Integration of thermochemical water splitting with CO₂ direct air capture

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Abundant production of fuels and chemicals from direct air capture (DAC) of CO₂ is a highly desired goal. Here, we report the integration of the DAC of CO₂ with the thermochemical splitting of water to produce CO₂, H₂, O₂, and electricity. The produced CO₂ and H₂ can be converted to value-added chemicals via existing technologies. The integrated process uses thermal solar energy as the only energy input and has the potential to provide the dual benefits of combating anthropogenic climate change while creating renewable chemicals. A sodium–manganese–carbonate (Mn–Na–CO₂) thermochemical water-splitting cycle that simultaneously drives renewable H₂ production and DAC of CO₂ is demonstrated. An integrated reactor is designed and fabricated to conduct all steps of the thermochemical water-splitting cycle that produces close to stoichiometric amounts (~90%) of H₂ and O₂ (illustrated with 6 consecutive cycles). The ability of the cycle to capture 75% of the ~400 ppm CO₂ from air is demonstrated also. A technoeconomic analysis of the integrated process for the renewable production of H₂, O₂, and electricity, as well as DAC of CO₂, shows that the proposed scheme of solar-driven H₂ production from thermochemical water splitting coupled with CO₂ DAC may be economically viable under certain circumstances.

sodium manganese oxide | process integration | technoeconomic analysis

Anthropogenic climate change is increasingly recognized as a serious and imminent threat to the prosperity of human society. A recent Intergovernmental Panel on Climate Change report claims that human activities have caused ~1 °C increase in global warming, and that net zero global CO₂ emissions is needed to cap the warming below 1.5 °C in the next few decades to avoid catastrophic consequences (1). Even if drastic measures are taken to completely halt anthropogenic CO₂ emissions by 2040, negative effects of high atmospheric CO₂ concentration [415 ppm as of May 2019 (2)] will still persist for decades afterward (1) due to the long lifetime of atmospheric CO₂ [thousands of years (3)]. Thus, it is not only imperative to reduce CO₂ emissions, but also to consider actively removing CO₂ from the atmosphere via direct air capture (DAC). Temperature/humidity swing adsorption with liquid amines, e.g., monoethanolamine (4, 5), is widely practiced in the natural gas industry and has been proposed for CO₂ removal from the flue gas of coal-fired power plants (6–8). The majority of the most mature DAC processes are based on amine absorbents (9); however, several undesirable features, such as low stability to oxygen environments at elevated temperatures of CO₂ removal (4), make the search for alternative absorbents necessary. Aqueous alkali hydroxides have been proposed as absorbents in DAC processes very similar to the Kraft Caustic Recovery Cycle commonly used in the paper mill/pulp industry (10) (left cycle in Fig. 1). In these processes, dilute CO₂ in air is absorbed to form carbonate ions in an aqueous solution of an alkalai hydroxide (Na⁺ or K⁺) via an acid–base reaction, followed by the introduction of calcium hydroxide to separate the carbonate in the form of solid calcium carbonate precipitate and regenerate the initial alkali hydroxide solution. The calcium carbonate is then calcined at high temperatures (600–900 °C) to produce a CO₂ stream and calcium oxide ready for use in the subsequent cycle (after reacting with water to form calcium hydroxide), thus completing the cycle (SI Appendix, Table S1). Two metal cations must be used in the cycle due to the different properties of the hydroxide and carbonate of Na⁺(K⁺) and Ca²⁺. The low solubility of Ca(OH)₂ in water makes it unsuitable as an absorbent, while the high thermal stability of K₂CO₃ and Na₂CO₃ makes their thermal decomposition energetically unfavorable (11). Despite recent research efforts, existing DAC processes have economics that could stand improvement (100–500 $/ton CO₂) (4, 9, 12–14). A comprehensive technoeconomic analysis of current DAC technologies is available (14).

