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Study of the NaOH(s)–CO₂(g) reaction creating value for industry: green natrite production, energy, and its potential in different sustainable scenarios†

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In this work, we present a comprehensive analysis of a solid-based NaOH technology for CO₂ capture. After CO₂ capture, high purity γ -Na₂CO₃ synthetic natrite is formed as demonstrated through exhaustive characterization using several chemical techniques (XRD, XPS, ¹³C and ²³Na MAS-NMR, ATR-FTIR, and TGA), and additionally a DFTB computational analysis was performed. The use of solid NaOH could offer some technical advantages such as high exothermicity (3445 MJ per tCO₂) and the production of high value carbonates (natrite). Furthermore, we consider either carbonate commercialization or regeneration of NaOH in our model. This novel technology has important advantages over currently available technologies regarding efficiency in capture and potential for heat exploitation. Additionally, environmental analysis reveals a negative LCA index (life cycle assessment index) for flue gas capture and the lowest operative costs for CO₂ capture. Two sustainable models are proposed that not only focus on CO₂ capture, but also on the sustainability of time. One model is based on sale of natrite, because the projections in the global market for carbonates will continue to increase, due to their non-traditional applications, e.g., in molten carbonate fuel cells, batteries and other technologies for generation and storage of energy. The second model considered NaOH-raw material regeneration, and pure CO₂ released can be used in obtaining of synthetic fuel, food packaging, and other industrial chemical procedures, and in this respect the current global demand for CO₂ is around 140 Mt per year.

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Environmental significance

Currently, carbon capture is one of the biggest challenges facing humanity. In this sense, it is mandatory to conduct advanced research focused on reducing, capturing and storing CO₂. Several methodologies are available for capturing carbon dioxide, but the great majority of them are not sustainable technologies. Consequently, it is necessary to develop CO₂ capture technologies that, in addition to being viable economically and technically, are sustainable and constitute a real possibility for the industrial and manufacturing sectors. For this reason, we have proposed a new technology that has important techno-economic advantages and as we have shown it is a sustainable technology; in fact, it is simple and strictly follows green chemistry principles. The first great advantage, in comparison with state-of-the-art available methodologies, is that our technology uses “solid-based NaOH”, with the lowest associated costs and very lowest carbon footprint impact and the second advantage is that it produces high-purity synthetic natrite, a product that has several outlets on the global marketplace, as a raw material for various chemical industrial processes and for the production and storage of energy (e.g., MCFs and batteries), stimulating a circular economy model. Here, comprehensive information about the advantages of the solid-based NaOH technology for CO₂ capture is shown and an exhaustive discussion of the environmental, energy and economic impact assessments of this transformative process is presented with promising results.

Introduction

CO₂ capture has been one of the most prevalent topics for the mitigation of climate change. Nowadays, the use of amine solutions is the most popular industrial process for CO₂ capture.¹ Recently, different types of amine processes have been developed, including the biphasic solvent's CO₂ absorption^{2,3} and related new technologies.⁴⁻⁷

However, the amine process has many drawbacks such as high energy consumption for amine recovery, evaporation of the solvent and degradation of the amine or amine-based sorbents.⁸

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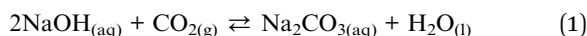
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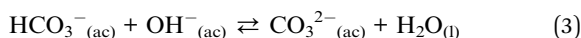


In contrast with the amine solution process, the use of solid hydroxide could offer some technical advantages such as high exothermicity, large reaction velocity and the production of high value carbonates.⁹ Here we perform an exhaustive study of CO₂ capture for a flue gas source using solid NaOH. To the best of the author's knowledge, there is no comprehensive study like this. In this introduction we start with the motivation for the present work.

Classical CO₂ capture can be carried out in alkaline aqueous solutions,¹ and it can be overlooked in eqn (1) in a liquid/gas medium ignoring the solid/gas possibility.



Eqn (1) is much more complex than is normally considered in basic chemistry,¹⁰ especially considering the equilibrium of the sodium solvated ions $[\text{Na}(\text{OH}_2)_x]^+$ _(aq).¹¹ On the other hand, it is necessary to consider the equilibrium of CO₂ in alkaline solutions (eqn (2) and (3)).¹



Clearly, the outcome of eqn (1) is combining the results of the carbonate equilibrium in an alkaline medium. Depending on the reactive conditions, in the solid state, it is possible to obtain not only sodium carbonate anhydrous, but other well-known forms of sodium carbonate hydrate such as thermonatrite, Na₂CO₃·H₂O;¹² natron, Na₂CO₃·10H₂O,¹³ and carbonate combinations such as Wegscheider salt, Na₂CO₃·3NaHCO₃ or trone, Na₂CO₃·3NaHCO₃·2H₂O as well.¹⁴ These products have been reported in the dry carbonation process for CO₂ capture (DCP).¹⁵

At a solid/gas interface,¹⁶ the process does not involve any solvation energy. Instead, it is driven by the NaOH lattice energy $U_{\text{NaOH}} = 550 \text{ kcal mol}^{-1}$.¹⁷ Consequently, the reaction with NaOH_(s) is a heterogeneous process, so the reaction may present small deviations from eqn (1) stoichiometry because sodium carbonate presents a sodium deficiency, previously reported by the XPS¹⁸ study, a medium that accumulates superficially by "nucleation". This mechanism has been previously proposed for alkali and alkaline earth metal oxides used as absorbers in CO₂ capture.¹⁹ The reaction, which is spontaneous ($\Delta G^\circ < 0$), is also exothermic due to a favourable balance between the U_{NaOH} and $U_{\text{Na}_2\text{CO}_3}$ lattice energies and will continue capturing CO_{2(g)} and generating anhydrous Na₂CO_{3(s)} until all the nucleophilic agents are consumed.

