Simulating carbon capture by enhanced weathering with croplands: an overview of key processes highlighting areas of future model development

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Enhanced weathering (EW) aims to amplify a natural sink for CO₂ by incorporating powdered silicate rock with high reactive surface area into agricultural soils. The goal is to achieve rapid dissolution of minerals and release of alkalinity with accompanying dissolution of CO₂ into soils and drainage waters. EW could counteract phosphorus limitation and greenhouse gas (GHG) emissions in tropical soils, and soil acidification, a common agricultural problem studied with numerical process models over several decades. Here, we review the processes leading to soil acidification in croplands and how the soil weathering CO₂ sink is represented in models. Mathematical models capturing the dominant processes and human interventions governing cropland soil chemistry and GHG emissions neglect weathering, while most weathering models neglect agricultural processes. We discuss current approaches to modelling EW and highlight several classes of model having the potential to simulate EW in croplands. Finally, we argue for further integration of process knowledge in mathematical models to capture feedbacks affecting both longer-term CO₂ consumption and crop growth and yields.

1. Introduction

Enhanced weathering (EW) is a ‘negative emissions technology’ receiving increasing attention because the Intergovernmental Panel on Climate Change’s recently adopted 2°C target may be difficult to attain by reducing fossil fuel emissions alone [1]. Natural chemical weathering is a small sink for atmospheric CO₂, consuming only approximately 0.25 PgC yr⁻¹ [2], with little effect on climate within our lifetimes. This is partly because many soils in warm, humid locations, where weathering should be favoured, are depleted in primary minerals. EW strategies aim to increase CO₂ consumption by adding silicate rock dust to soils in order to strengthen this sink for atmospheric CO₂, thus offsetting anthropogenic carbon emissions from fossil fuels.

Mass balance [3,4] and kinetics [5] calculations and a numerical modelling study [6] all suggest that terrestrial EW strategies could contribute to climate change mitigation by consuming significant atmospheric CO₂. Agricultural land, comprising 15 Tm² globally in 2000 [7], may be particularly suited for EW due to its ease of access and a range of ancillary benefits [8]. Indeed, basalt dust is increasingly suggested as an agricultural amendment on highly weathered nutrient-poor acidic soils in Brazil [9], where the climate is ideal for EW.
The potential of EW to consume CO$_2$ or provide co-benefits is difficult to quantify at a range of scales [10]. For example, the reliability of models for long-term forestry planning depends on input weathering rates [11], but weathering rate estimates vary greatly depending on how they are quantified. Several independent weathering rate estimates are therefore recommended when predicting the outcome of EW or assessing its effects [12]. These include empirical estimates based on different types of field data [12], upper CO$_2$ consumption limits based on mass balance approaches including assessments of downstream effects such as cation flux in surface water drainage from catchments [3], and rates from process-based numerical models (tables 1 and 2 [6, 13–21, 24]).

Currently, few models provide weathering estimates or predict co-benefits for stakeholders considering deployment of agricultural EW on any scale. However, different classes of models have already been developed to understand plant–soil–biogeochemical interactions, and they perform at least a subset of the calculations needed for EW (tables 1 and 2). With further development, they could provide a diverse set of tools for EW planning and assessment. This is particularly desirable given the increasing focus on multi-model ensembles for estimating uncertainties in model outputs, including water and carbon cycling [25, 26] and crop responses to climate change [27–29].

2. Calculating the weathering sink for atmospheric CO$_2$

When atmospheric CO$_2$ dissolves in water at pH values above the acid endpoint for H$_2$O–CO$_2$ proton balance, it forms carbonic acid that dissociates and lowers the pH of water, favouring mineral dissolution (weathering). As weathering of oxide minerals such as cation-bearing silicates progresses, the oxide ion acts as a strong base and consumes protons. The reaction releases a molar charge equivalent of base cations to solution and this increase in alkalinity raises pH; more CO$_2$ dissolves (is consumed) due to the increased solubility of inorganic carbon ions (HCO$_3^-$, CO$_3^{2-}$) at elevated pH in equilibrium with soil pCO$_2$ (gas).

Absolute weathering rates are calculated with mineral-specific rate laws capturing their dependence on reactive dissolved species such as H$^+$, and may vary over several orders of magnitude depending on mineralogy, temperature, pH and solute concentrations. Both protons and hydroxide ions are potent weathering agents, and rates are generally lowest at circumneutral pH where both ions are at relatively low molar activity. Rates increase at lower and higher pH as H$^+$ or OH$^-$ activity increases. Weathering mechanisms dominating at near-basic and higher pH can be important in arid to semi-arid climates, but have negligible effect in humid climates where pH is normally much lower. Solute concentrations and pH depend on hydrology and flushing rates along with weathering rates and nutrient cation and anion cycling. Existing numerical weathering models [6,15,20] calculate weathering and flushing rates and soil water chemistry, but require hydrological and soil texture parameters along with mineralogy and temperature.