We recently developed a thermochemical water-splitting (TWS) cycle based on a Mn–Na–CO₂ reaction network that is capable of splitting water into stoichiometric amounts of hydrogen and oxygen in 3 reaction steps, the highest temperature step operating at 850 °C (15). In contrast to sulfur–iodine TWS cycles (16, 17), no toxic/corrosive chemicals are involved in the Mn–Na–CO₂ system. The thermodynamic driving force of this TWS cycle is based on the divergent thermal and chemical stabilities of different manganese oxide species (α-NaMnO₂ and MnO₃) at different temperatures and environments (SI Appendix, Table S1 and Fig. S1) (15). A key step of interest in the context of DAC in this cycle is the extraction of Na⁺ from MnO₂ layers of α-NaMnO₂ with CO₂ to form H₂MnO₄ and Na₂CO₃. α-NaMnO₂ slurries are quite alkaline in nature (pH = 12–13), making them attractive as potential CO₂ adsorbents. There are 2 key parallels between the Mn–Na–CO₂ TWS and the Kraft cycles: 1) the use of CO₂ as an acid to extract alkali cations from an alkaline compound (α-NaMnO₂ and NaOH/KOH, blue in both cycles in Fig. 1), and 2) high-temperature production of CO₂ via thermal decomposition of sodium manganese oxide.
carbonates (Na$_2$CO$_3$ and CaCO$_3$, green in Fig. 1). These 2 parallels suggest that the Mn–Na–CO$_2$ TWS cycle has the potential to enable simultaneous DAC of CO$_2$ and hydrogen production.

In this work, we demonstrate a methodology that integrates the DAC of CO$_2$ and thermochemical water splitting that produces a mixture of CO$_2$ and H$_2$ (Fig. 2). The CO$_2$/H$_2$ stream can be converted to a variety of fuels and chemicals, e.g., to syngas via the reverse water–gas shift reaction (18, 19) or methanol via a number of established catalytic processes (20). Alternatively, the CO$_2$/H$_2$ stream can be directly upgraded to higher hydrocarbons via a number of established processes (18, 21). In this work we limit our analysis to CO$_2$ capture and water splitting without considering the economics of further upgrading in order to avoid added complexity and to allow our process to be directly compared to other DAC technologies.

Results and Discussion

Process Design and Implementation. While the chemistry of all 3 steps in the Mn–Na–CO$_2$ cycle has been established (15), several major challenges remain in coupling the cycle with a solar thermal power source and the DAC of CO$_2$: 1) solar energy is inherently intermittent, restricting the amount of time high-temperature reactions can be performed, and 2) DAC of CO$_2$ requires the utilization of massive airflow. Here, we provide a solution to these issues (summarized in Fig. 3). Heat for the cycle is supplied by a solar collector field that provides superheating steam at 1,000 °C. Concentrated solar thermal power is useful for generating high-quality heat for water-splitting cycles (22, 23). However, the diurnal cycle of available solar energy restricts the amount of time a water-splitting reactor can be held at high temperatures on a daily basis. Conveniently, the combined water splitting and CO$_2$ DAC process has 2 distinct operating regimes that pair well with this diurnal constraint. While high-quality heat is required for the high-temperature steps, sodium extraction is exothermic and time-consuming if dilute atmospheric CO$_2$ is used as a CO$_2$ source. Thus, cyclic quasibatch operation (SI Appendix, Fig. S2 shows a schedule of operation) where the high-temperature steps are performed during the day and Na$^+$ extraction is performed at night enables effective utilization of reactor time despite the limitations imposed by a diurnal power source. CO$_2$ DAC requires large flowrates of air due to the low concentration (~400 ppm) of CO$_2$. This leads to a correspondingly large power requirement. We address this issue by utilizing a solar updraft tower (24, 25). These systems have been proposed since the late 1990s for generating power from solar heating via thermal updraft through a power-generating turbine suspended in a large tower surrounded on ground level by a large solar collection basin (Fig. 3, Left). While this technology usually shows low efficiency in producing electricity (24), it is an excellent source of high flowrates of air without the corresponding electricity costs. Large thermal reservoirs are typically included in the base of updraft towers such that the airflow is relatively constant 24 h a day (24). This allows the tower to provide airflow for CO$_2$ capture at night, while also producing electricity during the day, when large air flowrates are not required. In general, diurnal systems suffer from low throughput and low efficiencies due to poor process intensification (26). We stress that this diurnal quasibatch operating schedule is only 1 possible implementation of this concept. With the implementation of a high-temperature heat storage system (27) and/or a system for the recirculation of solids (13) the cycle could be executed with higher frequencies or even continuously. We evaluate only our diurnal process in this work in order to simplify the necessary process assumptions.