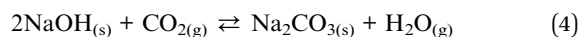
Physical adsorption of CO₂ requires about 324 kJ kg⁻¹ CO₂ with equal pressure in the feed and product streams, which is by far smaller than chemisorption, that requires about 1224 kJ kg⁻¹ CO₂.²⁰ Then, solid sorbents offer both the possibility of a lower cost, not only in energy, but also in operating costs, and application of a wide range of scales. The reaction mechanisms between a gas (CO₂) and a solid (metal hydroxides and oxides) are not very easy to discern and the reaction products analysed by X-ray diffraction vary depending on the temperature and the pressure at which they occur.²¹

The mechanistic and kinetic studies of the transformation of carbon dioxide in water and dilute aqueous solutions of sodium hydroxide have been studied since the beginning of the 20th century.^{22–27} Concisely, hydroxides have been used for direct CO₂ air capture with a variety of methods.^{28,29} Nikulshina *et al.*,³⁰ reported three Na-based thermochemical cycles for capturing CO₂ from a gas source of 500 ppm of CO₂, including the use of solid NaOH. However, they showed that with their experimental conditions, it is very difficult for the application of Na-based solid sorbents for capturing CO₂ from air. Furthermore, it is important to add that during the processes studied so far, carbonate and bicarbonate mixtures are produced.

In the present experiment, the molecular study of the reaction product between CO_{2(g)} and NaOH_(s) is performed, where an attempt is made to explain the formation of anhydrous form $\gamma\text{-Na}_2\text{CO}_{3(\text{s})}$ (synthetic natrite) with an incommensurately modulated structure and its sustainable advantages in a circular economy. The next section is divided into two parts: first, detailed mass, energy and chemical characterization of the solid-based process is presented. Second, the competitive advantages are presented at a technological level, in relation to favourable mass and energy balances that positively impact environmental (carbon footprint and LCA) and economic (techno-economic evaluation) sustainability during the CO₂ capture process.

Results and discussion

The amount of CO₂ captured (mean value in 28 experiments) was $1.941 \pm 0.057 \text{ g}$ of CO₂ captured using 3 g of NaOH. We consider the NaOH purity to obtain an accurate ratio between CO₂ moles per mol of NaOH. Based on the amount of hydrated NaOH and the scale factor (X-ray analysis) a purity of 92.5% is assumed. After considering the NaOH purity, a ratio of 0.66 CO₂ moles per mole of NaOH is obtained. The large percent of CO₂ capture per NaOH can be explained due to the increase in the amount of carbonate by sodium in the observed product. With this experimental evidence, the ratio between the moles of CO₂ (or Na₂CO₃) and moles of NaOH was found to be larger than 0.50, without any evidence of NaHCO₃ formation. A theoretical value of 127.6 kJ mol⁻¹ of CO₂ is predicted for the heat of reaction between solid NaOH and CO₂ (eqn (4)).



X-ray powder diffraction characterization

Fig. 1 shows the X-ray powder diffraction (XRD) pattern corresponding to a sample of the material obtained from the CO₂ capture process using the solid-based NaOH technology.¹⁶ Clearly, the sample is mainly crystalline and there is no evidence of the presence of non-crystalline/amorphous phases, containing only the crystalline phase $\gamma\text{-Na}_2\text{CO}_3$ natrite,³¹ monoclinic space group *C2/m* structure. This phase is an incommensurately modulated structure.³² Fig. 1a depicts the obtained X-ray powder diffraction diagram in the main angular



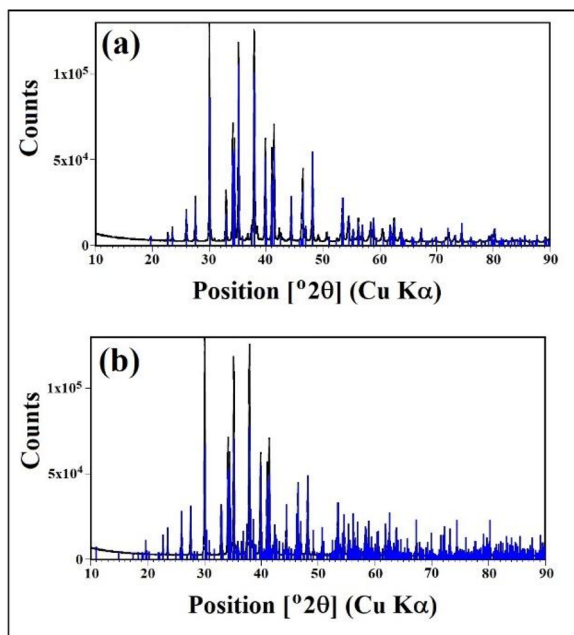


Fig. 1 X-ray powder diffraction (XRD) pattern corresponding to a γ - Na_2CO_3 natrite, monoclinic space group $C2/m$ structure, obtained by chemical transformation with solid-based NaOH technology for CO_2 capture. (a) and (b) Diagram with two patterns of γ - $\text{Na}_2(\text{CO}_3)$ natrite, PDF#4-11-4108 and PDF#0-61-1420 respectively.

range from 10° to 90° 2θ , with the intensity corrected to fixed 1° divergence slit values, with a main structure pattern of γ - Na_2CO_3 natrite (PDF#4-11-4108), with the reflections of the main structure superimposed. Fig. 1b depicts the same diagram with another pattern of γ - Na_2CO_3 natrite (PDF#0-61-1420), with additional satellite reflections superimposed.

