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Broader context

The cement industry is the second-largest industrial contributor to global greenhouse gas emissions. Portland cement is the largest regulated and ubiquitous commodity, second only to water. The main decarbonization challenge arises from the decomposition of limestone – the primary cement feedstock – which accounts for 60% of total CO₂ emissions. No technology has yet decarbonized this 200-year-old liming routine for producing cement. Here, we develop a novel decarbonization approach to utilize abundant, carbon-free calcium silicates from rocks and wastes. By coupling enhanced weathering of calcium silicates with water electrolysis, we demonstrate the production of carbon neutral/negative calcium carbonate, green hydrogen, and amorphous silica suitable for direct integration with contemporary cement manufacturing. A zero-gap electrolyzer configuration was determined to improve hydrogen productivity and durability. Life cycle assessment and techno-economic analysis indicate this process can maintain profitability while being carbon neutral/negative. This approach could enable gigaton-scale annual decarbonization of the cement industry, meeting regulatory compliance with minimal capital investment.

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- † Electronic supplementary information (ESI) available: Experimental methods, LCA and TEA methods, electrochemistry data, XRD, SEM, TEM, STXM, Raman spectra, LCA and TEA results. See DOI: <https://doi.org/10.1039/d4ee03529a> ‡ Equal contribution.

Scalable electrified cementitious materials production and recycling†

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The production of Portland cement, the industry-standard cement, contributes \sim 8% of global CO₂ emissions through fossil-fuel heating and decomposition of limestone (the primary cement raw material). Decarbonization, e.g., via direct electrification, of this 200-year-old liming routine is extremely challenging at the industry scale. We propose a scalable electrochemical decarbonization approach to circumvent the limestone use by switching to carbon-free calcium silicates from abundant minerals and recycled concrete. Water electrolysis produces protons and hydroxides to drive a pH gradient that accelerates Ca^{2+} ion leaching from calcium silicates and captures atmospheric $CO₂$ to form carbon-negative CaCO₃, which serves as the feedstock for cement manufacturing or as the carbon-mineralized product for cement substitution with permanent carbon storage. Value-added co-products amorphous silica and green H₂ further enhance cement performance and supplant fossil fuels for net-zero transition, respectively. The products readily meet presentday regulatory standards and demands, and the approach readily synergizes with business-as-usual cement manufacturing and concrete construction, which are important for upscaling and structural safety, promising ready reception by the public and industries. Blended Portland cement produced through our approach with carbon-negative CaCO₃ and silica demonstrates enhanced resilience and achieves carbon neutrality or negativity when incorporating storage or circulation of $CO₂$ from cement plant flue gas, respectively. This lowcost, electrochemical cement production approach using abundant ubiquitous raw materials enables electrification, transition to clean fuel, and decarbonization at a gigaton scale. PAPER

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Introduction

Cement, the essential binder of concrete – the most used material worldwide, only second to water, is produced at over 4 Gigatonnes per year (Gt per y) and contributes to a significant 8% of global CO_2 emissions.^{1,2} The annual global demand for cement is projected to increase by 50% by 2050 because urbanization and the surge in the renewable energy sector create significant demands for cement in the construction of building and infrastructure. Thus, it is urgent to incorporate carbon capture, utilization, and storage (CCUS) technologies

CO2 emissions of Portland cement (the most common cement and a ubiquitous commodity) production mainly arise from (i) limestone (CaCO₃) decomposition to CaO and $CO₂$ $\sim 60\%$ contribution) and (ii) fossil-fuel combustion for cement kiln heating for pyroprocessing (\sim 30% contribution).⁴ Despite extensive research, there has been no substantial breakthrough in decarbonization of this 200-year-old global industrial-standard liming routine, where the thermal efficiency of cement rotary kilns has been optimized over the 200 years to a ceiling. Most partial decarbonization strategies rely on partial Portland cement substitution by supplementary cementitious materials (SCMs). Common supplementary cementitious materials are vitreous silicates sourced from industrial byproducts, including fly ash from coal-fired power plants, which accounts for 50% of total share of the current supplementary cementitious materials market, and blast furnace slags from the iron/steel industry.⁵ However, these conventional supplies of supplementary cementitious materials are currently under shortage of over 0.5 Gt per y, while the shortage will be further exacerbated by the increasing cement demand and decommissioning of $CO₂$ -intensive traditional coal power plants and conventional steel manufacturing.⁶ Regardless, supplementary cementitious materials as a partial substitute for Portland cement are practically limited to $<$ 50% replacement level, therefore unable to achieve full cement decarbonization towards carbon-neutral/ negative Portland cement.4 Besides, studies on incorporating CCUS and/or clean energy with cement production are burgeoning, but a lack of focus on scalability and long-term $CO₂$ -storage in concrete precludes technology transfer, especially since the 1450 \degree C heating of massive cement kilns is beyond the feasibility of direct electrification, which also lacks economic incentives due to long payback periods of cement plant remodeling.⁷ Recent studies suggest that many emerging cement decarbonization techniques relying on $CO₂$ mineralization in concrete materials (e.g., $CO₂$ curing of concrete blocks and direct utilization of cement carbonation products as cementitious materials) may not effectively offset the lifecycle carbon footprint of cement at scale.^{8,9} The lack of international standards for these alternative low-carbon cements also discourages the deployment of low-carbon technologies in the construction sector for the concerns over liabilities and safety. Paper

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Herein, we demonstrate a carbon-neutral-to-negative, economically attractive Portland cement production scheme that utilizes abundant carbon-free natural/recycled materials and renewable electricity while co-producing carbon-neutral-or-negative supplementary cementitious materials and clean fuels. The strategy couples water electrolysis and $CO₂$ direct air capture with electrochemical generation of carbon-sequestered CaCO₃ – which substitutes limestone as the primary cement manufacturing feedstock and neutralizes limestone decomposition-induced $CO₂$ emissions – from non-carbonaceous precursors, which are highly available worldwide (e.g., basalt – half volume of the Earth's crust surface), and industry wastes (e.g., recycled concrete fines from construction and demolition waste).¹⁰ Our approach produces materials (Portland cement feedstock and supplementary cementitious materials) that are compatible with existing Portland cement manufacturing

Fig. 1 Scheme of the electro-geochemical cell powered by renewable electricity for converting Ca-bearing silicates (industrial/construction/ mine wastes and rocks) to $CaCO₃$ and high-value amorphous silica SCM for carbon neutral-to-negative Portland cement. Captured $CO₂$ can transform into dissolved bicarbonate (e.g., $Ca(HCO₃)₂$) to mitigate ocean acidification. Water electrolysis generates H_2 and O_2 as a green energy carrier for zero-emission infrastructure and oxyfuel for CCUS, respectively.