The integration of these 2 passive solar technologies with the Mn–Na–CO$_2$ water-splitting cycle yields a process ideal for coupling DAC of CO$_2$ and water splitting. High-temperature steps that produce hydrogen and oxygen operate during the daytime when concentrated solar energy is available with steam as a heating medium (Fig. 3, Top Right). The heat exchange/ separation system (Fig. 3, Bottom Right) receives either dilute streams of H$_2$ and CO$_2$ in steam (during the H$_2$ evolution step) or O$_2$ in steam (during the thermal reduction step) from the TWS reactor. These steps are separated by either the introduction of Na$_2$CO$_3$ or sodium extraction, such that these 2 gas streams are produced in distinct time periods throughout the cycle. This gas stream is first cooled to ~600 °C in a heat exchanger that provides heat to dry Na$_2$CO$_3$ solutions produced during sodium extraction. This 600 °C stream is then introduced to a steam turbine that cools the stream further to 100 °C and produces electricity. A water knockout vessel is used to condense the majority of the steam and separate the final gas.
mixture (H\textsubscript{2} and CO\textsubscript{2} or O\textsubscript{2}). The condensed water can then be reused as a source of steam such that each cycle consumes only the stoichiometric amount of water for water splitting. The use of steam as a reactant, heat-transfer medium, and carrier gas significantly simplifies the process as compared to other water-splitting cycles that require inert carrier gases (28, 29) and makes leveraging waste heat (via steam turbine) possible. At night, liquid water is supplied to the water-splitting reactor and a constant flow of air from the solar updraft tower (heat captured during the day by the solar tower allows operation at night; 24-h operation of solar towers is already proven in other applications) is introduced in order to extract sodium from Na\textsubscript{2}CO\textsubscript{3} solutions. Upon introduction of steam to the solid mixture at 850 °C at the end of 5 h (Fig. 4A, ii), MnO is oxidized by water in the presence of unreacted Na\textsubscript{2}CO\textsubscript{3} to form additional α-NaMnO\textsubscript{2} and a mixture of H\textsubscript{2} and CO\textsubscript{2} with a molar ratio of 1:1.

**Reactor Design.** We designed and fabricated an integrated reactor for the Mn-Na-CO\textsubscript{2} TWS cycle to demonstrate its feasibility. In our previous work, we elucidated the chemistry of each step in Mn–Na–CO\textsubscript{2} TWS in separate reactors on a test-tube scale (∼200-mg solid) (15). A key challenge in the implementation of this cycle is the drastically different operating conditions, i.e., solid–gas phase reactions at 850 °C for hydrogen and oxygen evolution, and solid–liquid phase Na\textsuperscript{+} extraction and CO\textsubscript{2} capture at 90 °C. The fabricated reactor eliminates the necessity of movement of solids with a capacity of handling ∼10-g solid, which is a 50-fold increase from our initial work (SI Appendix, Fig. S3, details provided in Experimental Procedures). The reactor consists of an alumina vessel with multiple inlets and outlets to control the flow of liquids and gases into and out of the reaction zone. We demonstrate below that all 3 steps in the Mn–Na–CO\textsubscript{2} TWS cycle can be conducted in the fabricated reactor with close to stoichiometric H\textsubscript{2} and O\textsubscript{2} yields in multiple cycles.

