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Riverine photosynthesis influences the carbon sequestration potential of enhanced rock weathering

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As climate mitigation efforts lag, dependence on anthropogenic CO₂ removal increases. Enhanced rock weathering (ERW) is a rapidly growing CO₂ removal approach. In terrestrial ERW, crushed rocks are spread on land where they react with CO₂ and water, forming dissolved inorganic carbon (DIC) and alkalinity. For long-term sequestration, these products must travel through rivers to oceans, where carbon remains stored for over 10,000 years. Carbon and alkalinity can be lost during river transport, reducing ERW efficacy. However, the ability of biological processes, such as aquatic photosynthesis, to affect the fate of DIC and alkalinity within rivers has been overlooked. Our analysis indicates that within a stream-order segment, aquatic photosynthesis uptakes 1%–30% of DIC delivered by flow for most locations. The effect of this uptake on ERW efficacy, however, depends on the cell-membrane transport mechanism and the fate of photosynthetic carbon. Different pathways can decrease just DIC, DIC and alkalinity, or just alkalinity, and the relative importance of each is unknown. Further, data show that expected river chemistry changes from ERW may stimulate photosynthesis, amplifying the importance of these biological processes. We argue that estimating ERW's carbon sequestration potential requires consideration and better understanding of biological processes in rivers.

KEYWORDS

carbon dioxide removal (CDR), enhanced rock weathering, aquatic photosynthesis, enhanced mineral weathering, carbon, alkalinity, dissolved inorganic carbon

Introduction

Carbon dioxide removal (CDR) has become an increasingly prevalent tool in IPCC modeled mitigation pathways that limit global warming to 1.5°C or 2°C relative to the preindustrial period (IPCC, 2022). There is still vast uncertainty and debate around the need, cost, efficacy, scalability, and ethics of using CDR to meet climate objectives (Anderson and Peters, 2016; Rockström et al., 2016; Hollnaicher, 2022). Nonetheless, the likelihood of CDR dependence grows with the ongoing failure of pledged and actualized emissions reductions to meet what is needed to limit warming (Boehm et al., 2022). In response, the CDR industry is growing rapidly as it prepares to meet future demand (Smith et al., 2024; CDR.fyi, 2025).

In the durable CDR landscape—i.e., carbon removal for >1,000 years—enhanced rock weathering (ERW) is an emerging and rapidly growing approach. Currently, 18 ERW companies exist and ~19% of the total tons of carbon sold in the durable carbon market

is attributed to ERW (CDR.fyi, 2025). However, the CDR potential of ERW at scale is uncertain (Buma et al., 2024). Given the proliferating CDR landscape and increasing use of ERW as a carbon offset tool, it is crucial to fully understand the approach's efficacy in order to avoid emitting more carbon than is actually sequestered.

In ERW, silicate or carbonate rocks are crushed and spread on land—usually agricultural fields—where the minerals react with CO₂ and water to form bicarbonate (HCO₃[−]) and cations. For ERW to successfully and durably remove CO₂, the weathering products must reach a long-term reservoir. For most deployments, that reservoir is the ocean where the residence time of carbon is long enough [$>10,000$ years (Middelburg et al., 2020)] to offset fossil emissions, which stay in the atmosphere for thousands of years (Archer et al., 2009).

The journey from fields, where weathering occurs, to the ocean, where sequestration occurs, involves streams, rivers, lakes, and reservoirs. In these freshwater systems, many processes can pull or release weathering products from water, altering the amount delivered to the ocean. Thus far, the ERW community has only considered the ability of abiotic processes—namely carbonate precipitation (Knapp and Tipper, 2022; Zhang et al., 2022; Harrington et al., 2023) and CO₂ degassing (Zhang et al., 2025)—to facilitate the loss of ERW products within rivers.

We contend that biological processes—namely photosynthesis—by submerged plants and algae represent an important but overlooked loss pathway for ERW products in freshwater systems. Quantifying the impact of biological processes on these products is necessary to correctly estimate net CDR associated with ERW.

Here we briefly review ERW as a CDR technology, use existing data from river systems to explore the importance of aquatic photosynthesis on carbon budgets, and consider how large-scale deployment of ERW could feedback and alter biological processes.