X-ray photoelectron spectroscopy analysis

The XPS spectra (see ESI, Fig. S1a†) show the expected lines for the constituent's elements in the sample previously identified by DRX as natrite, an anhydrous form of sodium carbonate, γ - Na_2CO_3 . However, a peak corresponding to silicon was observed, a plausible impurity resulting from the material manipulation technique. The C_{1s} spectral region (see ESI, Fig. S1b†) contains two lines at around 290 eV and 284 eV respectively. The literature reported one line centered at 289.3 eV corresponding to γ - Na_2CO_3 .³³ In this case, the second line to lower BE is discardable as an Auger peak, satellite peak, adventitious carbon³⁴ (C_xH_y) or signal of the H-CO group by the sodium bicarbonate impurity, previously reported at 289.6 eV. Consequently, it is possible to propose a C_{1s} high-resolution signal splitting into a doublet, attributable to more than one type of C-O bonding present in the crystalline structure of natrite. Multiplet splitting was observed in other carbonaceous materials as an additional sp^3 C peak in the C_{1s} XPS spectra region.³⁵ In this instance, an extreme situation is possible for carbonate ions present in the natrite structure. First, a carbon bond with a strong sp^3 component. Second, a carbon atom with a strong sp^2 component, as shown in the X-ray structure.³⁶ In

a nutshell, two XPS C_{1s} peaks are observed: an sp^2 C peak with high binding energy and an sp^3 C peak with low BE. The O_{1s} spectral region for natrite (see ESI, Fig. S1c†) contains one distinguishable oxygen peak centered at 530.5 eV, previously reported at 531.8 eV.³³ Additionally, we see the Auger lines NaKLL1 (a3) at 537 eV and NaKLL (a4) at 531 eV. The characteristic form of the O_{1s} peak indicates that it can be decomposed into various components, a typical XPS signal behaviour in the presence of different oxygen bondings. This result is consistent with that observed for the C_{1s} line, demonstrating more than one type of C-O bonding present in the incommensurately modulated X-ray structure of natrite.³² 1071.5 eV BE was observed at the Na_{1s} peak (see ESI, Fig. S1d†) corresponding to sodium atoms with octahedral coordination present in the natrite structure; however, a minoritarian signal at around 1070 eV suggests another type of Na atom too. The X-ray structure reported for natritic sodium carbonate, with graphite-like layers formed by three types of Na^+ and CO_3^{2-} ions³⁷ and a distorted-octahedral coordination sphere around the sodium atom, is in accordance with the observed Na_{1s} signal.

Differential thermal and thermogravimetric analysis

Differential thermal and thermogravimetric analysis (see ESI, Fig. S2a†) carried out on the γ - Na_2CO_3 sample shows a loss of 0.0776% ubicated at 114.93 °C, associated with water in the lattice. Also observed were two endothermic peaks at around 232.97 °C and 337.178 °C, with losses of 0.1410% and 0.2360% respectively. These values are below the temperature range of the phase transition $\gamma \rightarrow \beta$, but this can be attributed to carbon species (C_xH_y), previously reported in the TGA-DSC-MS experiment on a similar natrite sample,¹⁶ yet cannot be discarded as a polymorphism effect. The thermogram showed an endothermic steady fall from 200 °C to 700 °C, a range where we note the three phase transitions γ -, β - and α - Na_2CO_3 .³⁶ In fact, the γ -phase is stable until 361 °C, β -phase at around 361 °C and 489 °C, and α - Na_2CO_3 above 489 °C to the melting point. Finally, it was observed that the γ - Na_2CO_3 melting point was 851 °C, a 0.28% difference in respect to 854 °C, a value reported in the literature and corresponding to pure sodium carbonate or natrite.³² The thermogravimetric analysis result indicates that sodium carbonate obtained by chemical transformation with solid-based NaOH technology for CO_2 capture has a purity level of 99.55%.

ATR-FTIR analysis

The infrared spectrum with attenuated total reflectance (see ESI, Fig. S2b†) showed seven bands from 4000 to 400 cm^{-1} , similarly, to that observed in the infrared spectrum of γ - Na_2CO_3 and the natrite.³⁸ No signal in the 3500 to 3000 cm^{-1} region indicates an absence of water molecules, discarding the presence of thermonatrite, a typical impurity in natrite samples. In order to carry out a detailed study of the spectrum, an ideal XY_3 system is considered with D_{3h} symmetry. The application of the selection rules shows that there are four vibrations expected in these types of molecules: one nondegenerate (A'_1) vibration, two



doubly degenerate (E') vibrations, and one out-of-plane (A_2'') vibration. Consequently, the carbonate ion exhibits four normal vibrations:³⁹ ν_1 (A_1'), symmetric C–O stretching mode; ν_2 (A_2''), CO_3^{2-} out-of-plane deformation mode; ν_3 (E'), asymmetric C–O stretching mode and ν_4 (E'), O–C–O bending in-plane deformation mode. Furthermore, in the ATR-FTIR spectrum of the sample, a weak band around the 702–695 cm^{-1} region was assigned to ν_4 vibrational mode. The medium intensity band near the 877–854 cm^{-1} region must be due to ν_2 normal mode. A strong broad band around the 1418–1409 cm^{-1} region corresponds to ν_3 vibrational mode. The ν_1 (A_1') vibrational mode is inactive in infrared according to selection rules; however, a distortion of the punctual symmetry from carbonate ions, justified on the basis of the structure of natrite obtained by X-ray data, indicates a possible relaxation of the selection rule mechanism, and therefore now it is possible to observe ν_1 (A_1') mode as a very weak shoulder at 1079 cm^{-1} . Previously, this ν_1 mode was reported at 1063 cm^{-1} and 1056 cm^{-1} , in the γ - Na_2CO_3 and natrite spectrum,^{38,40} respectively, using a traditional infrared technique. The comparative analysis of the four vibrational modes observed in the spectrum indicates a reduction of the D_{3h} ideal symmetry in the carbonate ion, a consequence of position modulations of carbon and oxygen atom in phase γ into an incommensurately modulated structure of natrite.³⁷ This observation would justify the appearance of more than one component in each vibrational mode band.