**Hydrogen Evolution.** During hydrogen evolution, a stoichiometric mixture of Mn\textsubscript{3}O\textsubscript{4} and Na\textsubscript{2}CO\textsubscript{3} (2:3 molar ratio) is initially heated to 850 °C under an inert gas flow and then under steam to form NaMnO\textsubscript{2}, CO\textsubscript{2}, and H\textsubscript{2}. Prior to the introduction of steam (Fig. 4A, i), the only gas-phase product formed as the solid mixture is heated to 850 °C is CO\textsubscript{2}. This reaction leads to the formation of a 2:1 ratio of α-NaMnO\textsubscript{2} and MnO as we have shown in our previous work (15). No oxidation state change of manganese occurs in this step, as Mn\textsuperscript{III} and Mn\textsuperscript{II} in Mn\textsubscript{3}O\textsubscript{4} form α-NaMnO\textsubscript{2} and MnO, respectively. Upon introduction of steam to the solid mixture at 850 °C at the end of 5 h (Fig. 4A, ii), MnO is oxidized by water in the presence of unreacted Na\textsubscript{2}CO\textsubscript{3} to form additional α-NaMnO\textsubscript{2}, and a mixture of H\textsubscript{2} and CO\textsubscript{2} with a molar ratio of 1:1.

**Sodium Extraction.** Na\textsuperscript{+} intercalated in the MnO\textsubscript{2} layers of α-NaMnO\textsubscript{2} must be removed before Mn\textsubscript{3}O\textsubscript{4} can be thermally reduced to complete the TWS cycle. Sodium extraction is performed by cooling the solids produced in the hydrogen evolution step to 90 °C, followed by the introduction of water to the reactor via the liquid inlet and bubbling of 5% CO\textsubscript{2} in N\textsubscript{2} via the gas inlet into the liquid–solid slurry. Initially complete CO\textsubscript{2} absorption is observed (Fig. 4B), demonstrating the high binding affinity of Na\textsuperscript{+} to CO\textsubscript{3}\textsuperscript{2–} and facile sodium extraction from the birnessite phase formed upon contact of α-NaMnO\textsubscript{2} with liquid water (30). A detectable amount of CO\textsubscript{2} is observed after 2.3 h in the breakthrough curve, followed by a quick increase of the CO\textsubscript{2} concentration to the level in the feed. Periodic pH measurements of the slurry during extraction reveal that the slurry is very alkaline in nature, with an initial pH of 12.8, that decreases gradually as CO\textsubscript{2} is adsorbed into the liquid. This indicates that Na\textsuperscript{+} is gradually extracted from the birnessite phase by CO\textsubscript{2}. The onset of incomplete CO\textsubscript{2} adsorption occurs when the pH of the slurry decreases to ∼11.4, that corresponds to ∼63% of Na\textsuperscript{+} extracted based on the amount of CO\textsubscript{2} absorbed. The final pH of the slurry is ∼9.6, which is roughly consistent with the formation of a Na\textsubscript{2}CO\textsubscript{3}–NaHCO\textsubscript{3} buffer (31) after the majority of Na\textsuperscript{+} has been extracted from the solid. Upon the completion of sodium extraction, the Na\textsubscript{2}CO\textsubscript{3}–NaHCO\textsubscript{3} solution is extracted from the liquid outlet equipped with an inline filter.

**Fig. 3.** A schematic of the proposed integrated DAC and TWS process.
Thus, higher weight loadings of Na$_2$CO$_3$ solution. Evaporating the water yields a solid mixture of reacto ri sc o o l e dt o only take place during the H$_2$ production step as mixtures of CO$_2$ and O$_2$ are produced in each cycle (Fig. 5). This is likely a consequence of the lower initial pH from the NaMnO$_2$ layers (15). Thus, each cycle has a fresh start without any “memory” from previous cycles. Further, roughly 90% of the stoichiometric amount of CO$_2$ is produced in the hydrogen evolution step in each cycle (Fig. 5B), with a negligible amount of CO$_2$ produced in the thermal reduction step. Powder X-ray diffraction patterns of solids after all 3 steps in the 6th cycle were collected (SI Appendix, Fig. S4A) and the data confirm that the expected phases are formed based on our previous work (15). Analysis of the X-ray diffraction patterns of produced Mn$_3$O$_4$ at several points in the cycle (SI Appendix, Fig. S4B) reveals that the average particle size decreases slightly during the 6-cycle test. Typically, CaO/Co$_2$O$_3$ particles in Kraft-based cycles increase in size dramatically over the course of many cycles, limiting practical reuse (13). The fact that our Na–Mn–CO$_2$ system does not suffer from the same issue may allow substantially longer times between the replacement of solids. Results from these cycling experiments demonstrate that the fabricated reactor can be used to facilitate the Mn–Na–CO$_2$ TWS cycle.