Enhanced rock weathering as CDR technology

Carbonic-acid weathering of carbonate and silicate minerals is one of the natural processes that controls CO₂ concentrations in the atmosphere. It is estimated that, currently, natural rock weathering on land removes ~ 1 GtCO₂ yr^{−1} from the atmosphere (Ciais et al., 2014). The goal of ERW is to augment and speed up this natural weathering process by crushing and spreading rocks. Assuming the weathered products enter the ocean, both reactions hold potential to remove CO₂ from the atmosphere on timescales of interest to humans (i.e., hundreds to thousands of years) (Middelburg et al., 2020; Zhang et al., 2022).

Ultimately, carbonic-acid weathering converts CO₂ gas into dissolved inorganic carbon (DIC) and generates alkalinity. We distinguish between DIC and alkalinity because, while the two are tightly coupled, there are processes that can affect one while not influencing the other (Biedunkova and Kuznetsov, 2024). Both are important for CDR. ERW-generated DIC delivered to the deep ocean is the metric of interest for sequestration. Alkalinity affects DIC when waters equilibrate with CO₂ in the atmosphere, but full equilibration does not always occur (Raymond et al., 2013; Campeau et al., 2017; Zhou et al., 2025).

There have been multiple theoretical assessments of the carbon-sequestration potential of ERW on agricultural fields (Kantola et al., 2017; Beerling et al., 2020; Kantzas et al., 2022; Baek et al., 2023) as well as analyses of potential life-cycle carbon budgets associated with mining, transporting, crushing, and spreading rock (Moosdorf et al., 2014; Zhang et al., 2023; Li et al., 2024). These analyses have concluded that ERW on agricultural fields has potential to sequester a meaningful amount of carbon (0.5 to 3 Gt CO₂ yr^{−1}). However, many of these ERW CDR estimates did not account for loss of carbon and/or alkalinity in rivers as ERW products travel from fields to ocean. Estimates that included river losses only considered abiotic processes or were process-agnostic. The impact of river biology was neglected.

Inorganic carbon in river systems

Measurements of total DIC in rivers represent a snapshot of dynamic bio-physical and chemical interactions that are constrained by the buffering capacity of the aquatic environment (Campeau et al., 2017). Upon entering open waters, DIC can be lost to the atmosphere as CO₂ (Butman and Raymond, 2011), incorporated into submerged aquatic biomass through the uptake of either CO_{2(aq)} or HCO₃[−] during photosynthesis (Hotchkiss et al., 2015), remain dissolved and exported downstream, or removed from solution through carbonate precipitation (Tobias and Böhlke, 2011) (Figures 1A–C).

The relative importance of each loss pathway is a function of the physical environment (light, temperature, gas exchange velocity, channel slope, and stream flow), the biological composition of aquatic and riparian primary producers that alter aqueous chemical conditions (pH, alkalinity), and the inputs of geochemically complex groundwater. Perhaps the strongest control on the fate and transport of DIC is hydrology. Water velocity controls the residence time of carbon (Rehn et al., 2023) and the rate of CO₂ degassing (Hall and Ulseth, 2020). Seasonal changes in run-off can drive extreme shifts in the magnitude of CO₂ emissions across stream networks (Conroy et al., 2023).

Here we recognize that rivers drive toward equilibrium but often never reach it (Raymond et al., 2013; Campeau et al., 2017), and are often not at steady state, particularly smaller tributaries (Durighetto et al., 2020; Brinkerhoff et al., 2024). Thus, our discussion considers how riverine biological processes can affect DIC and alkalinity within rivers, but it does not presume how these biologically driven changes in turn affect other processes occurring within the river, such as pH shifts, degassing, and carbonate precipitation.

Aquatic photosynthesis and respiration

Aquatic photosynthesis requires uptake of either dissolved CO₂ or bicarbonate—submerged plants and algae cannot access atmospheric CO₂ for photosynthesis. The bicarbonate carbon fixation pathway dominates over the CO_{2(aq)} fixation pathway in river systems with high bicarbonate concentrations (Aho et al., 2021; Kaijser et al., 2021). In fact, the proportion of HCO₃[−] vs. CO_{2(aq)} affects aquatic plant assemblages. Systems that experienced increases in available HCO₃[−] through agricultural liming shifted species composition to those most