Acidity, alkalinity and pH are strongly influenced by base cations and their removal rate by hydrological flushing. The source strength for base cations (mol ha$^{-1}$ y$^{-1}$) arises by multiplying mineral-specific weathering rates (mol m$^{-2}$ mineral s$^{-1}$)
Table 2. Additional model details. DON, dissolved organic nitrogen.

<table>
<thead>
<tr>
<th>soil texture</th>
<th>KIZ</th>
<th>Sheffield</th>
<th>WITCH</th>
<th>PROFILE</th>
<th>SAFE</th>
<th>MAGIC</th>
<th>APSIM</th>
<th>DayCent -Chem</th>
</tr>
</thead>
<tbody>
<tr>
<td>uniform or by layer</td>
<td>uniform</td>
<td>by layer</td>
<td>by layer</td>
<td>by layer</td>
<td>by layer</td>
<td>by layer</td>
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</tr>
<tr>
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<td>static</td>
<td>static</td>
<td>static</td>
<td>static</td>
<td>static</td>
<td>static</td>
<td>static (bulk density dynamic)</td>
</tr>
<tr>
<td>number of layers</td>
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<td>10</td>
<td>6 or 3 (B-WITCH)</td>
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<td>input</td>
<td>1–2</td>
<td>input</td>
<td>up to 10 or organic + groundwater</td>
</tr>
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<td>equilibrium</td>
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<td>input, and negative uptake of N</td>
<td>input, and negative uptake of N</td>
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<td>input, and negative uptake of N</td>
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<td>stems, branches, bark, fine roots</td>
<td>whole plant</td>
<td>root, stems, leaves, shoots, stolers</td>
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</tr>
<tr>
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<td>lumped</td>
<td>one pool</td>
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<td>upper layers</td>
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<td>dynamic root growth</td>
<td>input by layer</td>
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<td>no</td>
<td>no</td>
<td>no</td>
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<tr>
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<td>two bacterial guilds</td>
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<td>no</td>
<td>no</td>
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<td>no</td>
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<td>lumped</td>
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<td>lumped</td>
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<td>input by layer</td>
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<td>no</td>
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<td>lumped</td>
<td>lumped</td>
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(Continued.)
Table 2. (Continued.)

<table>
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<th>Model</th>
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<th>PROFILE</th>
<th>SAFE</th>
<th>MAGIC</th>
<th>APSIM</th>
<th>DayKent -Chem</th>
</tr>
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<td>Mineralogy</td>
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<td>lithological map, uniform</td>
<td>site and layer data</td>
<td>site and layer data</td>
<td>site and layer data</td>
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<td>n.a.</td>
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<td>yes</td>
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<td>n.a.</td>
<td>n.a.</td>
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<td>kinetics</td>
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<td>none</td>
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<td>n.a.</td>
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</tr>
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<td>no</td>
<td>with deposition</td>
<td>with deposition</td>
<td>with deposition</td>
<td>N, P from manure or fertilizer, residues</td>
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<td>Deposition</td>
<td>NH$_4^+$, NO$_3^-$, K$, \text{low molecular weight N, P}$</td>
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<td>none</td>
<td>SO$_4^{2-}$, NO$_3^-$, H$_2$O, NH$_4^+$, NH$_3$ base cations</td>
<td>SO$_4^{2-}$, NH$_4^+$, NO$_3^-$, lumped CaMgK, Ca$^{2+}$, Mg$^{2+}$, K$, Na$, NH$_4$ base cations, NH$_4^+$, NO$_3^-$, SO$_4^{2-}$, Cl$^-$, F$^-$</td>
<td>none</td>
<td>wet and dry H$, Ca^{2+}$, Mg$^{2+}$, K$, Na$, O$, NH$_4^+$, NO$_3^-$, SO$_4^{2-}$</td>
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</tr>
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<td>Pyrite dissolution</td>
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<td>specified based on lithology</td>
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<td>none</td>
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<td>input</td>
<td>none</td>
<td>input annual flux</td>
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<tr>
<td>Secondary minerals</td>
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<td>equilibria only</td>
<td>kinetics</td>
<td>gibbsite equilibria</td>
<td>gibbsite equilibria</td>
<td>aggregated Al(OH)$_3$</td>
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<td>input annual flux</td>
</tr>
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<td>Weathering agents</td>
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<td>none</td>
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</tr>
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<td>kinetics</td>
<td>kinetics</td>
<td>kinetics</td>
<td>kinetics</td>
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<tr>
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<td>implicit</td>
<td>NH$_4^+$, NO$_3^-$</td>
<td>NH$_4^+$, NO$_3^-$</td>
<td>NH$_4^+$, NO$_3^-$</td>
<td>NH$_4^+$, NO$_3^-$, NH$_4^+$, NO$_3^-$, N$_2$O, NOx, N$_2$, DON</td>
<td></td>
</tr>
<tr>
<td>N immobilized by microbes</td>
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<td>no</td>
<td>net uptake</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
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<tr>
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<td>kinetic</td>
<td>kinetic</td>
<td>input</td>
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<td>no</td>
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<td>no</td>
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<td>yes</td>
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<td>no</td>
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<td>NO$_x$</td>
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<td>no</td>
<td>no</td>
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<td>no</td>
<td>yes</td>
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</tr>
<tr>
<td>Al species</td>
<td>13 species</td>
<td>lumped</td>
<td>Al$^{3+}$, Al(OH)$_3^{2-}$</td>
<td>Al$^{3+}$, Al(OH)$_3^{2-}$</td>
<td>Al$^{3+}$, Al(OH)$_3^{2-}$</td>
<td>13 species</td>
<td>none</td>
<td>input</td>
</tr>
<tr>
<td>P species</td>
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<td>lumped</td>
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<td>none</td>
<td>none</td>
<td>none</td>
<td>lumped</td>
<td>input</td>
</tr>
<tr>
<td>Organic acids</td>
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<td>lumped</td>
<td>oxalate</td>
<td>lumped</td>
<td>monovalent</td>
<td>lumped</td>
<td>monovalent</td>
<td>lumped</td>
</tr>
</tbody>
</table>