**Thermal Reduction.** The solid component remaining in the reactor from the sodium extraction step is dried and heated to 850 °C to regenerate Mn$_3$O$_4$. O$_2$ is produced during the temperature ramp and hold (Fig. 4C) due to the thermal reduction of manganese oxides produced in the previous steps to Mn$_3$O$_4$ (15). Additionally, a negligible amount of CO$_2$ is also produced during this step via decomposition of MnCO$_3$ [formed during sodium extraction (15)] during heating. Higher amounts of CO$_2$ formation were observed in our previous work (15). The difference may be due to the significantly higher α-NaMnO$_2$ weight percentage in the slurry during sodium extraction (~15 wt %) employed in this work, as compared to our previous work (~5 wt %) (15). Sodium extraction takes place via competing redox extraction (2NaMnO$_2$ + 4H$^+$ → MnV$^0$O$_2$ + Mn$^{2+}$ + 2Na$^+$ + 2H$_2$O) and ion exchange (NaMnIIIO$_2$ + H$^+$ → H MnIV$^0$O$_2$ + Na$^+$) where the redox mechanism dominates at low pH (32). MnCO$_3$ formation is attributed to the reaction between the aqueous Mn$^{2+}$ formed via the redox mechanism and CO$_2$ (15). Thus, higher weight loadings of α-NaMnO$_2$ in the slurry suppress the redox mechanism by maintaining a higher pH, which in turn reduces the MnCO$_3$ formation. It is desirable for CO$_2$ evolution to only take place during the H$_2$ production step as mixtures of CO$_2$ and H$_2$ are valuable while mixtures of CO$_2$ and O$_2$ would require costly separation. At the end of the thermal reduction step, the reactor is cooled to ~100 °C, followed by the introduction of a Na$_2$CO$_3$ solution. Evaporating the water yields a solid mixture of Na$_2$CO$_3$ and Mn$_3$O$_4$ ready for the hydrogen evolution step in the subsequent cycle.

**Overall Cycling Performance.** Near-theoretical yields of H$_2$ and O$_2$ are produced from the Mn–Na–CO$_2$ TWS cycle in the fabricated reactor over the course of 6 consecutive cycles (Fig. 5A). A stoichiometric mixture of Mn$_3$O$_4$/Na$_2$CO$_3$ is charged in the reactor prior to the hydrogen evolution step of the first cycle, and no additional manganese oxide is added to or removed from the reactor in all consecutive cycles. Roughly 90% of the theoretical amounts of H$_2$ and O$_2$ are produced in each cycle (Fig. 5A) with no discernible decrease in production with increasing cycles. The lack of deactivation is in part due to the fact that complete phase change is involved in every step, e.g., the intercalation of Na$^+$ into and extraction of Na$^+$ from MnO$_x$ layers (15). Thus, each cycle has a fresh start with no “memory” from previous cycles. Further, roughly 90% of the stoichiometric amount of CO$_2$ is produced in the hydrogen evolution step in each cycle (Fig. 5B), with a negligible amount of CO$_2$ produced in the thermal reduction step. Powder X-ray diffraction patterns of solids after all 3 steps in the 6th cycle were collected (SI Appendix, Fig. S4A) and the data confirm that the expected phases are formed based on our previous work (15). Analysis of the X-ray diffraction patterns of produced Mn$_3$O$_4$ at several points in the cycle (SI Appendix, Fig. S4B) reveals that the average particle size decreases slightly during the 6-cycle test. Typically, CaO/Co$_2$O$_3$ particles in Kraft-based cycles increase in size dramatically over the course of many cycles, limiting practical reuse (13). The fact that our Na–Mn–CO$_2$ system does not suffer from the same issue may allow substantially longer times between the replacement of solids. Results from these cycling experiments demonstrate that the fabricated reactor can be used to facilitate the Mn–Na–CO$_2$ TWS cycle.