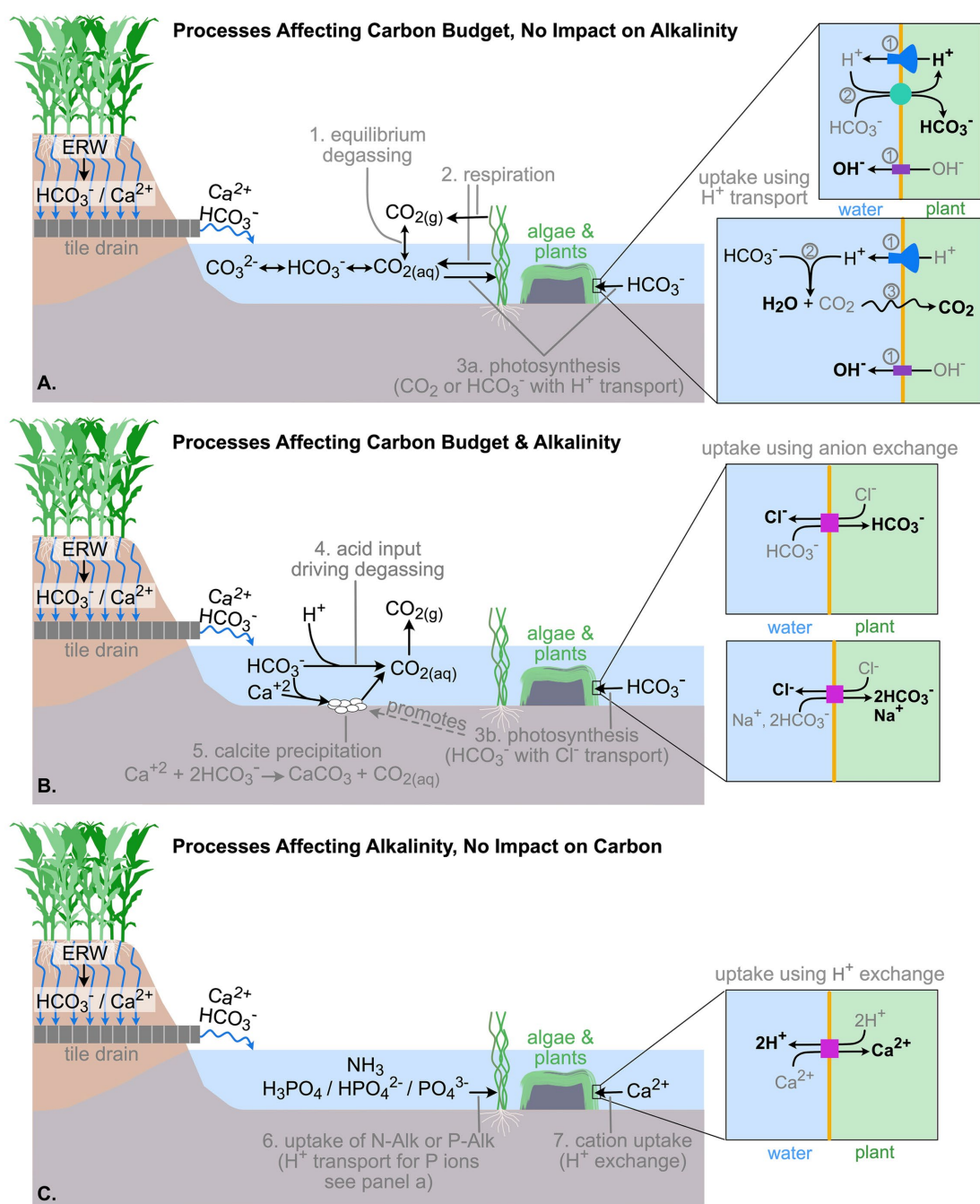


FIGURE 1

Riverine processes affecting weathering products. Processes are numbered for reference. **(A)** Processes that affect dissolved inorganic carbon concentrations, but not alkalinity: 1. Equilibrium degassing, 2. Respiration, and 3a. Photosynthesis where either dissolved CO_2 is used or bicarbonate use is coupled with H^+ transport across the cell membrane. **(B)** Processes that affect both dissolved inorganic carbon concentrations and alkalinity: 4. Acid inputs that drive degassing, 5. Calcite precipitation, and 3b. Photosynthesis where bicarbonate uptake occurs via anion exchange. **(C)** Process that do not affect dissolved inorganic carbon concentrations but do affect alkalinity: 6. Uptake of nitrogen and phosphorous species that contribute to alkalinity (i.e., N-alkalinity and P-alkalinity)—phosphorous ions are transported across the cell membrane using H^+ transport, as illustrated in panel A—and 7. Cation uptake by aquatic plants and algae that is coupled with H^+ transport across the cell membrane.

able to actively take up HCO_3^- for photosynthesis (Brandrud, 2002; Iversen et al., 2019).