infrastructure and decarbonization technologies, 11 avoiding regulatory limits on this safety-sensitive structural material. Portland cement developed through our approach complies with existing international and national standards, and its usage does not require new training for millions of materials and civil engineers or builders globally. Furthermore, the decreasing cost of renewable electricity, now below that of conventional fossil-fuel-powered electricity, together with potential carbon credit savings enabled by the present carbon-neutral-to-negative cement manufacturing scheme, provides further economic competitiveness beyond environmental benefits.¹¹

Fig. 1 demonstrates the potential integration of our approach with cement manufacturing as well as CCUS processes and a circular CO₂ pathway. Our room temperature electro-geochemical cell takes in electricity (possibly in the form of renewable energy) as well as captured atmospheric $CO₂$ (or circulated concentrated CO2) and supports water electrolysis, which incurs a pH gradient enabling Ca^{2+} extraction from precursors calcium silicates to produce CaCO₃, amorphous $SiO₂$, and green H₂ gas. By delivering the same primary feedstock as current Portland cement production, $CaCO₃$, we provide a cost-efficient, scalable decarbonization and energy transition pathway to the cement industry, which has previously been reluctant to embrace emerging marginal decarbonization technologies due to their costly, long payback periods for retrofitting or remodeling modern cement plants. 12

Our strategy offers the following benefits: (1) full carbonneutralization of limestone decomposition through $CO₂$ circulation or direct air-captured $CO₂$ mineralization and storage at scale, (2) production of high-value supplementary cementitious materials (carbon-negative reactive $CaCO₃$ and highly pure amorphous silica) to further decarbonize cement for carbon credit savings, enhance life-time concrete performance, and lower production and operational costs, (3) H₂ generation for industrial heating, power generation, and chemical manufacturing, (4) O₂ generation for oxy-fuel combustion aiding in sequestering concentrated $CO₂$ flue gas at cement plants, and (5) generation of calcium (bi)carbonate water to mitigate ocean acidification. $13-15$ We studied this process in three electrochemical reactor configurations, evaluated the products and performances, and performed life cycle assessment and technoeconomic analysis to probe the embodied carbon, energy use, and economic viability.

Results and discussion

Electrolytic dissolution of calcium silicates

H-cell configuration. Ambient-temperature water electrolysis in an H-cell was performed to convert a model carbon-free calcium silicate, wollastonite (CaSiO₃), to CaCO₃ precipitates (e-CaCO₃) and amorphous silica (a-SiO₂) (Fig. 2A). The precursor $CaSiO₃$ is placed in the anodic compartment, and two pieces of porous cellulose separators are used to prevent cathode

passivation.¹⁶ In a near-neutral 1 M NaNO₃ electrolyte, H_2O dissociation is required for both the hydrogen evolution reaction (HER, eqn (1)) and the oxygen evolution reaction (OER, eqn (2)), thereby producing H^+ and OH⁻ for subsequent reactions.

$$
2H_2O + 2e^- \to 2OH^- + H_{2(g)}
$$
 (1)

$$
2H_2O \to 4H^+ + O_{2(g)} + 4e^-
$$
 (2)

Note that $NO₃⁻$ is the chosen anion as Cl species trigger the corrosion of carbon steel rebars in reinforced concrete structures.¹⁷ As electrolysis proceeds, the cathodic and anodic compartments become increasingly alkaline and acidic, respectively. The precursor $CaSiO₃$ in the anodic compartment is decalcified by H^+ into solid a-SiO₂ (eqn (3)), which remains in the anodic compartment; and Ca^{2+} ions, which diffuse towards the alkaline catholyte due to electric potential and concentration gradient and then precipitate with air-captured $CO₂$ at $pH > 9$ to form e-CaCO₃ in the center and cathodic compartments (eqn (4)).

Fig. 2 (A) Scheme of reactions occurring in the H-cell for weathering of CaSiO₃ at room temperature; cation exchange membrane (CEM), anion exchange membrane (AEM), and bipolar membranes (BPM) are used as ion separators. (B) e-CaCO₃-to-CaSiO₃ product/precursor weight ratio post electrolysis. (C) Schematic of flow cell electrolyzer operation using ion exchange membranes and non-precious metal catalysts. (D) Current density of constant potential hold at 7 V. (E) Schematic of zero-gap electrolyzer and cascade cement recycling process. (F) Current density reached with zero-gap electrolyzer at 3.7 V for 12 distinct runs.

$$
CaSiO_{3(s)} + 2H^{+} \rightarrow Ca^{2+} + SiO_{2(s)} + H_{2}O \tag{3}
$$

$$
Ca^{2+} + 2OH^{-} + CO_{2(g)} \rightarrow CaCO_{3(s)} + H_2O \tag{4}
$$

The series of reactions occurring in the H-cell sums up to eqn (5).

$$
\begin{aligned}[t] 2CaSiO_{3(s)} + 2CO_{2(g)} + 2H_2O &\rightarrow 2CaCO_{3(s)} + 2H_{2(g)} + O_{2(g)} + \\ &\quad 2SiO_{2(s)} \end{aligned} \qquad \qquad \begin{aligned}[t] \text{(5)} \end{aligned}
$$

At cell potentials ranging from 3 to 9 V, the current during electrolysis increases over initial operation due to increased electrical conductivity from H^+ and OH $^-$ generation (Fig. S2A and B, ESI†) and decreases due to decreased electrolyte levels over longtime operation (Fig. S2, ESI†). The resulting mass ratio of e-CaCO₃-to-CaSiO₃ (*i.e.*, calcium conversion rate) linearly scales with charge passed until the theoretical limit of ~ 0.9 (Fig. 2B). Dissolved $Ca²⁺$ ion content in the anolyte increases linearly until \sim 50 000 C has passed, at which point the precursor no longer provides dissolvable Ca, and most of the $Ca²⁺$ ions have diffused to the cathodic compartment (Fig. S4B, ESI†).

Flow-cell configuration. A flow cell electrolyzer is next employed to validate the scalability of this process (Fig. 2C). Two additional compartments are used to separate the solid products from the electrodes with ion exchange membranes to prevent product crossover. A bipolar membrane is used between the anodic and precursor compartments for water dissociation at the anode for non-precious metal (here Ni) alkaline OER catalysts.18 Starting with the precursor placed inside the acidic compartment, the current during chronoamperometric hold (7 V) gradually increases from \sim 30 to \sim 60 mA cm⁻² due to increased H^+ , Ca²⁺, and OH⁻ ion concentration similar to our H-cell operation (Fig. 2D). With a higher H^+ production rate, an e-CaCO₃-to-CaSiO₃ mass ratio of 0.7 is achieved at 24 h, approaching \sim 75% of the theoretical limit and outperforming the H-cell at the same potential applied, which results in a ratio of \sim 0.5.