**DAC of CO$_2$ with Mn–Na–CO$_2$ TWS Cycle.** The alkaline nature of aqueous suspensions of α-NaMnO$_2$ (Fig. 4B) suggests that it could be an effective adsorbent for CO$_2$ DAC. To evaluate the feasibility of this we employ ambient air with ~400 ppm of CO$_2$ as a CO$_2$ source to extract sodium from α-NaMnO$_2$ rather than 5% CO$_2$ that was used in our cycling test. The breakthrough curves for CO$_2$ absorption in several α-NaMnO$_2$ slurries with varying NaMnO$_2$ loadings (Fig. 6A) show an excellent initial performance, absorbing more than 75% of CO$_2$ from air. The fraction of sodium extracted by CO$_2$ before the air effluent contains more than 100 ppm of CO$_2$ increases (up to 72%) with the amount of α-NaMnO$_2$ in the slurry (Fig. 6B). This is likely a consequence of the lower initial pH of slurries with low weight loading of α-NaMnO$_2$ (0.2–0.8 wt %) caused by near-complete extraction of Na$^+$ from the MnO$_x$ layers upon initial introduction of water. In contrast, ~15 wt % of NaMnO$_2$ slurry in water is used in the cycling experiments. The primary reason to use a relatively low NaMnO$_2$ wt % slurry in the proof of concept CO$_2$ DAC experiments is to measure these

![Diagram](Image)
breakthrough curves in reasonable timescales. The amount of CO₂ captured when the CO₂ concentration in the effluent recovers to 400 ppm is in all cases consistent with ∼90% of intercalated Na⁺ being converted to Na₂CO₃ (similar to the case when 5% of CO₂ is used). As the weight fraction of NaNMnO₂ in the slurry increases, the fraction of Na⁺ extracted before the effluent air contains more than 100 ppm of CO₂ is expected to increase.

Technoeconomic Analysis. A preliminary technoeconomic analysis is performed on a large-scale design based on the proposed flow-sheet (Fig. 3) on a plant that removes 25,000 metric tons of CO₂ from the atmosphere per year. The major unit processes used are 1) a water-splitting reactor, 2) a field of heliostats focused on a central tower, 3) a solar updraft tower to generate the airflow during Na⁺ extraction at night and provide electricity during the day, and 4) a series of steam turbines to utilize waste heat in cooling the water-splitting effluent. The system is simulated as a quasibatch system where a water-splitting reactor is cycled through all of the required steps of the cycle once a day, performing the high-temperatures step during the day and the low-temperature step at night (SI Appendix, Fig. S2). In order to determine the extent to which integration of the updraft tower and steam turbines affects the feasibilities of the plant, we consider 4 alternate cases. Case 1 considers a plant that is not integrated with solar airflow or steam turbines for power recuperation. Case 2 considers a plant that is integrated with a solar updraft tower for airflow and power generation. Case 3 considers the construction of a full plant

Fig. 5. Amounts of (A) H₂ and O₂ produced, and (B) CO₂ captured/produced in 6 consecutive TWS cycles. All gas products are quantified with an SD of ∼4%.

Fig. 6. CO₂ direct air capture experiments: (A) CO₂ in air breakthrough curves during DAC tests. (B) The fraction of the Na⁺ extracted during DAC of CO₂ while CO₂ concentrations are below 100 ppm. All gas products are quantified with an SD of ∼4%.
with the implementation of a series of turbines but no solar updraft tower. Case 4 takes into account the implementation of both a solar updraft tower and a series of steam turbines. A detailed account of all 4 cases and all relevant assumptions are provided in SI Appendix, Fig. S5. For each case we evaluate the economic viability using an optimistic CO$_2$ price of $200/ton. Cases 1–3 exhibit negative net present values (NPVs) even at this quite optimistic CO$_2$ price, highlighting the importance of proper integration of the water-splitting cycle with other technologies for power generation. The NPV of case 4 is dependent on financial assumptions but is positive, assuming a CO$_2$ price of $200/ton. Thus, we focus on the fully integrated system for further analysis.