Uptake of charged ions, such as bicarbonate, by plants requires transport of other ions across cell walls to maintain internal pH balance and ionic equilibrium. Figure 1 illustrates the various HCO_3^- uptake pathways that are known to exist for aquatic macroalgae and

seagrass (Poschenrieder et al., 2018). Membrane transport of bicarbonate by vegetation is not well documented (Poschenrieder et al., 2018). One set of known uptake pathways involves active transport of H^+ out of the cell followed by passive outward transport of OH^- (Figure 1A). The H^+ then is either co-transported back into the cell with HCO_3^- or it reacts with HCO_3^- and generates $\text{CO}_2(\text{aq})$ that

then diffuses through the cell membrane. Both carbon uptake pathways remove DIC from the river water but have no impact on alkalinity.

The other set of known carbon uptake pathways involves anion exchange across the cell membrane (Figure 1B). A chloride ion exits the cell while either a single bicarbonate or two bicarbonates and a sodium ion enter the cell. This uptake pathway reduces DIC and alkalinity in the surrounding river water.

Aquatic plant productivity has additional indirect impacts on river DIC and alkalinity. First, benthic algae promote calcite precipitation, which decreases both DIC and alkalinity (Figure 1B); the algal mats provide a surface for crystal growth while mat photosynthesis increases pH and the saturation index of calcite in local microenvironments (Hoffer-French and Herman, 1989; Hayashi et al., 2012). Second, associated uptake of nitrogen (NH_3) and phosphorous (H_3PO_4 , H_2PO_4^- , HPO_4^{2-} , PO_4^{3-}) by plants can affect the non-carbonate components of alkalinity. Finally, in addition to nitrogen and phosphorous, plants require cations and uptake of these can be coupled with the release of H^+ out of the cell (Babourina and Rengel, 2010) (Figure 1C). If H^+ is released, it reduces alkalinity in the surrounding water. However, cation uptake is not always coupled with H^+ release. Sometimes, uptake of a given cation, like Ca^{2+} , is coupled with the release of another non-acidic cation, like Na^+ (Babourina and Rengel, 2010), which will not alter alkalinity.

Uptake pathways used by aquatic vegetation, as illustrated by Figure 1, strongly determine the impact plant productivity has on DIC and alkalinity in rivers. However, it is extremely difficult to discern the actual membrane transport pathways used in a field setting; detailed, cell-level investigations are most likely required. In the context of ERW, a conservative approach is to assume that all photosynthetic activity reduces both DIC and alkalinity (i.e., plants are using bicarbonate uptake via anion exchange) and all cation uptake reduces alkalinity (i.e., plants take up cations using H^+ exchange) in rivers.

The magnitude of aquatic photosynthesis relative to DIC transport

Recent advances in sensor technology and modeling approaches have allowed scientists to estimate river gross primary production (GPP) across hundreds of rivers (Appling et al., 2018). We harnessed this dataset along with USGS stream chemistry data (US Geological Survey, 2016) to determine the magnitude of plant productivity and to put its influence in context.

The two datasets shared at least three overlapping time points at 70 USGS gaging stations; the median number of overlapping points was 16. A majority of the stations overlapped with agricultural areas (Supplementary Figure S1) and had balanced carbonate systems (i.e., DIC-to-alkalinity ratios near 1) (Supplementary Figure S2). The combined dataset included 1,600 datapoints spanning from 2007 to 2016. The GPP dataset reports GPP in terms of O_2 concentrations. Although photosynthetic quotients have been shown to vary between 0.8 and 1.2 (Trentman et al., 2023), here we assumed a 1:1 molar ratio between CO_2 and O_2 .

We used a stream-order framework (Figure 2B) to compare the rate of carbon transport to that of photosynthesis. We harnessed the power-law relationship between average stream-order length and median discharge (Downing et al., 2012; McManamay and DeRolph, 2019) (Figure 2A) to calculate a stream-order length for each gage location (Figure 2D). With this length set, we calculated a stream-order photosynthesis fraction for each time point. The stream-order photosynthesis fraction was defined as the ratio between the amount of carbon removed by photosynthesis within the stream-order section to the amount of carbon delivered to the stream-order section by water flow (Figure 2C). Resulting fractions are plotted in Figures 2E,F.