²³Na and ¹³C MAS-NMR analysis

The ²³Na MAS-NMR spectrum (see ESI, Fig. S2c†) presented two distinct signals, at 3.15 ppm, a singlet and around –15.85 to –23.47, a broad multiplet. These signals indicate the unambiguous existence of more than one type of sodium atom, or preferably, different coordination symmetries in the lattice of the sodium atom. The X-ray structure reported for natrite showed two octahedral sodium atoms, ³²Na₁ and Na₂, occupying interstitial sites with a highly symmetric coordination sphere in the natritic structure, a position that gives a 3.4 ppm signal. A third-sodium type (Na₃) and carbonate ions form graphite-like layers, parallel to the [001] direction³⁶ in a position with lower symmetry and surrounded by six oxygen atoms in distorted equatorial coordination with one oxygen atom above and two equidistant oxygen atoms below this plane. These types of atoms, in positions of low symmetry, appear at lower fields consequently,^{41,42} and the Na₃ atom is observed as a broad signal below –15.85 ppm.

The ¹³C MAS-NMR spectrum showed a singlet at 172.19 ppm, typical of the carbonyl group of the carbonate ion.⁴³ The signal amplification into the 170–175 ppm region indicates unsymmetric slight broadening due to a variation of the CO_3 -group geometry coherent with X-ray details observed in the incommensurately modulated structure and with ATR-FTIR analysis which indicates a reduction of the D_{3h} ideal symmetry of the carbonate ion in natritic chemical environments.

Clearly, the ²³Na, ¹³C MAS-NMR spectrum according to X-ray data and XPS suggests a sodium deficiency especially on the natrite surface. This means a deviation in the ideal sodium-to-

carbonate ratio; in fact, this value was previously reported as 1.92 in bulk³⁸ and 1.75³³ and 1.70 on the surface.¹⁸ Normally, stoichiometric calculations that involve sodium carbonate with a value of 2 are normal, and a value between 1.92–1.70 is possible by approximation to 2, especially in fine chemicals. But, in mass balance for industrial chemistry where tons of raw materials are present, this deviation must be considered.

DFTB calculations

To compare different structures again, we generated the average values for the most important geometric parameters. From Table S1 (see ESI†), we can conclude: (1) the stability of the Na_2CO_3 structures increases when the C–O distance increases and the Na–C distance decreases. This keeps the number of neighbourhoods relatively constant. (2) The non-planarity of the CO_3^{2-} fragment stabilizes the structure. This non-planarity is necessary for the observed decrease in the Na–C distance. (3) The charge on the sodium atom decreases as the bond is stronger. (4) These results show that possible vacancies of Na (perhaps on the surfaces) can stabilize the Na_2CO_3 surface.

Life cycle assessment

The regeneration of NaOH is modelled according to three different methods: (i) electrolysis of sodium carbonate. The energy cost of this technology is similar to that of the industrial

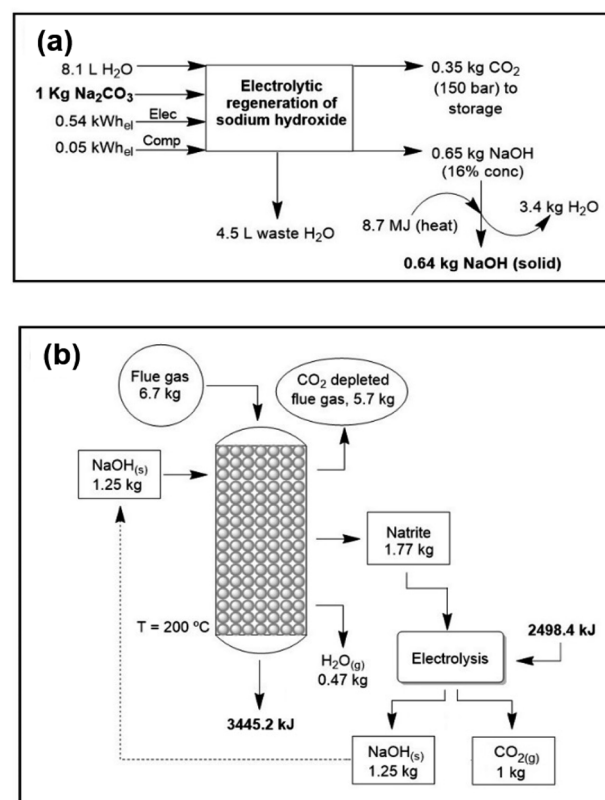


Fig. 2 (a) Electrolytic regeneration of sodium hydroxide and pure CO₂ production.⁴⁴ (b) Mass and energy balance of the complete cycle with NaOH-solid technology. In this figure the complete regeneration of natrite is studied.