For more efficient continuous operation, the precursor is placed in an agitated anolyte reservoir, so that continuous operation is achieved by replacing the precursor reservoir once the leaching of $Ca²⁺$ ions completes. Two consecutive batches of 2 g CaSiO₃ yielded a total of 2.65 g e-CaCO₃ (74% Ca converted) after 44 h of electrolysis. Although the flow-cell electrolyzers are operated continuously, membrane fouling due to $CaCO₃$ precipitation on the cation exchange membrane (CEM) decreases current density by 30% from the first to the second cycle (Fig. S5 and S6, ESI†). Such fouling is common in state-of-theart processes; therefore, for flow cell electrolyzers to be viable, engineering of membrane permeability for Ca^{2+} ions and mass transport to rapidly remove precipitates at the membrane is essential.^{19,20}

Zero-gap configuration. To eliminate the membrane degradation and electrode passivation, the weathering was redirected to perform in distinct chemical reactors following the electrochemical production of $HNO₃$ (e-HNO₃) and NaOH (e-NaOH) (Fig. 2E). This approach releases the heavy requirements on ion exchange membrane robustness and the need for an additional

electrolyte compartment, which are proposed by literature on electrified cement production with limestone feeds.^{19,21} By modifying our weathering process as a series of cascade reactions, using a 5 cm^2 zero-gap electrolyzer plus chemical steps in separate reactors, the Ohmic resistance is minimized to 0.5 Ω at room temperature, or 110 m Ω at 80 °C with a catalyst coated membrane (CCM), compared to the \sim 10 Ω of a flow cell electrolyzer, leading to significantly improved energy efficiency (Fig. S7 and S9, ESI†). An order-of-magnitude higher current density of $>$ 300 mA cm⁻² at 3.7 V was achieved with a room temperature zero gap electrolyzer (Fig. S8, ESI†), and 2 A cm⁻² with a CCM zero-gap electrolyzer operated at 80 °C, which translates to 38% energy efficiency and suffices the requirement for commercial-relevancy of > 200 mA cm⁻².²² The present zero-gap electrolyzer remarkably outperforms the flow cell electrolyzer and literature precedents purposed for cement-related productions (Fig. S10 and S11, ESI†). In addition, the zero-gap configuration exhibits improved stability for continuous operation with minimal performance decay after 12 startup-shutdown cycles, surpassing previous cement electrolyzers and showing promises for upscaling. **Paper**

CaSIO₄₄1 211⁻ - Ca^x⁻ : SO₄₅₁⁻ II₄O (3) electrophy ecompetent product on the internation and paperion commons becomes Case¹³⁴⁻¹⁴ grand of the commons are competited on 5 not article is licensed u

As-produced e -HNO₃ and e -NaOH were also utilized to recycle commercial ordinary Portland cement hardened paste (mainly consisting of hydrated calcium silicates), as a model for recycled concrete fines from construction and demolition waste. Ground paste was treated with e-HNO₃ at $pH < 1$ and rapidly exhibited complete dissolution. Si in solution was recovered by adjusting the pH to 4.5 with e-NaOH (pH > 13) and filtered as a-SiO₂ precipitates. Ca²⁺ ions in the solution carbon-mineralize to form e-CaCO₃ with e-NaOH and air CO₂, or solution with captured $CO₂$ (e.g., 0.1 M Na₂CO₃). Performing dissolution and precipitation of cementitious materials allows for precise control over crystallization parameters²³ and avoids heterogeneity in the electrochemical system, as real-world precursor feeds often contain impurities that are detrimental to membrane and electrode stability.²⁴

Product characterization and performance

Phase and morphology. The solid electrochemical products were first characterized with powder X-ray diffraction (XRD) as shown in Fig. 3A. The carbon-mineralized e-CaCO₃ comprises of both vaterite and calcite in the absence of the intermediate portlandite $(Ca(OH)_2)$ phase, which, if uncarbonated in the electrolyzer, would rapidly capture atmospheric $CO₂$ to form $CaCO₃$ during the drying process.²⁵ The existence of two types of CaCO₃ polymorphs suggests various conditions for nucleation and growth. Particularly, vaterite is a metastable polymorph, which may be stabilized by the anion in the present electrochemical reactions.²³ Scanning electron microscopy (SEM) identifies three morphologies in the e-CaCO₃ product: aggregated vaterite ranging from submicron to \sim 10 µm, isolated vaterite spheres up to \sim 90 µm, and angular calcite particles of \sim 10 s µm (Fig. 3Bi–iii). These particle size distributions are on the same order of magnitude as anhydrous Portland cement, suggesting their readiness for direct use as supplementary cementitious materials without grinding or as a replacement for bulky limestone rocks to feed cement production.

Fig. 3 (A) XRD of e-CaCO₃, a-SiO₂, and references. (B) Scanning electron microscopy images of (i) aggregated vaterite, (ii) angular calcite, (iii) vaterite sphere, and (iv) a-SiO₂. (C) Particle size distribution of a-SiO₂ product after 1-day electrolysis at 5 V. Five measurements were recorded on the same sample mixture. (D) and (E) Raman spectrum of e-CaCO₃ and a-SiO₂ products. See Table S5 (ESI†) for assignment of peaks a–h. (F) and (G) STXM image and Ca L-edge NEXAFS spectrum of an e-CaCO₃ particle. (H) and (I) STXM Ca mapping and Ca L-edge NEXAFS spectrum of a-SiO₂ sample

As for the major decalcified coproduct, $a-SiO₂$, XRD demonstrates that a-SiO₂ from both precursors are amorphous according to the diffuse peak around d^{-1} of 0.25–0.3 \AA^{-1} (Fig. 3A). Coproduced a-SiO₂ from CaSiO₃ (initial Si/Ca atomic ratio of 1) leaching retains the morphology of the precursor $CaSiO₃$ and that from dissolution and re-precipitation of recycled cement paste (initial Si/Ca atomic ratio of 0.3) exhibits submicron sizes of the precipitates (Fig. 3Biv and Fig. S12–S14, ESI†). Energy dispersive X-ray spectroscopy (EDX) reveals high Si/Ca atomic ratios of 120 from leaching $CaSiO₃$ and 24 from re-precipitation of recycled cement paste (Tables S6, S7 and Fig. S15, S16, ESI†), promising the superior quality of the present a- $SiO₂$ as a siliceous supplementary cementitious material. The laser diffraction-based particle size characterization demonstrates that precursor $CaSiO₃$ particles range \sim 1–30 µm in size with a mean value of 7.2 µm (Fig. S17, ESI[†]), while the enhanced weathering product a-SiO₂ particles range 0.2–20 μ m in size with a mean value of 6.8 μ m (Fig. 3C), primarily following the particle size distribution of the precursor with a small fraction of finer particles likely caused by fracturing

upon Ca-leaching, agreeing with the measured specific area increase from 0.9 m^2 g^{-1} of CaSiO₃ to 1.7 m^2 g^{-1} of decalcified $a-SiO₂$. Again, both the particle size distribution and specific area of the electrochemical product fall closely to that of anhydrous Portland cement, suggesting their direct usage without the need for grinding or flowability adjustment via adding polymeric superplasticizers, whose manufacturing is intensive in cost, energy, and carbon emissions.²⁶⁻²⁹

Chemical coordination. The chemical orderings of the electrochemical products are further characterized using Raman spectroscopy and spatially resolved near edge X-ray absorption fine structures (NEXAFS) spectroscopy with scanning transmission X-ray microscopy (STXM).