The stream-order photosynthesis fraction for individual time points ranged from less than 0.01% to greater than 1,000%. The median fraction for gage sites ranged from 0.2 to 125%, with most sites falling between 1% and 30% (Figure 2F). These results indicate that photosynthesis can process a notable portion of the carbon moving through a stream order. The fraction processed increased as photosynthesis rates increased and DIC concentrations decreased. In fact, the stream-order photosynthesis fraction and photosynthesis rate had a positive power-law relationship ($R^2 = 0.48$ for individual points, solid black line, $R^2 = 0.30$ for site medians, dashed black line, Figure 2E). The photosynthesis fraction did not systematically vary across stream orders; longer stream-order lengths were not associated with a greater fraction of carbon processed by photosynthesis.

The calculated fractions apply to a single stream order. A river is composed of series of stream orders. For a given carbon input, it is possible to estimate the total fraction taken up by photosynthesis across a series stream orders by assuming a perfectly ideal river network (e.g., Figure 2B) and no cycling of carbon taken up by plants. In this simplified situation, the fraction leaving a given stream order is 1 minus the stream-order photosynthesis fraction (Figure 2C). This is then the amount entering the next higher stream order within which photosynthesis can uptake an additional fraction of the original carbon input. Thus, total fractional uptake of a given carbon input due to photosynthesis across n stream orders can be calculated as: $F_n = 1 - (1-f_1)(1-f_2)(1-f_3)\dots(1-f_n)$, where f_n is the photosynthesis fraction within the n^{th} stream order. With this framework, relatively small fractions within individual stream orders can grow to a large total photosynthesis fraction across a river network. For example, if a single carbon input travels through 10 stream orders and plants take up 5% of the received input in each stream order (i.e., the network has a consistent 0.05 stream-order photosynthesis fraction), by the end of the journey, 40% of the original input will be lost from the water due to plant uptake.

The effect of aquatic photosynthesis on ERW-generated DIC and alkalinity depends on the membrane transport pathway used by the plant during carbon uptake (Figures 1A,B) as well as on the occurrence of processes associated with photosynthesis, such as calcite precipitation (Figure 1B) and uptake of nutrients or cations (Figure 1C). The net effect, in turn, that these biologically driven DIC and alkalinity shifts have on CDR efficacy depends on what happens to the vegetation (discussed in the next section) and the impact these DIC and alkalinity shifts have on other riverine processes (e.g., CO_2 air-water exchange).