“chloralkali” process for the production of NaOH from aqueous NaCl solutions, that is, about 4500–5500 MJ per ton of NaOH, without considering water evaporation. In this method,⁴⁴ pure CO₂ is produced and sodium hydroxide is recycled back into the process through an electrolytic cell with hydrogen recycling (Fig. 2a). A theoretical maximum concentration of sodium hydroxide of 16% (w/w) is achievable due to the solubility limitations of sodium carbonate. The voltage to be applied is that of water electrolysis, slightly less than 2 V per cell. The achieved current efficiency is 85% (see Methods). If we exclude water evaporation this process is near the thermodynamics minimum. (ii) Lime cycle. Conversion of sodium carbonate into NaOH, also called “causticization”, is one of the oldest processes in the chemical industry. This process has been used by the paper industry for over a century to recover NaOH from aqueous solutions of sodium carbonate and a lignocellulosic material, known in the industry as “black liquor”. It consists of treating the sodium carbonate solution with lime (Ca(OH)₂) and precipitating CaCO₃. Subsequently, calcium hydroxide is regenerated by heating calcium carbonate to 900 °C. Some studies on CO₂ air capture with aqueous NaOH have focused on this regeneration process.^{45–47} (iii) Direct causticization using titanates. Unlike the lime cycle, this process requires solid carbonate which is subsequently heated with sodium trititanate to form sodium pentatitanate and CO₂. Sodium pentatitanate in water spontaneously produces NaOH and regenerates trititanate. Heating up to 900 °C of sodium trititanate and carbonate is the energy consuming step in this process. The titanate process requires 50% less energy than the lime process and 50% less water evaporation than electrolytic recovery.

From the regeneration point of view, the less energy intensive per CO₂ ton is the electrolysis (around 40 euros per CO₂ ton); however, the cost of water evaporation is high; in contrast, heating in direct caustification is costly but since the final solution is more concentrated the evaporation cost is much lower. Finally, even if costly, the lime cycle is the most mature and well-known of all technologies. Depending on water evaporation and the temperature the energetic cost of electrolysis is between 44 and 120 euros. However, due to less water evaporation the cost of using titanates can be well below that of electrolysis.

At this stage a full cost benefit analysis would be speculative due to the uncertainty in parts of the investment costs and operating costs; for this reason the analysis is focused on the energetic cost and carbon footprint.

Mass and energy balance

The final energy and mass balances of the NaOH system with flue gas are represented in Fig. 2b. Considering this, the functional unit (*i.e.*, the unit to which the process inventory is referred to) selected was 1 kg of captured CO₂ (*i.e.*, 1 kg CO₂ at the point of capture). As mentioned before, the last part (regeneration of NaOH) can be performed using an electrochemical cell. In addition to the large energy gain, solid-based processes achieve a large reaction rate when compared with the reported rate for other technologies.

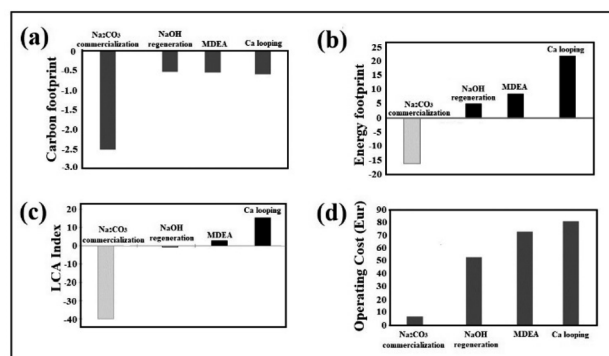


Fig. 3 (a) Carbon footprint (kg CO₂ eq. per kg CO₂ captured). (b) Energy footprint (MJ per kg CO₂ captured). (c) LCA index considering MDEA technologies, looping carbonate and NaOH(solid)-based solid technology. Both scenarios are taken into consideration: full Na₂CO₃ commercialization and full NaOH regeneration without the sale of natrite. (d) Operating cost per ton of CO₂ capture.

Carbon footprint (CF)

The CF assessment of the process not including NaOH regeneration (Fig. 3a) is -2.51 using green NaOH (see Methods) and -1.66 using the commercial one. When NaOH recovery is performed, the source of NaOH is less important, but the energy source for heating and the efficiency of the heat transfer play a major role. Assuming a conservative 50% energy recovery in the reaction the CF is estimated to be -0.57 kg CO₂ for electrolytic recovery. Energy recovery in the form of electricity can be achieved using an organic Rankine cycle (ORC) using temperatures below 200 °C with high efficiency. Therefore, the assumption of a 50% heat is very reasonable in this scenario of low temperature. Under our assumptions, the CF is negative in all scenarios. Thus, there is a net decrease in greenhouse gas emissions in the atmosphere (see ESI, Table S2†).

Energy footprint (EF)

This environmental category focuses on the balance of primary non-renewable energy consumption. Hence, an EF is the assessment of the energy consumption related to the whole supply chain and processing of the captured CO₂. The results observed for the net balance of fossil resources are presented in Fig. 3b. The net values, in this case, are always positive, except for the scenario where CO₂ is captured and sequestered in Na₂CO₃ for sale in the carbonate market. The other options involve an increase in the consumption of fossil fuel resources: air-based processes produce a large depletion of fossil resources, while for flue gas, our NaOH(s)-regeneration process produces the lowest impact. Indirect calcium looping increases the demand for fossil resources with respect to amines; as a matter of fact, all the alkali regeneration approaches assessed in this study (electrolytic or indirect Ca-looping) have significant energy footprints (see ESI, Table S3†).

LCA index

As shown in Fig. 3c, it is possible to quantitatively appreciate the significant impact in relation to the carbon and energy



footprints of the three capture technologies. Competitive advantages are not seen in a conclusive way in the previous graphs where both variables were evaluated separately.