The total costs of the fully integrated system investigated in case 4 are dominated by initial capital costs, mainly the solar energy harvesting facilities (Fig. 7A). The combination of concentrated solar power, water splitting, CO$_2$ removal, and turbines to produce electricity makes this process different from other DAC processes in its reliance on several revenue streams (Fig. 7B). The generation of H$_2$, O$_2$, CO$_2$, and electricity contributes to the plant’s revenue, although the vast majority of the revenue is derived from the generation of electricity. As a result, the economic viability of the proposed process depends on the value of all these outputs (Fig. 7C). For example, assuming H$_2$, O$_2$, and electricity prices of $2/kg (33), $0.04/kg (34), and $0.167/kWh [average cost of electricity in California in June 2019 (35)], respectively, a CO$_2$ price of $83/ton is needed for the proposed plant to break even. This hypothetical price point is lower than previous reports (4, 12–14). This relatively low projected break-even CO$_2$ price illustrates the economic feasibility of such an integrated process, despite its dependence on the assumed electricity price (SI Appendix, Fig. S6). It is noted that no carbon subsidy is assumed in the analysis.

Due to the nature of the proposed cycle, the degree of uncertainty of the techno-economic analysis will be greater than those on DAC schemes leveraging years of pilot plant data and vast industrial knowledge on similar processes. Our process is leveraging 2 reasonably mature solar-to-power technologies (updraft towers and solar thermal concentrators coupled to heat engines) to offset the costs of CO$_2$ DAC via electricity generation; thus, the comparison to processes that aim only to capture CO$_2$ may not be a fair comparison. Although the current analysis focusses on CO$_2$ DAC, the proposed cycle can also use more concentrated sources of CO$_2$, which is expected to yield improved economics as less inlet gas flow is needed for a given amount of CO$_2$ captured. As the majority of the capital cost of the process is derived from the solar-harvesting facilities, it is expected that as these technologies mature and improve, these costs will decrease. There is precedence in literature for dramatic cost reduction after the construction of a pioneer plant as a result of improvements in manufacturing/ experience in both solar-harvesting technologies (36) and for direct air capture (13).

**Conclusions**

We established the concept of using a Mn–Na–CO$_2$ water-splitting cycle combined with the DAC of CO$_2$ to create an integrated methodology that under certain circumstances may be economically viable for the production of renewable chemicals. We have designed a complete process schematic, constructed a 1-vessel test reactor capable of performing a complete TWS cycle, performed proof of concept CO$_2$ direct air capture tests, and assessed the economic viability of the approach. We have demonstrated that our reactor design can facilitate several cycles of the complete water-splitting reaction system. Direct air capture tests illustrated that alkaline slurries of NaMnO$_2$ could remove CO$_2$ from air with high efficacy even at ultradilute CO$_2$ concentrations. While our techno-economic analysis was preliminary, it demonstrated that the proposed process may be economically feasible for combining CO$_2$ removal from the atmosphere with thermochemical water splitting to produce a gas stream (CO$_2$H$_2$) that could be readily integrated with existing chemical processes to create renewable chemicals.