Fig. 3D and E show the Raman spectra of e-CaCO₃ and a- $SiO₂$ with peak assignments given in Table S8 (ESI†). The two $CaCO₃$ polymorphs were differentiated by the position of the peaks corresponding to in-plane bending and symmetric stretching of CO_3^2 ⁻, *i.e.*, peaks d and g at 708 and 1082 cm^{-1} , respectively, as the signatures of the vaterite polymorph versus

peaks e and f at 720 and 1063 cm^{-1} , respectively, as the signatures of the calcite polymorph (Fig. 3D). Both calcite and vaterite-rich e-CaCO₃ regions display peaks below 300 cm^{-1} , which are assigned to the translational and rotational lattice modes. The Raman spectrum of a-SiO₂ exhibits a prominent peak at 1049 cm $^{-1}$, attributed to the symmetric stretching of Q^3 (silicate tetrahedra connected at three corners into the network). This dominant peak reveals full dissolution of precursor CaSiO₃, which contains solely Q² (silicate tetrahedra connected at two corners as a chain), and that acid-stable a-SiO₂ is left with $Q³$ structure that is the most resistant to acid leaching (Fig. 3E).³⁰ This Q^3 signal also suggests the silica product to be amorphous.

STXM-NEXAFS analysis further confirms the presence of the vaterite polymorph in e-CaCO₃ produced (Fig. 3F and G). In the Ca $L_{2,3}$ -edge NEXAFS spectrum, an a_1 peak (L_3 pre-edge) position of 348.1 eV and a ΔE of 1.3 eV between the b₂ peak (L_2 edge) and b_1 peak (L_2 pre-edge) are characteristic of the vaterite phase (Fig. 3G).³¹ The absence of the calcite polymorph under STXM was due to the absorption saturation limit of the thicker calcite particles. The a-SiO₂ resulting from precursor $CaSiO₃$ contains trace amounts of Ca structures indicated by bright spots in Ca element mapping (Fig. 3H), which shows the spatially resolved difference in the absorbance level pre- versus on-Ca L3-edge, with the bright spot indicating large difference thus abundance of Ca (Fig. S18, ESI†). NEXAFS spectrum obtained from the Ca-remaining region on $a-SiO₂$ exhibits multiple L_2 and L_3 peaks, suggesting disordered Ca coordination and low Ca content (Fig. 3I).

Cement performance enhancement via substitution. Both electrochemical products, e-CaCO₃ and a-SiO₂, outperform their existing counterparts – conventional supplementary cementitious materials, *i.e.*, ground limestone and coal fly ash of comparable particle sizes, due to improved purity and reactivity of electrochemical products. We examined blended Portland cements with e- $CaCO₃$ and a-SiO₂ powders partially substituting commercial ordinary Portland cement $(i.e.,$ Portland cement without supplementary cementitious materials) by 5–35 wt% for the hydration behavior using isothermal calorimetry before 1-day curing age and the mechanical strength development starting 1-day age up to 90-day.

The cement hydration reaction is exothermic, where its main heat evolution peak (at \sim 5 h for the ordinary Portland cement reference in Fig. 4A) signifies the rapid growth of calcium silicate hydrate (C–S–H, the primary binding phase and strength contributor of concrete) and of ettringite (the minor phase contributing to the setting of concrete).32,33 During this stage, the hydration products grow and interlock, allowing cement paste to harden and gain strength, which continues to increase asymptotically in the long term with continued hydration. In electrochemicalproducts-blended Portland cement, the main heat evolution peak is \sim 1 h earlier with significantly increased area under the curve, meaning accelerated hydration reaction onset and greater heat flow compared to ordinary Portland cement (Fig. 4A). Both e-CaCO₃ and a-SiO₂ promote Portland cement hydration, $34,35$ accelerating the formation of C–S–H and the setting and strength development of cement. In detail, e -CaCO₃ reacts with aluminates in Portland cement to form calcium carboaluminate hydrates for additional strength gain. e -CaCO₃ also provides nucleation sites to facilitate C–S–H growth.^{36,37} a-SiO₂ consumes portlandite $(Ca(OH)₂)$ – a chemically and mechanically vulnerable phase in hydrated Portland cement – to facilitate further C–S–H formation via the pozzolanic reaction.^{36,37} Paper

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Compared to ordinary Portland cement and conventional limestone powder-and-fly ash-blended Porland cements, the electrochemical-products-blended Portland cements achieve greater compressive strengths at all curing ages, with especially pronounced enhancement at late ages (Fig. 4B), explained by the higher reactivity of metastable vaterite in e -CaCO₃ and the enhanced pozzolanic reactivity of $a-SiO₂$. The enhancement effect of vaterite over calcite has also been found in other cement systems.³⁸ Using the present electrochemical products overcomes the typical problem of low early-age strength of blended Portland cement due to the limited reactivity of common industrial byproduct supplementary cementitious materials, as manifested by the conventional blends compared to ordinary Portland cement in terms of their lower 28-day strengths, the most critical metric in practical construction applications. Moreover, while it is widely agreed that ordinary Portland cement gains the majority of compressive strength by 28 days, 39 the electrochemical-productsblended Portland cement demonstrates 15% continued strength

Fig. 4 (A) Calorimetry of 5 g of (blended) cement paste. (B) Compressive strength of (blended) cement paste using electrochemical products (e-CaCO₃ and a-SiO₂) or conventional supplementary cementitious materials (limestone powder and fly ash) over 90 days. OPC = ordinary Portland cement; LS = limestone; $FA = fly$ ash. (C) XRD of synthesized C_2S and reference.

gain between 28 to 90 days at 30 wt% a-SiO₂ substitution in contrast to just 6% gain for the ordinary Portland cement reference, benefiting from the pozzolanic reactivity of $a-SiO₂$. The $a-SiO₂$ blended cement typically shows higher durability (e.g., resistance to sulfate and acidic environments) due to the consumption of $Ca(OH)_{2}$ and refined pore structure, meaning reduced repair/maintenance and longer lifetime, further decreasing the carbon footprint, especially when normalized by the service life of concrete structures.⁶