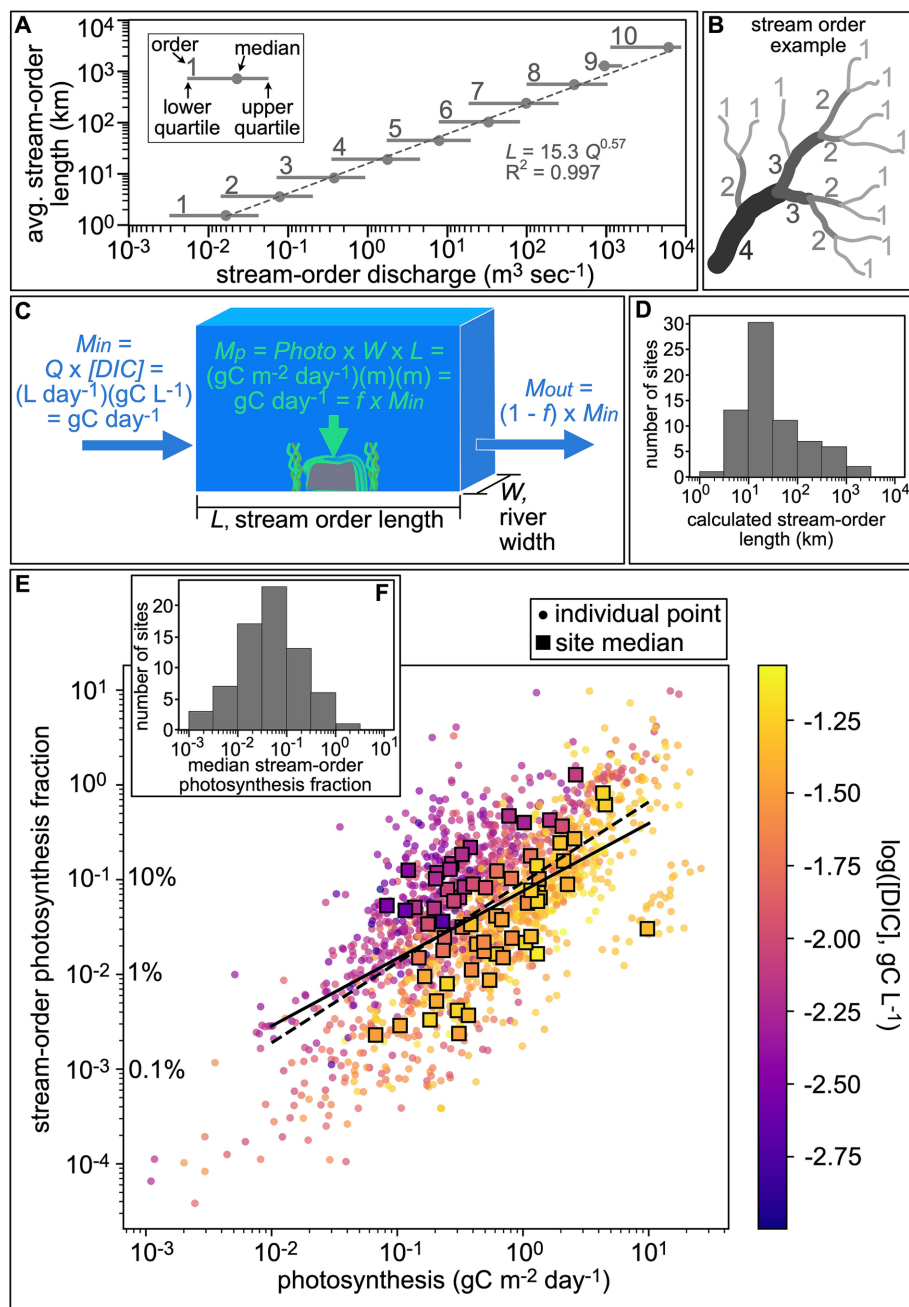


FIGURE 2

Magnitude of aquatic photosynthesis quantified using a stream-order mass-balance framework. (A) Power-law relationship between average stream-order length and stream-order discharge based on published literature (Downing et al., 2012; McManamay and DeRolph, 2019). (B) Example stream-order network. (C) Mass balance calculation producing stream-order photosynthesis fraction (f), which is the ratio between the amount of carbon removed by photosynthesis within the stream-order section (M_p) to the amount of carbon delivered to the stream-order section by water flow (M_{in}). M_p was calculated as the photosynthesis rate or GPP ($photo$) \times river width (W) \times stream order length (L). M_{in} was calculated as discharge (Q) \times DIC concentration. (D) Histogram of stream-order lengths calculated for the dataset using median discharge rates and the relationship in panel A. (E) Resulting stream-order photosynthesis fraction versus photosynthesis rate. Colors indicate DIC concentrations. Circles are individual data points and squares are site medians. The solid line marks the power-law relationship for individual points ($R^2 = 0.5$). Dashed line marks the power-law relationship for site medians ($R^2 = 0.3$). (F) Histogram of stream-order photosynthesis fraction for site medians.

Fate of carbon and alkalinity in aquatic vegetation

Carbon and alkalinity taken up by aquatic vegetation is stored as plant biomass. Biomass is subsequently lost from plants through

respiration, degradation, or consumption by herbivores. The fate of plant carbon and alkalinity differs depending on which of these processes occurs and if the vegetation is covered by water. Aquatic vegetation can cycle between submersion and exposure to the atmosphere as river levels change; remarkably more than half of

USA runoff is sourced from ephemeral streams (Brinkerhoff et al., 2024).