Techno-economic assessment

Fig. 3d shows the competitive advantage in relation to operating costs of the technology using NaOH_(s), being the most favoured when natrite (anhydrous high purity Na₂CO₃) is sold at the price of conventional industrial carbonate. However, as the global carbonate market is relatively small, almost all of it must be regenerated and converted to NaOH (feedstock). 99% of raw material regeneration is considered in this study, which causes higher CO₂-capture operating costs. Considering this additional process, the operating cost of the NaOH technology is still significantly lower (52 Eur per tCO₂) than the operating costs of the technologies considered (≥ 73 Eur per tCO₂) (see ESI, Table S4†).

Sustainable model I

Sale of natrite and use of energy

The natrite produced with our technology represents an added value that stimulates a sustainable economy by providing value to CO₂ in an economically viable way for several other industrial applications (see ESI, Fig. S3†). Using natrite would provide environmental and economic sustainability for a relatively lower cost, high purity, green carbonate industry. On the other hand, the projections in the global market for carbonates will continue to increase, among other aspects, due to their application in the technology of molten carbonate fuel cells (MCFCs),⁴⁸ which require raw materials of high purity to produce clean energy.

These MCFCs are a growing technology, with global potential in evaluated pilots, that meet energy needs efficiently and with great prospects in the residential, commercial, and shipping industries and capture of CO₂ from industrial emissions, among other areas of energy application. Although the market for anhydrous green carbonate and molten carbonate fuel cells is not intensive, international pressures for the use of more environmentally friendly alternatives are increasingly driving these emerging markets, as is happening in maritime transport. For example, Fig. S3 (see the ESI†) shows an important release of energy (3445 MJ per tCO₂ captured) which corresponds to energy released in the form of heat and can be used to feed an organic Rankine cycle (ORC) or the conventional Rankine cycle. Due to the high temperatures produced, obtaining this increase in the production of electricity or even heating water represents additional benefits to the global economic balance.

Sustainable model II

Raw material regeneration

Excess natrite can be processed to regenerate chemical converters (NaOH) which would start the capture cycle all over again with a lower carbon footprint at each regeneration step. The separated pure CO₂ can be injected into geological sinks

and a part could be used for enhanced oil recovery (EOR), food packaging, and synthetic fuel synthesis, among other uses. Current global demand for CO₂ is around 140 Mt per year, with the oil industry as the main consumer.

This regeneration has the advantage that both carbonate and solid hydroxide are stable under industrial operating conditions and easy to handle, which allows the use of existing transport infrastructure for a lower cost of transporting CO₂ to regenerating locations that may be advantageous for storage or use. There may also be potential benefits to compensating the costs of the energy consumed in the raw material regeneration (NaOH) with the energy released during CO₂ capture. In other words, in this model, industry has alternative evaluation: (1) if the benefits from the sale of high purity Na₂CO₃ are attractive, (2) which in turn would give benefits in energy savings during the regeneration process by electrolysis and (3) in turn an “additional advantage” that involves the use of energy that occurs during the capture process. These three aspects must be considered for the global economic balance of the system, which will have a direct impact on the improvement of the estimated operating costs.

In the same way, achievable savings in costs associated with CO₂ allowance units (≈ 80 Eur per tCO₂ in EU/ETS) would be an interesting incentive for industrial operations in Europe.

Conclusions

(1) The system of CO₂ capture using NaOH in a solid state is a robust, efficient, and flexible technology and very environmentally friendly and additionally can be considered a green chemical technology. By means of XRD, XPS, DSC-TGA, ATR-FTIR, and ²³Na and ¹³C MAS-NMR it was possible to confirm the sodium carbonate anhydrous production with high purity (>99.55%), present in the form of γ -Na₂CO₃ (synthetic natrite) with a sodium deficiency especially on the surface. It is important to highlight that in stoichiometric calculations with sodium carbonate, a value of 2 is normal and a value of 1.92–1.70 is possible to find by approximation to 2, which are valid conditions in fine chemicals, but in mass balance for industrial chemistry that involves tons of raw materials, this sodium deficiency must necessarily be considered. All results obtained in the structural characterization of natrite are consistent with those exhibited in the DFT calculations and with the X-ray results previously reported, where an incommensurately modulated structure was described.

(2) The techno-economic assessment considered two sustainable models; ‘model I’ is based on sale of natrite. Furthermore, the release of energy in the form of heat during the captured process, 3445 MJ per tCO₂, can be used for electric energy production. Sustainable ‘model II’ considered the NaOH-raw material regeneration and the pure CO₂ obtained in the process, which can be used in other industrial applications.

(3) High exothermicity of the reaction allows for taking the energy released during the capture and making use of it in the regeneration process. This has benefits for the sustainability of the global process. Additionally, even if it is not strictly necessary for it to be a chemical conversion and regeneration, this



technology is meaningfully more favorable than those offered by the technology market. Concisely, a balance that considers the sale, completed regeneration or partial regeneration of nitrite, to which we may add the benefits in the carbon allowance markets, would give this technology competitive advantages over traditional CO₂ capture technologies.

Methods

Gas emission concentration

The gas concentration was determined using an analyser type HORIBA Multi-Component Gas Analyzer/VA-5112G, consisting of a non-dispersive infrared method of a double beam (NDIR-nondispersive infrared) to measure gases such as SO₂, CO, CO₂, CH₄, N₂O, NO, and NH₃) and the chemiluminescence method (CLA) that allows measuring NO_x at low concentrations. The instrumentation bench also has a mixer EnviroNics gas dilution system/Series 4040, a sample conditioning M&C gas conditioning unit with temperature controller/CSS Series, two flow controller systems precision gas mass flow controller/MCS-Series and M-Series, a differential pressure transducer OMEGA differential pressure transmitter/Model PX2300 and a trace detector of humidity in gases based on a P₂O₅ sensor DKS Aquatrace IV/Typ-R. These instruments were operated using CO₂ REMO software from TCA Spain.