Using supplementary cementitious materials as Portland cement substitutes (typically up to 35 wt%) has been successfully applied in the cement industry for over 100 years, penetrating the market for our process to be scaled up.⁴⁰ Electrochemically produced a-SiO₂ is more chemically homogeneous than the highly heterogeneous conventional supplementary cementitious materials (e.g., fly ash and volcanic ash) and emerging supplementary cementitious materials (e.g., municipal solid waste incineration ash). These industrial waste-sourced supplementary cementitious materials experience varying pozzolanic reactivity due to intermixed inert impurities (e.g., mullite and quartz^{41,42}) and are prone to cause cracking failures of concrete due to other detrimental impurities (e.g., aluminum fines).⁶ Thus, our a-SiO₂ addresses the challenging quality control of blended Portland cement incorporating existing supplementary cementitious materials regarding mechanical and durability performances. Therefore, by superseding conventional supplementary cementitious materials with electrochemical products, our approach further contributes to the decarbonization and sustainability of built environment by enhancing the lifetime of concrete structures. Experience on 13 notes As 200, substitution in 14- for contract kind original interactions are the substitution of the control of the control of the creative commons are the substitution of the control of the control of th

Cement production. Besides direct Portland cement substitution, carbon-negative e-CaCO₃ can replace conventional limestone as the primary feedstock for Portland cement manufacturing. We calcined a mixture of quartz and e -CaCO₃ to demonstrate the formation of belite (i.e., larnite; β -2CaO·SiO₂), a major calcium silicate in anhydrous Portland cement. Belite is a reactive mineral and the primary contributor to long-term strength of Portland cement and many alternative cements (e.g., calcium sulfoaluiminate cement) (Fig. 4C and Fig. S19, ESI†). Hence, beyond the broad Portland cement market, our electrochemical approach also applies to specialized and low-carbon alternative cement for niche markets.

Life cycle assessment and techno-economic analysis

A cradle-to-gate life cycle assessment (LCA) compares the carbon footprint of industrial-standard Portland cement manufacturing (Fig. 5A and F) and the electrochemical Portland cement production schemes (Fig. 5B–E and G and H) under three treatment models for different destinations of concentrated $CO₂$ flue gas from the cement kiln: "emission" model assuming $CO₂$ release to the atmosphere, ''circulation'' model assuming circulating $CO₂$ to feed the electrochemical reaction, and "CCS" assuming CO2 capture and storage. For each model, two or three scenarios are considered: ''-conv'' indicates a conventional cement manufacturing scheme business-as-usual; "FF" or "H2" indicate an electrochemical manufacturing scheme using fossil fuel or green

 $H₂$ for cement kiln fueling, respectively. Thus, "FF" scenarios represent a low capital-intensive scheme using our electrochemical products to feed existing Portland cement plants; ''H2'' scenarios represent a moderately capital-intensive scheme additionally using our electrochemical co-product green $H₂$ to fuel a cement kiln. Note S1 (ESI†) provides the energy consumption analysis, and Note S2 (ESI†) provides the full description of our LCA methodology and assumptions. Conventional ordinary Portland cement manufacturing (scenario "emission-conv") incurs global warming potential (GWP) of 0.93 kg $CO₂$ -eq per kg ordinary Portland cement produced (Fig. 6Ai and Fig. S20 and Table S9, ESI†): 0.51 kg from limestone decomposition at the kiln, 0.39 kg from combustion of conventional fuels (mainly coal/coke) at the kiln, and 0.03 kg from electricity or transportation during other processes (e.g., quarrying, transporting, grinding, and in-plant conveying). By blending ordinary Portland cement with conventional supplementary cementitious materials (limestone powder and fly ash) up to 35 wt%, the total GWP is abated by up to 34%.

In all electrochemical manufacturing scenarios, $CO₂$ mineralized e-CaCO₃ decarbonizes cement manufacturing in two-fold: (i) as the Portland cement feedstock, e-CaCO₃ fully decarbonizes the decomposition-induced $CO₂$ emissions; (ii) as a supplementary cementitious material (up to 15 wt% of Portland cement substitution), e-CaCO₃ allows permanent $CO₂$ storage and utilization in concrete, which alone, is a carbonnegative process. Besides, $a-SiO₂$, the carbon-neutral electrochemical coproduct, is directly incorporated as a supplementary cementitious material up to 30 wt% of Portland cement substitution, or together with e-CaCO₃ up to 35 wt%.

In scenario ''emission-FF'' with business-as-usual cement plant operation (Fig. 5B and 6Ai), wollastonite-based, electrochemical-products-blended Portland cement at 0–35 wt% substitution levels achieve 45–66% $CO₂$ abatement. Switching the precursor from wollastonite to recycled cement paste increases GWP moderately, owing to the increased electrical energy demand to fully dissolve recycled cement paste compared to partial dissolution of wollastonite (5.40 vs. 2.86 MJ kg^{-1} e-CaCO₃ produced). In scenario ''emission-H2'' (Fig. 5C and 6Ai), additionally, the co-product green H_2 replaces fossil fuels for kiln heating, further offsetting the rest combustion-induced emissions, leading to approximately net zero emission of the cement kiln and total $CO₂$ abatement by 86–95%. Such fuel switching is practical as the industry has adopted coal/coke as the primary fuel for current kilns for cost saving, superseding the dominance of natural gas as the primary fuel in the 1970s.⁴³ Specifically, the 15 wt% e-CaCO₃ blend achieves the lowest carbon intensity 0.049 kg $CO₂$ -eq per kg blended Portland cement produced, owing to the direct utilization of carbon-mineralized e-CaCO₃ as a partial substitute for cement. Meanwhile, electrochemical production-associated processes, e.g., filtering of solid electrochemical products and pumping of $CO₂$ and H_2 gases, contribute marginally to the overall GWP.

Furthermore, under the ''circulation'' model (Fig. 5D, E and 6Aii), the flue gas with concentrated $CO₂$, from e-CaCO₃ decomposition (and fossil fuel combustion in ''FF'' scenarios), does not exit the system but is cycled as the electrochemical feedstock.

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Fig. 5 (A)–(H) Schematics of comparative LCA scenarios. In scenarios (B)–(E), (G) and (H), precursor refers to wollastonite or recycled cement paste. RE = renewable electricity; DAC = direct air capture; CC(U)S = carbon capture, (utilization,) and storage. Note that to clearly illustrate the electrochemical products and conventional equivalents' flow in the manufacturing process, other cement production raw materials, e.g., clay, are not plotted in the diagrams but accounted for.