*aRecommended.  
*bRequired for cropland EW.
by reactive mineral surface area (m$^2$ ha$^{-1}$ or m$^2$ m$^{-3}$ soil). Reactive surface area is, therefore, a critical parameter in determining weathering rates and pH, usually estimated with empirical functions based on soil texture and mineralogy [6,20]. EW models must account for changing surface area as the applied particles dissolve; one method is to assume these particles are shrinking spheres [6,30].

Soil chemistry also depends on the in situ pCO$_2$ that suppresses pH and arises from microbial mineralisation of soil organic matter (SOM). SOM, largely derived from decaying vegetation [31], is a carbon and energy source supporting communities of respiring organisms that also take up nutrient ions. SOM and microbial activity also affect soil structure and the related moisture retention and water permeability. Because soil pore space architecture and water retention slow upward diffusion of respired CO$_2$ in soils, respiration causes order-of-magnitude increases in soil CO$_2$ levels beyond those of the atmosphere. Although degassing will occur when CO$_2$-charged soil drainage water reaches rivers, within the soil profile elevated CO$_2$ accelerates weathering through increased H$^+$-activity.

3. Agricultural soil chemistry depends on nitrogen cycling

Agricultural practices have a profound impact on the water, carbon and nitrogen [32] cycles, and by altering soil chemistry can create acidic, nutrient-depleted conditions affecting weathering regimes and crop growth. Nutrients in harvested plant material cannot enter SOM pools or be mineralised to release alkalinity. Soil acidification leads to Al and Mn toxicity [33], limiting nutrient uptake, root growth and crop yields, so alleviation of this global agricultural problem is therefore, a potential EW co-benefit.

Plants acidify soil by several mechanisms. For example, roots exude low-molecular-weight organic acids providing protons at soil pH values high enough for the acids to be dissociated, and the anions (oxalate/citrate/malate) are chelating ligands accelerating weathering [34]. However, nutrient demand is the main source of acidification from living plants, because protons are exchanged for nutrient cations in the ‘rhizosphere’ soil pore waters surrounding roots and mycorrhizal hyphae [35].

Rhizosphere acidification occurs due to cation uptake in excess of anion uptake, which depends on availability of nitrate (NO$_3^-$) and ammonium (NH$_4^+$). Decomposition of organic N in SOM produces ammonium [35], which is converted to nitrate at a rate depending on soil pH, temperature, moisture and the existing nitrate and ammonium concentrations [36]. Oxidation of organic N to nitrate is accompanied by proton release and acidification of soil. Ammonium may be retained on cation exchange surfaces and on cell walls in the apoplast [37], but nitrate is highly mobile and prone to transport from the soil profile. Because of these transformations, application of ammonium or organic fertilizers also contributes to nitrification and generation of protons remaining in the soil following nitrate leaching [35]. These protons reduce the solubility of inorganic carbon ions, leading to undersaturated pore waters with respect to dolomite (CaMg(CO$_3$)$_2$) and calcite (CaCO$_3$). Dissolution of these ‘liming’ minerals can then release CO$_2$ to the atmosphere [38].