Chemicals

Sodium hydroxide was purchased from Gie-Tec GmbH (Eiterfeld, Germany) and Merck KGaA, Darmstadt, Germany. Industrial NaOH was also purchased from Barcelona Global Chemical Solutions (Cornella de Llobregat – Spain). Sodium carbonate was obtained from PanReac AppliChem (Darmstadt, Germany). The gases CO₂ and N₂ were purchased from Air Liquid (Madrid, Spain). All chemicals were used directly without further purification.

Experimental capture system

We employ a laboratory scale model converter to test the amount of CO₂ captured using a solid NaOH chemical converter. A 50 mL Falcon conical tube was packed with three layers of solid NaOH, 1.00 g each. The system was purged using N₂ to remove any traces of remaining gases. Following this 5% CO₂ was captured in three steps at room temperature (≈ 20 °C). The solid component was recovered after each step and ground up to pack the Falcon tube with 2 layers (step 2) or 1 layer (step 3) of 1.00 g chemical converter. The captured CO₂ was determined by using an instrumental method (see ESI, Fig. S4†).

CO₂ determination method

The amount of CO₂ captured was obtained following the concentration of CO₂ at the end of the reactor until the system was saturated and no more CO₂ was adsorbed in each capture step. From the gas flow (1.3 L min⁻¹), we obtained the volume of CO₂ captured. From this volume, we calculated the moles of CO₂. The typical error in moles of CO₂ from this method is

about 1%. This error was estimated assuming a 1% in the CO₂ profile (see ESI, Fig. S5†).

Reproducibility and repeatability of studies

For this evaluation, 4 new NaOH flasks were considered, from different provider batches (Gietec (Germany), Merck (Germany) and Barcelonesa Group (Spain)) and experiments were carried out independently for each sample for a total of 28 trials. To estimate the purity of NaOH, we use information from the X-ray analysis of fresh NaOH based on the amount of hydrated NaOH and the scale factor.

The power X-ray diffraction and XPS studies

X-ray powder diffraction characterization was carried out using a PANalytical X'Pert PRO MPD apha1 powder diffractometer in Bragg–Brentano $\theta/2\theta$ geometry of 240 millimeters of radius. A focalizing Ge (111) primary monochromator was used with Cu K α 1 radiation: $\lambda = 1.5406$ Å and work power: 45 kV to 40 mA. A variable divergence slit aperture was used to get an irradiated length over the sample of 10 millimeters. A mask defining the length of the beam over the sample in the axial direction of 12 millimeters was used. A diffracted beam with 0.04 radians Soller slits was used. Sample spinning was performed at 2 revolutions per second. A pixel detector was used; active length = $3.247^\circ / 2\theta$ scans from 4 to 100° 2 θ with a step size of 0.026° and measuring time of 100 seconds per step (six repeated consecutive scans). Qualitative crystalline phase analysis was performed by means of the PDF (Powder Diffraction File) database, ICDD-JCPDS (International Center for Diffraction Data – Joint Committee of Powder Diffraction Standards, 2021).

X-ray photoelectron spectroscopy

XPS was carried out using a Physical Electronics spectrometer, model PHI 5500. The experiment covers all elements of the periodic table, except hydrogen and helium. The depth of analysis <10 nm was performed in a chamber of ultra-high vacuum (UHV) working in the range of 1×10^{-8} to 1×10^{-9} torr (from 1.3×10^{-6} to 1.3×10^{-7} Pa) pressure. The system was calibrated in accordance with PHI procedures, with photo-emission lines Eb of Ag_{3d5/2} = 368.3 eV and with 0.8 eV of amplitude at half peak height (FWHM), using the monochromatized aluminium source for excitation.

Differential thermal and thermogravimetric analysis

DSC-TGA was performed simultaneously with an SDT Q600 V20.9 Build 20 at a heating rate of 5.00 °C min⁻¹ in the temperature range 33–1100 °C. Approximately 15 mg of the sample was used in each run.

ATR-FTIR analysis

The infrared spectrum has been obtained using a Thermo IZ10 instrument with an attenuated total reflectance accessory (ATR) of a diamond. The instrument has a DTGS detector, a KBr beam-splitter and a glow-bar source. The spectral range analysed has



been 4000 to 525 cm^{-1} with a 4 cm^{-1} resolution and 32 accumulations.

Nuclear magnetic resonance

Solid-state magic angle spinning (MAS) NMR spectroscopy experiments were performed on a Bruker AV 400 MHz WB (89 mm; 9.39 T) spectrometer at spinning speeds of 12 kHz at a ^{13}C frequency of 100.61 MHz and referenced to glycine CO at 176.03 ppm and ^{23}Na frequency of 105.84 MHz and referenced to NaCl at 7.28 ppm. The ^{13}C direct excitation experiments were carried out with a 90° pulse with a length of 4 μs averaging 256 scans with a recycle delay of 300 s and 600 s in the MAS NMR sequence. The ^{23}Na direct excitation experiments were performed using a 90° pulse with a length of 4 μs and a recycle delay of 5 s in the MAS NMR sequence.

The DRX, XPS, TGA-DSC, IR-ATR and elemental analysis (% C and % Na) experiments were carried out in Centres Científics i Tecnològics (CCiT), Universitat de Barcelona, Spain. The NMR spectra were obtained at the Unidad de Resonancia Magnética, Facultad de Ciencias Químicas, Universidad Complutense de Madrid, Spain.