Using the concentrated $CO₂$ provides great energy benefits and $CO₂$ abatement by avoiding the entropic penalty of gas separation in direct air capture (DAC) and low efficiency \sim 10%, which invokes DAC energy demand of 2 MJ kg⁻¹ e-CaCO₃ produced. Note that the excess $CO₂$ emissions from fossil fuel combustion in scenario ''circulation-FF'' is considered for CCS, while the net $CO₂$ removal via CCUS in e -CaCO₃ as a cement partial substitute in scenario "circulationH2" is accounted for by introducing small amount of supplemental DAC-CO₂ to keep a steady CO_2 supply. Overall, by curtailing both the cement pyroprocessing $CO₂$ emissions and energy-intensive DAC, wollastonite precursor-based ''circulation'' model leads to 83–92% GWP abatement without cement kiln modification (''FF'') or 89–97% with fuel switching, achieving a minimum of 0.029 kg $CO₂$ -eq per kg blended Portland cement produced.

Fig. 6 (A) Cumulative GWP of four blended cements (100 wt% ordinary Portland cement (OPC); 90 wt% OPC with 10 wt% CaCO₃; 85 wt% OPC with 15 wt% CaCO₃; 65 wt% OPC with 5 wt% CaCO₃ and 30 wt% SiO₂) under eight manufacturing scenarios following three CO₂ treatment models (i) "emission"; (ii) "circulation"; (iii) ("CCS"); legends are shared between (i)–(iii) with grey for conventional manufacturing, green for wollastonite (CaSiO₃) as the precursor, and purple for recycled cement paste as the precursor. (B) Cumulative economic benefits of ordinary Portland cement manufacturing under four TEA-models relative to conventional manufacturing (black dashed line); green for wollastonite, W, as the precursor and purple for recycled cement paste, RC, as the precursor. (C) Radar plots for comprehensive comparison of ''conventional'' and three wollastonite-based ''large business'' scenarios under different CO₂ treatment models; RE = renewable energy; FF = fossil fuel; E = energy

Alternatively, under the ''CCS'' model (Fig. 5F–H and 6Aiii), the flue gas from cement pyroprocessing is not released or circulated but directly captured, transported, and geologically stored. Thus, the DAC-CO₂ intake and mineralization at the electrolyzer lead to carbon-negative cement in all examined ''CCS-FF'' and ''CCS-H2'' scenarios, achieving as low as -0.342 kg CO₂-eq per kg blended Portland cement produced (wollastonite-based scenario ''CCS-H2" at 15 wt% e-CaCO₃ as a partial cement substitute). On the other hand, scenario ''CCS-conv'' is associated with a higher GWP than ''circulation-FF'' and ''circulation-H2,'' suggesting the significant merit of a circular $CO₂$ scheme, which enables almost netzero cement manufacturing at low-to-moderate cement plant modifications and eliminates the need for the long-distance CO2 pipeline transport to geological CCS reservoirs that raise liability and infrastructure rollout issues.

Based on the life cycle inventory analysis of energy consumption, the wollastonite-based electrolyzer operation is associated with 2.9 MJ kg^{-1} e-CaCO₃ produced, according to the thermodynamic energy requirement for the electrochemical production of 20 mol H⁺ required for leaching of Ca^{2+} per kg e-CaCO₃ produced. This amounts to 4.5 MJ kg^{-1} ordinary Portland cement produced at a typical industrial electrolyzer efficiency of 75% (Notes S1 and S2, ESI†). This electrolyzer electrical energy consumption is moderately higher than the equivalent cement kiln thermal energy consumption of 3.0 MJ kg^{-1} ordinary Portland cement produced. Although the energy requirement in the conventional cement manufacturing scheme is lower than our electrochemistry-based counterparts, the inaccessibility to electrified heating in conventional manufacturing schemes always inevitably results in higher GWP. DAC

and CCS derive additional demands of 2.3 and 0.4 MJ kg^{-1} ordinary Portland cement produced, respectively. Overall, scenario ''circulation-H2'' has the lowest energy consumption among all electrochemical manufacturing schemes, achieving a minimum of 3.96 MJ kg^{-1} blended Portland cement produced (Fig. S21, ESI†), comparable to the conventional cement manufacturing cradle-togate energy consumption of 3.84 MJ kg^{-1} ordinary Portland cement produced – but importantly, the former fully supplants the fossil fuel use by renewable energy (renewable electricity and green H_2), accelerating cement manufacturing transition to clean electricity at the industry scale. Using recycled cement paste as the precursor instead, the electricity usage of the electrolyzer increases to 5.40 MJ kg^{-1} e-CaCO₃ produced. Thus, the cradle-to-gate lifecycle energy consumption with recycled cement paste as the precursor increases by \sim 3 MJ kg $^{-1}$ blended Portland cement produced compared to wollastonite as the precursor under equivalent scenarios (Fig. S21, ESI†).

A techno-economic analysis (TEA) for ordinary Portland cement manufacturing via the present electrochemical scheme is conducted in relative to the ''conventional'' manufacturing (Fig. 6B and Table S5, ESI†). Note S3 (ESI†) contains a description of our TEA methodology and assumptions. TEA-model0 is a ''small business'' that operates only the electrochemical production with direct sale of all electrochemical products, e.g., to existing Portland cement plants. Maximum benefits of \$172/t $CO₂$ utilized at the electrolyzer, resulted from: sales of \$170, \$91, and \$23 from a-SiO₂, green H_2 , and e-CaCO₃, respectively; carbon credit saving of \$130; costs of \$41, \$48, and \$23 from raw materials, electrolyzer operational cost, and DAC operation cost, respectively; and \$27 and \$100 from capital expenditure of electrolyzer and DAC, respectively.

TEA-model1, 2, and 3 consider a ''large business'' that runs both the electrochemical production and an existing Portland cement plant switched to green H_2 kiln fueling (\$15/t CO₂ utilized from cement plant fossil fuels saving), consuming e-CaCO₃ within the system boundary and treating green H_2 surplus and a-SiO₂ for sale. These models provide large corporations, who already own Portland cement plants, an economically attractive, near-term pathway towards clean energy transition and decarbonization without establishing remarkably capital-intensive new Portland cement plants. Among the three models for managing flue gas $CO₂$ from cement pyroprocessing, ''circulation'' leads to the greatest economic benefits of maximumly $$77/t$ CO₂ utilized at the electrolyzer. Even though the ''circulation'' model does not allow for claiming carbon credits, its reduction of DAC operational and capital expenses is substantial. Future reduction of the DAC capital expenditure and increase of energy efficiency is expected to increase the economic benefits of ''emission'' and ''CCS'' models, but under the current assumptions, the margins are at least \$55/t and \$72/ t $CO₂$ utilized at the electrolyzer, respectively, lower than the economic benefit of the ''circulation'' model.

While scenarios involving wollastonite as the precursor are mostly economically beneficial at low to intermediate electricity prices, the scenarios involving recycled cement paste as the precursor are generally not as profitable as conventional cement manufacturing. However, the present TEA does not estimate the eliminated cost of waste disposal for recycled cement paste. Moreover, future opportunities with lower DAC and CCS costs and lower renewable electricity prices could increase economic competitiveness. Future studies are suggested to expand the system boundary to comprehensively evaluate the benefits of recycled cement paste versus wollastonite considering their different sources of industrial waste versus natural reserves.