While the vegetation is alive, it releases carbon as CO₂ during respiration. If the vegetation is covered by water, respired CO₂ re-enters the water, returning carbon, but not alkalinity, to the river system (Figure 1A). If instead respiration occurs when the vegetation is not covered by water, respired CO₂ directly enters the atmosphere. Degradation is similar to respiration in that CO₂ is released either back into river water or the atmosphere depending on if the plant matter is covered by water. However, full degradation also releases alkalinity, represented by the excess base cations stored in the plant biomass (Figure 1C), back into the river channel. If instead aquatic biomass is consumed by aquatic or terrestrial herbivores, its carbon and alkalinity are transferred to the consuming organism. At this point, tracking the carbon and alkalinity becomes highly challenging because they can continue moving through either the aquatic or terrestrial food web.

Of these fates, only aquatic degradation fully reverses the removal of DIC and alkalinity from the stream associated with aquatic photosynthesis. The other fates either drive carbon emissions (directly or indirectly) or move DIC and alkalinity out of the stream and into food webs where they are nearly impossible to track. Given these outcomes, we contend that when aquatic plants uptake DIC and alkalinity generated by ERW, a portion of it likely never reaches the ocean. The remainder that does reach the ocean is delayed relative to water flow due to time spent incorporated in aquatic biomass.

These results are relevant for ERW carbon crediting. Credits should only be awarded once the carbon is durably stored or when there is a high degree of confidence that it will not be released before it reaches the durable reservoir. Carbon that is returned to the “fast” carbon cycle, where CO₂ is taken up and released by organic matter, should not count as removal until more is known about the fate of photosynthetic carbon. Standard practice today assumes a ~15% loss of ERW-generated carbon during transport between the field and ocean (Puro Earth, 2024); our analysis indicates the actual losses are potentially higher and hard to predict.

Response of photosynthesis to ERW-induced river chemistry changes

Today, ERW deployments are small and sparse enough that weathering products are essentially undetectable once they reach a river. In this situation, we can assume that ERW products behave and partition similarly to background riverine DIC and alkalinity. However, as ERW scales, it will begin to have measurable impacts on river chemistry, in particular, increases in DIC concentrations and alkalinity, and changes in pH (Oh and Raymond, 2006). Such chemical changes can feedback and impact the biological processes discussed above.

The existing USGS chemistry data show that photosynthesis rates tend to increase as pH and DIC concentrations increase

(Supplementary Figure S3). This trend could reflect a general photosynthesis response to higher DIC concentrations, or a specific response to increased bicarbonate concentrations. However, other explanations exist. For example, pH can affect sorption of elements to sediment. In particular, phosphorous, a key nutrient for biological activity, has been shown to sorb less to sediment and remain more dissolved in water at higher pH values (Temporetti et al., 2019). It is also possible that the data are showing the existing impact of agricultural runoff, which has higher pH, DIC, and nutrient concentrations (Barnes and Raymond, 2009), on aquatic photosynthesis. Regardless of the mechanisms, the analysis indicates that chemical changes in rivers induced by ERW hold potential to enhance the rate at which plants process carbon and alkalinity. The net impact of this chemical-biological feedback on the fate of ERW products in rivers is hard to predict.

Summary

The analyses demonstrate that riverine biological processes should not be ignored by the ERW community. At most locations within our dataset, aquatic photosynthesis currently processes 1 to 30% of DIC moving through a stream-order section (Figure 2). The cumulative proportion processed by plants quickly increases as photosynthesis operates across an entire stream network composed of multiple stream orders. Our analysis also indicated that as ERW scales, the anticipated changes to downstream water chemistry will likely feedback and enhance aquatic photosynthesis. The fate of DIC and alkalinity removed from the stream by aquatic photosynthesis is unclear and should be an area of further study. We contend that, most likely, a portion of DIC and alkalinity transformed into aquatic biomass never makes it into a durable storage reservoir, introducing uncertainty to the estimates of downstream losses that are currently used for ERW carbon crediting.

Data availability statement

Publicly available datasets were analyzed in this study. Data can be found at: <https://www.sciencebase.gov/catalog/item/59eb9c0ae4b0026a55ffe389>; <https://waterdata.usgs.gov/nwis/qw>.

Author contributions

RN: Conceptualization, Data curation, Formal analysis, Funding acquisition, Investigation, Methodology, Project administration, Visualization, Writing – original draft, Writing – review & editing. TK: Formal analysis, Methodology, Validation, Writing – review & editing. SZ: Data curation, Funding acquisition, Writing – review & editing. DB: Conceptualization, Funding acquisition, Writing – review & editing.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fclim.2025.1582786/full#supplementary-material>

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