**Experimental Procedures**

**Cycling Tests.** The experimental test reactor is constructed from readily available parts. The reactor tube is an 8.5-in.-long closed-ended Al$_2$O$_3$ tube with an outer diameter of 1 in. and an inner diameter of 0.75 in. The gas-handling and liquid-handling tubes are Al$_2$O$_3$ tubes with outer diameters of 1/4 and 1/8 in. and inner diameters of 3/16 and 1/16 in., respectively. All Al$_2$O$_3$ tubes are purchased from AdValue Technologies. An Inconel clathromouple (Omega) is threaded through the gas-handling tube in order to accurately measure temperatures in the reactor tube. All gas flows are controlled with Brooks mass-flow controllers. Steam is introduced via a Cole Parmer syringe pump with a 60-ml syringe. All other liquid additions or extractions are performed using the reactor syringe port using a Corning 0.20-μm nylon filter as an inline filter. The reactor is heated by a clamshell furnace (Lingberg/Blue M) and the gas distribution section is heated to 150 °C to prevent condensation of steam. All heaters and mass flow controllers are controlled via an integrated LabView program made in-house. An online differentially pumped quadrupole mass spectrometer (Stanford Research Instruments RGA100) is used to quantify gas products and a 100-ml condenser suspended in an ice bath is positioned upstream of the residual gas analyzer (RGA) to remove excess water.

Cycling experiments are performed by first preparing a physical mixture of Mn$_2$O$_3$ and Na$_2$CO$_3$ (Sigma-Aldrich) via grinding with a mortar and pestle. This mixture is then loaded into the reactor with care so as not to clog the gas-handling tube with solids. The water-splitting step is performed by first heating the physical mixture to 850 °C under a flowrate of 50 scm of nitrogen at

![Image](image.png)

**Fig. 7.** Technoeconomic analysis of the process: (A) A summary of installed costs of major unit processes, (B) the annual revenue and costs of operating the DAC and water-splitting process assuming CO$_2$ is sold at 83 $/ton, and (C) the net present value of the process as a function of the selling prices of H$_2$ and CO$_2$ (the line represents the prices required for the plant to break even).
a ramping rate of 10 °C/min. After the set temperature is reached, steam is introduced to the heated inlet gas line by syringe pump at a rate of 2 mL/h. Once the reaction is completed (hydrogen production ceases) the steam flowrate is stopped, and the reactor is cooled to room temperature under nitrogen.

Once cool, sodium extraction is performed. First ~40 mL of deionized (DI) water is introduced to the reactor using the syringe port on the side of the reactor to 850 °C at a rate of 10 °C/min under a flow of 50 sccm of nitrogen. After each step flowrate of CO2 is introduced into the nitrogen carrier gas such that a 5% CO2 in N2 stream at 50 sccm is fed to the reactor. The online RGA is used to measure CO2 concentration during this time to establish a rough degree of extraction. Sodium extraction is performed in 3 steps starting from 5% CO2 in N2, 10% CO2 in N2, and finally pure CO2 to insure complete sodium extraction. Between each step ~40 mL of liquid is extracted, filtered, and replaced with the same volume of DI water using the inline filter attached to the syringe port. After the pH of the extracted solution is ~7.5–8 the reactor is heated to and held at 100 °C for several hours to remove the remaining water.

Thermal reduction of the postextraction solid is performed by heating the reactor to 850 °C at a rate of 10 °C/min under a flow of 50 sccm of nitrogen. The RGA is used to quantify produced O2 and the reactor is cooled once oxygen production ceases. After cooling to room temperature an aqueous solution of Na2CO3 is introduced to the reactor via the syringe port and the reactor is heated to 100 °C, removing water and leaving a physical mixture of MnO2 and Na2CO3 which is used in the next cycle.

**Air-Scrubbing Tests.** Air-scrubbing tests are performed using a small 4-in. closed-end Al2O3 tube with an outer diameter of 1 in. and an inner diameter of 0.75 in. A 60-sccm flowrate of air is supplied with a small SVDC diaphragm pump with a potentiometer to control the flowrate. The same RGA as above is used to track CO2 concentration during Na extraction. A ppm level CO2 calibration is obtained for the RGA by diluting atmospheric air with nitrogen in various ratios. Na2MnO4 used in tests was synthesized via solid-state synthesis using stoichiometric amounts of Mn2O3 (Sigma-Aldrich) and Na2CO3 heated in flowing air at 700 °C for 6 h.

**Data availability.** All data are available in the main text and SI Appendix.

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