Fig. 6C provides a comprehensive comparison between the models based on LCA and TEA using five metrics, where greater values and the area enclosed indicate model performances superior to the ''conventional'' reference. Total GWP mitigation and fossil fuel GWP mitigation evaluates the percentage of total and fossil fuel-induced GWP reduction relative to the ''conventional'' model, respectively, demonstrating that all three electrochemical production models are able to achieve $>90\%$ mitigation at $>90\%$ of renewable energy usage rate. Particularly, the ''CCS'' model is carbon-negative and reaches 135% total GWP mitigation. Relative energy efficiency evaluates the total energy demand of electrochemical cement manufacturing schemes relative to the ''conventional'' reference: values of \sim 47% for "emission" and "CCS" models and 65% for ''circulation'' model result from the high energy demand for DAC and electrolyzer, despite fundamentally transforming cement manufacturing from fossil fuel-intensive to renewable energy-dominant. Lastly, the ''circulation'' model gains the greatest economic benefits relative to business-as-usual cement manufacturing, while ''emission'' and ''CCS'' models have intermediate to low profit margins, which could be improved with the maturity of carbon management and renewable electricity technologies in foreseeable future. In general, while ''CCS'' model achieves substantial carbon-negativity, ''circulation'' model is overall highly rated for the highest economic benefits, greater energy efficiency, and approximate carbon-neutrality. Paper
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Our sensitivity analysis primarily examines the influence of the renewable electricity's embodied carbon footprints on the total GWP of the electrochemical manufacturing scenarios. Fig. 7A lists the carbon footprint of renewable electricity resources up to 0.04 kg $CO₂$ -eq. per MJ of electricity generated as well as the projected carbon footprint of \sim 0.022 kg CO₂-eq. per MJ for Global 2050 electricity generation based on the forecasted 2050 mixed grid. The data points in Fig. 7A–D mark three representative cases of low, medium, and high renewable electricity carbon footprints from, respectively, land-based wind power, photovoltaic (PV) without and with Li-ion battery energy storage for improved stability and steady energy supply. Note that unlike alternative high-temperature electrolyzer requiring continuous electrical heating, the present electrolyzer can operate entirely at low temperatures ($<$ 100 °C), requiring significantly less strict operating environments. Therefore, the present electrochemical scheme is potentially more compatible with the use of waste heat from cement kilns and lower-carbon footprint, intermittent renewable electricity resources, relaxing the reliance on high-carbon footprint battery energy storage systems thus offering additional $CO₂$ abatement compared to the alternative hightemperature electrochemical cement manufacturing techniques. In Fig. 7A–D, the slope is dictated by the average specific electricity consumption of blended Portland cement production, equaling

Fig. 7 Sensitivity analysis – influence of renewable electricity carbon footprint and electrolyzer energy efficiency on total GWP of conventional and electrochemical cement manufacturing schemes. The data point value is averaged GWP of the four blend designs, and the error bar shows the min-tomax range of the four values. $43,44$

4.3–6.1 and 6.1–7.9 MJ kg^{-1} blended Portland cement produced for wollastonite-based electrolysis at high and low energy efficiency, respectively, or 5.3-7.0 and 7.6-9.5 MJ kg^{-1} blended Portland cement produced if recycled cement paste-based. Overall, the relative trends between the scenarios are consistent across the different electricity sources and precursors: ''CCS-FF'' and ''CCS-H2'' are mostly carbon-negative; ''circulation-FF,'' ''circulation-H2,'' and ''emission-H2'' achieve nearly carbon-neutral at low renewable electricity carbon footprint and maintain $>50\%$ CO₂ abatement even at high renewable electricity carbon footprint; ''emission-FF'' leads up to \sim 50% CO₂ abatement but catch up conventional manufacturing (''emission-conv'') at higher renewable electricity carbon footprint due to the high electrical energy demand from DAC and electrolyzer operation. Nevertheless, the blended Portland cement produced in ''emission-FF'' at high renewable electricity carbon footprint would be still more industrially favorable due to the shortage of coal fly ash for conventional blended cement. Furthermore, the influence of electrolyzer energy efficiency on the total energy consumption is evaluated (Fig. S22, ESI†). At high industrial-electrolyzer energy efficiency (75%), relative energy efficiency of electrochemical cement manufacturing, averaged across the various blended cement designs, ranges 35–66% (wollastonite as the precursor) and 25–42% (recycled cement paste as the precursor), by normalizing to the total energy consumption of conventional cement manufacturing, nevertheless, enabling the cement industry to transition from fossil fuels to renewable resources. Lower efficiency (50% – low industrial-electrolyzer energy efficiency) causes decreases to 29–48% and 20–29%, respectively. On the other hand, improving electrolyzer energy efficiency towards 85% and 95% can achieve respectively 72% and 78% relative energy efficiency through wollastonite-based scenario ''circulation-H2,'' where the electrolyzer electricity demand becomes comparable to conventional cement kiln thermal energy use.

Discussions

Our TEA estimation is conservative as we consider the low-value fate of e-product $CaCO₃$ for cement manufacturing at \$10/t. If the carbon-mineralized $CaCO₃$ is not sourced for liming but treated as a conventional CCS product, this carbon-negative electrochemical product can directly claim carbon credit at \$57/ t or \$79/t CaCO₃ in the U.S. at \$130/t of direct atmospheric $CO₂$ capture and utilization or at \$180/t for direct atmosphere $CO₂$ capture and storage. Although appearing even more profitable, this business model does not compete with the alternative conventional DAC with CCS techniques from the cost perspective due to its greater energy intensity and, more importantly, it sacrifices the substantial economic and decarbonization benefits to the cement/concrete industry and incurs extra concerns for landfill costs and impacts because end users, like concrete, with gigaton capability to the massively produced e-CaCO₃ is rare. It is noteworthy that e -CaCO₃ consumption (as well as the proposed circular $CO₂$ scheme) within cement manufacturing potentially transforms the concrete industry from a gigaton carbon emitter to a gigaton CCUS enabler and furthermore largely alleviates the tremendous challenge and inertia behind large-scale underground carbon storage faced by conventional CCS technologies, not to mention their accompanying issues including long-distance transport, capital expenditure for new infrastructure, leakage/ contamination liability, and more. Meanwhile, implementing the present electrochemical manufacturing scheme at scale can solve the industry-wide shortage of supplementary cementitious materials, particularly, coal fly ash, for conventional blended cements, facilitating net-zero transitions in both the energy and concrete sectors.

