO \rightleftharpoons Ba_3Fe_2O_6 + 0.5O_2 \tag{15}$$

$$2BaFeO_3 + 3BaO \rightleftharpoons Ba_5Fe_2O_8 + 0.5O_2 \tag{16}$$



Fig. 9. Cyclic oxygen uncoupling in a coupled temperature and pressure swing between 550 °C in air and 950 °C in N₂.

Based on the stoichiometries shown in Eqs. (11) and (12), the theoretical gravimetric capacities of Ba3 and Ba5 are 2.52 wt% and 1.70 wt%, respectively. Based on these theoretical capacities, the cycleaveraged oxygen release efficiency, no2 rel of Ba3 was 49.60 %. The gravimetric O2 uptake and release by Ba5 were much lower than those by Ba₃, partly owing to the less favorable reaction stoichiometry, as shown in reactions (15) and (16), which is largely governed by the Fe content in the ternary oxide. The observed mean mass change of Ba₅ is 0.67 wt% over the 10 redox cycles. Based on Eq. (16), the cycle-averaged oxygen release efficiency $\eta_{O2,rel}$ of Ba_5 is 39.41 %. This low $\eta_{O2,rel},$ as well as the apparently sluggish kinetics of both the O2 uptake and release, as shown in Fig. 9, may be attributed to the ability of Ba₅ to easily take up CO₂ to form BaCO₃ (instead of ternary oxides such as BaFeO₃), which hinders the CLOU activity of Ba₅. Therefore, between the two Ba-rich barium ferrite phases, Ba₃Fe₂O₆ shows more promising CLOU activity at high temperature, making it a potential candidate for applications such as CLOU and chemical looping air separation (CLAS).

3.3.2. Morphological changes over CLOU cycles

Fig. 10(a) & (b) shows the SEM images of the spent Ba_3 and Ba_5 , respectively, over 20 cycles of CLOU between 550 and 950 °C. Interestingly, spent Ba₃ after CLOU experiments showed a surface morphology resembling that of the fresh sample (Fig. 10(a)). The preservation of the pore structure is corroborated by the high BET surface area of the spent Ba_3 of 15.02 m²/g compared to that of the fresh Ba_3 $(8.80 \text{ m}^2/\text{g}, \text{ as shown in Table 3})$. However, for Ba₅, the spent sample exhibits signs of sintering, as characterised by large grains with smooth surfaces (annotated by the solid arrow shown in Fig. 10(b)), which are surrounded by small grains (annotated by the dashed arrow in Fig. 10 (b)). These large, non-porous grains could be assigned to the lowmelting BaCO3 phases, while the smaller grains could be BaFeO3 or Ba₂Fe₂O₅. Overall, a slight decrease in surface area is observed for Ba₅ (12.11 to 10.96 m^2/g), shown in Table 3. The apparent presence of BaCO₃, which are also suggested by the TGA results, indicate that Ba₅ is not a suitable oxygen carrier for CLOU-related applications.

4. Conclusions

 CO_2 looping and CLOU processes rely on the availability of highperformance CO_2 sorbents and oxygen carriers, respectively. In this paper, two barium ferrite samples, namely Ba_3 and Ba_5 with nominal compositions of $Ba_3Fe_2O_6$ and $Ba_5Fe_2O_8$ were investigated for their potential applications in CO_2 looping and CLOU, both of which are process schemes to capture CO_2 using the principle of chemical looping. The two compounds were unexplored for chemical looping applications Table 3

BET surface area and pore size of the fresh and spent samples post oxygen release in TGA.

Sample	Surface area (m ² /g)	Pore size (nm)
Fresh Ba ₃ Fe ₂ O ₆	8.80	8.30
Fresh Ba ₅ Fe ₂ O ₈	12.11	10.96
Spent O2 release Ba3Fe2O6	15.02	5.31
Spent O ₂ release Ba ₅ Fe ₂ O ₈	10.96	13.79

in the literature. Specifically, the crystal structure of ${\rm Ba}_5{\rm Fe}_2{\rm O}_8$ remains unresolved.

Based on the results of TGA experiments, both Ba₃ and Ba₅ have demonstrated satisfactory performance to cyclically capture CO₂ at 900 °C and 1000 °C, forming mixtures of BaCO₃ and BaFe₂O₄ after carbonation. Between the two barium ferrites, Ba₅ shows superior and more stable CO₂ capture performance with average gravimetric CO₂ uptake capacities of 4.37 wt% and 13.39 wt% at 900 °C and 1000 °C, respectively. In comparison, Ba₃ shows gravimetric CO₂ uptake capacities of 7.79 wt% and 11.99 wt% at 900 °C and 1000 °C, respectively. The decay in the cyclic CO₂ capture capacity of Ba₃ was attributed to sintering. Both barium ferrites are only effective in capturing CO₂ above 800 °C. Below 800 °C, the Ba-rich barium ferrites are stable in the form of BaCO₃.

Because of the outstanding ability of Ba₅ to take up CO₂, it does not appear to be a suitable oxygen carrier for CLOU-related applications, which are often conducted in the presence of a high CO₂ partial pressure (e.g., for oxy-fuel combustion or the Allam cycle). Ba₃Fe₂O₆, on the other hand, has shown an oxygen carrying capacity of 1.25 wt%, i.e., 49.60 mol% of the Fe species in Ba3Fe2O6 could undergo reversible redox transitions between Fe⁴⁺ and Fe³⁺ in each cycle, with fast redox kinetics and good cyclic stability. Given that the performance and reactivity of ternary oxides could be modified by modifying the A site and B site occupancies, there is potential to further improve formulation of Ba₃Fe₂O₆ as an oxygen carrier for advanced chemical looping applications such as chemical looping air separation. Therefore, in-depth studies of the oxygen uncoupling (and air separation) properties of Ba₃Fe₂O₆ should be carried out in the future to fully probe and exploit the potential of this ternary oxide system for chemical looping applications.

CRediT authorship contribution statement

Syed Saqline: Conceptualization, Methodology, Investigation, Writing – original draft, Writing – review & editing. **Haiming Wang:**



Fig. 10. SEM images of (a) and (b) - after 20 CLOU cycles.

Conceptualization, Methodology, Investigation, Writing – review & editing. Qianwenhao Fan: Investigation, Writing – review & editing. Felix Donat: Methodology, Investigation, Writing – review & editing. Christoph Müller: Supervision, Writing – review & editing. Wen Liu: Conceptualization, Supervision, Funding acquisition, Writing – review & editing.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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