DFTB studies

We performed non-spin polarized density functional tight binding (DFTB) calculations with the dftb+ package. The bulk Brillouin zones were sampled with $1 \times 2 \times 2$ and gamma point Monkhorst-Pack k -point meshes. Self-consistent field calculations were converged to 1×10^{-5} eV and geometries were optimized using a force converge of 0.01 eV \AA^{-1} . We initially optimized the atomic positions and lattice constants starting from the reported γ - Na_2CO_3 bulk parameters ($a = 8.920$ \AA , $b = 5.245$ \AA , and $c = 6.050$ \AA). The structure of Na_2CO_3 used as a starting point can be described as graphite-like layers formed by one Na^+ for CO_3^{2-} ions, which are stacked along the hexagonal axis. An additional Na^+ ion is in the hexagonal interchannels. The optimized Na_2CO_3 bulk structure was further revised to better represent the real material which is incommensurable. We generated 20 additional candidate starting structures by randomly perturbing the locations of Na^+ ions. The 20 revised bulk structures are divided into three groups according to their relative energy differences. Group 1 contains structures that relax back to the same geometry as the initial structure. The optimized structures from these groups are all energetically more favourable than the initial structure. Group 2 consists of structures with energies lower than by about 0.5–1 kcal per formula unit. Group 3 structures are lower in energy than the initial structure by about 1–2 kcal per formula unit. The existence of several different structures that are nearly isoenergetic means that the bulk material likely consists of domains having different orientations of the carbonate ions at room temperature and above. The theoretical DFTB Na_2CO_3 structure results are similar to those obtained using DFT-PBE.⁴⁹

Mass energy balance

In this process, pure CO_2 is produced and NaOH is recycled back into the process through an electrolytic cell with hydrogen

recycling. This process was originally patented to produce NaOH without the production of chlorine. A theoretical maximum concentration of sodium hydroxide of 16% w/w is achievable due to the solubility limitations of sodium carbonate. The voltage to be applied is that of water electrolysis, slightly less than 2 V per cell.⁴⁸ The reported current efficiency is 85%.

Green sodium hydroxide assumes the use of renewable energy in the production of low-impact sodium hydroxide. The basic concept of green sodium hydroxide is the substitution of grid electricity with a renewable source (see ESI, Tables S2–S4†).

Life cycle assessment impact

In this part, two reference systems have been studied: absorption of CO_2 with amines and indirect calcium looping using NaOH. Traditionally, monoethanolamine (MEA) has been the benchmark solvent for this application. Here, we consider the use of methyldiethanolamine (MDEA), which, when compared to MEA, presents a higher maximum CO_2 loading capacity (1.00 mol CO_2 mol⁻¹ MDEA vs. 0.50 mol CO_2 mol⁻¹ MEA). Furthermore, thermal regeneration, which has a high energy demand and accounts for 80% of the operating cost for MEA-based capture, can be reduced if MDEA is chosen instead. The energy demand for MEA and MDEA regeneration is experimentally determined. They respectively reported 4.4 GJ per t CO_2 captured and 2 GJ per t CO_2 captured for MEA and MDEA.⁵⁰

The capture of CO_2 from air is based on indirect calcium looping and causticization.⁹ The original publication assumed KOH for the cycle, and NaOH has been analyzed in this work. Since the process utilizes an aqueous solution of NaOH and further includes its regeneration through a thermal route using CaCO_3 produced in a causticization reactor, it can be compared to other options for NaOH regeneration. In total, five scenarios for solid NaOH technology for carbon capture are considered based on the final use of the technology and the source of NaOH: (1) commercialization of Na_2CO_3 from commercial NaOH; (2) commercialization of Na_2CO_3 from green NaOH; (3) regeneration of NaOH by electrolysis and CO_2 storage from green NaOH.

Carbon footprint

The CF is the total greenhouse gas emissions caused by the process, expressed as kg CO_2 equivalents, characterized through the factors for chemical species of the IPCC (Intergovernmental Panel on Climate Change). If no NaOH regeneration is employed, the production of NaOH is likely to be the main contributor to the CF of the present technology. In this case, two sources of sodium hydroxide can be considered: commercial NaOH and green NaOH. For green NaOH, it was assumed that wind power is used instead of electricity from the grid. Also, the utilization of Na_2CO_3 based products such as sodium carbonate, reduces the carbon footprint of our technology, as it replaces the carbon footprint of current mining and chemical manufacturing processes of high-purity carbonates. For the carbon footprint, we estimated that a value of 0.50 kg CO_2 eq. per kg of NaOH was used for green sodium hydroxide and 0.83



kg CO₂ eq. per kg of NaOH for commercial sodium hydroxide.⁵⁰ The production of green Na₂CO₃ has an impact of −1.01 kg of CO₂ per kg of Na₂CO₃ produced, as it is avoided.

LCA index

It is often convenient to combine both parameters, carbon and energy footprints, into a single score, in particular, if one needs a simple way to assess several CO₂ capture technologies. Here, we integrated both values into a single equation (LCA index). This index allows us to appreciate, through a quantitative value, the competitive advantages of any technology in relation to the environmental impact of carbon dioxide capture. Since the energy footprint is an order of magnitude larger than the carbon footprint, the index is calculated as $LCA_{\text{index}} = EF + 10 CF$.

Mass energy balance

The composition of the flue gas was 14.90% CO₂, 80.50% N₂, and 3.60% O₂. This balance considers two steps: (1) a fluid-bed reactor of solid NaOH as the chemical sorbent of CO₂, a 100% conversion reaction of NaOH can be assumed; (2) NaOH regeneration and the CO₂ concentration.

Conflicts of interest

There are no conflicts to declare.

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