A potential alternative solution is to harness e -CaCO₃ for ocean acidification mitigation by enhancing alkalinity and promoting marine direct air capture and storage.^{45,46} Every t of e-CaCO₃ ejected into oceans can capture up to 440 kg of air $CO₂$ to $Ca(HCO₃)₂$.⁴⁷ However, the location of marine electrochemical reactors is limited to onshore or offshore, imposing transportation optimization questions to co-produced a-SiO₂ in order to be distributed to various regions (coastal areas benefit from barging with lower cost and $CO₂$ intensity relative to rail and highway). Presently, this alternative electrolysis-based air CO2 capture scheme lacks optimal prototypes at pilot scale or higher, and the true economic and environmental benefits of this alternative electrolysis-based atmospheric $CO₂$ capture scheme require further validation.

Besides, electrochemically produced $O₂$ can power oxyfuel cement kilns as a superior source than air $O₂$, which requires additional energy for N_2 separation from air. Oxyfuel improves the thermal efficiency of combustion and negates the formation of NO_x , toxic greenhouse gases. After initial fossil-fuel combustion, flue gas is recirculated to mix with pure $O₂$ for subsequent combustion, allowing for more efficient CCUS from higher $CO₂$ concentration $(>90\%)^{48}$ and integration with mature CO₂ capture technologies, e.g., NO_x-sensitive, amine-based CO₂ sorption⁴⁹ – transforming CO₂intensive cement products into gigatons of carbon sink. Moreover, the electrochemical process can separate Ca from Mg impurities (common in precursor minerals but forbidden in Portland cement) via precipitation pH difference, while precipitated $Mg(OH)_2$ can directly capture and store $CO₂$ from air.⁵⁰

Prior to the present study, electrochemical production for cement manufacturing has focused on the electrified production of $Ca(OH)_2$,^{16,19,21} which similarly avoids limestone decomposition-induced $CO₂$ emissions but has the following drawbacks: (i) when using $CaSiO₃$ as the precursor, the standard enthalpy of reaction is 113 kJ mol^{-1} CaSiO₃ higher without introducing CO_2 reactant to form $CaCO_3$ (without CO_2 , $Ca(OH)_2$ is formed instead), meaning greater energy demand; (ii) the utilization of $Ca(OH)_2$ for Portland cement production requires modification of existing industrial cement plants, particularly to the preheater and precalciner prior to the cement kiln, which is unfavored by large cement manufacturing businesses, who prefer modification-free strategies due to capital investment concerns; (iii) $Ca(OH)_{2}$ could absorb air CO_{2} during post-electrolysis processing and transportation, suggesting strict storage requirements for calcination at lower temperature in modified precalciners or compromised efficacy; (iv) the approach does not involve carbonmineralized products for concrete carbon storage, thus infeasible to achieve carbon-neutral/negative. Therefore, the present electrochemical scheme via e-CaCO₃ represents a more efficient, prompt, and preferable solution to cement decarbonization at the industry scale. Paper

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Indeed, clean energy sources with a low embodied carbon footprint and low prices are a prerequisite to the carbonnegativity or carbon-neutrality of the present electrochemical cement manufacturing scheme and positive economic benefit margins compared to business-as-usual conventional cement plants. While the large availability of low-carbon clean energy may have already been achieved in countries like Iceland, Norway, and Sweden, whose power grids are dominated by hydroelectric and geothermal sources, progress in decarbonizing electricity generation is still anticipated for the rest of the world toward approaching the Global 2050 goal, which would widely allow for the present green electrochemical manufacturing scheme with versatility. Ongoing research is dedicated to (1) upscaling the present laboratory gram-scale experiments into industrial-style systems, e.g., larger electrolyzers and stirred reactors and (2) examining the efficacy for multitudes of feedstock profiles including various industrial solid wastes, in order to better assess the efficiencies and challenges and evaluate the

environmental impacts more comprehensively. Besides, due to experimental limitations in measuring the current $CO₂$ capture and utilization processes, the present LCA and TEA use average values of energy demand and cost from references for a generic DAC process. Future studies are encouraged to carefully evaluate the direct capture and utilization of atmospheric $CO₂$ under our electrochemical reaction scheme through experimental electrochemical reaction scheme through approaches to assess the energy consumption, efficiency, operational costs, and capital expenditure for more comprehensive LCA and TEA refinement. Additionally, it is crucial to compare these results with the utilization of circulated concentrated $CO₂$, which demonstrates greater environmental and economic competitiveness in the present study.

Currently, the U.S. is estimated to generate \sim 35–40 Mt y⁻¹ waste hydrated cement paste $(i.e.,$ the reactive component in recycled concrete fines) from construction and demolition wastes generated at >600 Mt y^{-1,51-53} These waste fines have been commonly landfilled after concrete recycling and aggregates reclamation due to the high water demand of hydrated cement in the fines.⁵¹⁻⁵³ Our work encourages research on recycled cement paste separation and treatment by providing a scalable pathway towards its valued use. At full scale in the U.S., the recycled cement paste may be converted to \sim 21–24 Mt y⁻¹ green ordinary Portland cement via the present strategy, accounting for \sim 25% of annual U.S. cement production.⁵¹⁻⁵³ Globally, at full capacity, our strategy can achieve $CO₂$ abatement by 1.2 Gt per y without cement plant modification, 3 Gt per y with green H_2 and CO_2 circulation integrated, or 4.7 Gt per y with further CCS incorporated, equivalent to over 5% of total annual global $CO₂$ emissions.⁵⁴

Conclusion

We demonstrated an electrochemical approach potentially incorporating $CO₂$ circulation as well as capture and storage for carbonneutral/negative cement manufacturing that can be readily integrated to the existing cement industry and rapidly scaled up in the near term. Calcium silicates as naturally abundant rocks and industrial/municipal solid wastes undergo accelerated weathering and capture atmospheric $CO₂$ to form carbon-negative $CaCO₃$ to feed cement kilns, neutralizing the 200-year-old liming routine without modifying the conventional cement manufacturing process. The electrochemical products allow for direct cement substitution for long-term carbon storage and enhanced concrete lifetime; the co-produced green hydrogen provides an economically competitive solution to $CO₂$ abatement for existing capitalintensive cement plants. These results elucidate a promising pathway for the fundamental decarbonization and clean energy transition of the cement industry, which can transform from a gigaton $CO₂$ emitter to a gigaton-scale enabler for renewable energy, direct air carbon capture and storage, and green hydrogen.

Author contributions

X. K. L. contributed to performing experiments, analyzing the data, writing and editing the manuscript. W. Z. contributed to performing experiments, analyzing the data, performing LCA and TEA, writing and editing of the manuscript. B. N. R. contributed to performing experiments, analyzing the data, editing the manuscript. L. C. S. contributed to editing the manuscript. J. L. contributed to conceiving the idea, performing experiments, writing and editing the manuscript.

Data availability

The data supporting this article has been included as part of the ESI.†

Conflicts of interest

The authors declare no conflict of interests.

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