Legumes with symbiotic nitrogen-fixing bacteria in root nodules are less likely to take up nitrate. They generate protons and exchange these for base cations from the soil solution, but the extent of rhizosphere acidification depends in part on the forms of the organic acids produced during carbon assimilation and is therefore species-specific [35].

Because nutrient demand, fertiliser usage, root distributions and rhizosphere processes are crop-specific, proper representation of crop rotations will be critical for capturing EW effects. Teixeira et al. [39] showed that modelled soil nitrogen differed substantially between four simulation methods for wheat, maize and kale rotations.

Models simulating soil chemistry and EW in croplands need to capture the most important processes of the nitrogen cycle, including mineralisation, nitrification and nutrient uptake by plants and microbes. These comprise the four major nitrogen transformations in agricultural ecosystems [32]. Key loss pathways are nitrate transport in drainage waters, harvests and NH$_3$ volatilization, while denitrification, burning, erosion and trace gas (N$_2$O, NO) fluxes are often smaller [32].

4. Adapting existing models to simulate enhanced weathering in croplands

The need to assess potential management strategies for remediating soil acidification has driven the development of nutrient cycling modules within crop models. A well-established example, APSIM [40], simulates soil pH based on proton budgets, calculating ‘excess’ cation uptake along with carbon and nitrogen cycle imbalances due to removal of biomass, organic matter accumulation and leaching of organic anions. APSIM also calculates crop growth and soil water cycling. This model ignores both cation exchange and weathering [41], but if these were incorporated it could provide detailed predictions of CO$_2$ consumption during crop growth because it considers changes in multiple plant alkalinity pools with time. APSIM can simulate management practices including crop rotations [39], most of the major nitrogen-cycling processes [22] and greenhouse gases (GHGs) including N$_2$O [42]. Crop models such as APSIM are, therefore, already well placed to inform emissions policy [43].

Another example, DayCent [44], simulates a similar range of processes [43] and has been combined with the aqueous geochemistry model pH redox equilibrium C (PHREEQC) [45] to simulate GHG fluxes and stream water chemistry in forested and alpine catchments [46]. Although computationally intensive [24], DayCent-Chem requires input weathering fluxes and ignores base cation cycling, assuming uptake is balanced by release from SOM. Nevertheless, with further development this integrated model could be a powerful predictive tool for EW, even capturing the fate of heavy metals that might be released from the applied dust [47].

Acid rain and resulting S and N deposition drove the development of policy-relevant [48] models to estimate the ‘critical loads’ of acid that ecosystems can tolerate without damage; linkages to plant or biodiversity models have allowed calculation of soil chemistry thresholds for different vegetation types [49]. These models typically include equilibria with solid phases, but often require initial weathering rate and nutrient uptake data as inputs. A widely used dynamic example is MAGIC [50], which calculates changes
in soil chemistry with time at catchment scale. MAGIC includes carbonate equilibria, carbon and nitrogen cycling, and adjusts the input exchange capacities and weathering rates to match observed calibration data.

PROFILE is a steady-state soil chemistry model [18,19] with a long history of use in forestry and agriculture, and its dynamic version, SAFE [20], calculates changes in acid–base balance over time. These models have been employed at field and catchment scale and could easily calculate CO₂ consumption due to EW by incorporating appropriate rate and surface area expressions for the applied minerals.

Several process-based weathering models calculate CO₂ consumption and weathering at regional or global scale given climate, soil water flow and respiration, lithology and a suite of weathering rate laws. One such model, WITCH [15], has been applied to large watersheds such as the Mackenzie River basin [16]. Another, the less computationally complex Sheffield weathering model [6], has previously been employed globally in a non-agricultural EW context. These models rely on the output of dynamic global vegetation models (DGVMs) simulating plant productivity, soil respiration and hydrological flows and could be adapted to assess EW in agricultural practices at farm scale.

A recently developed code to assess the interaction of soil structure with vegetation and mineral weathering is the Integrated Critical Zone (ICZ) model incorporated into the Soil and Water Assessment Tool software package [51]. The ICZ model simulates the plant–soil–water system with one-dimensional water flow and reactive transport in the soil profile. It incorporates a vegetation model, SOM dynamics, nutrient transformations of N and P, mineral weathering kinetics based on the SAFE code noted above, and geochemical speciation equilibria between solutes, mineral and gas phases and ion-exchange surfaces [13,14]. The ICZ includes a sub-model for changes in soil structure with changing biological activity, organic carbon inputs and mineralization, and the resulting changes in bulk soil hydraulic and transport parameters feedback to the equations describing hydrological flow and transport processes. Simulating EW with the code still requires process descriptions accounting for changing reactive surface area as rock powder dissolves; e.g. shrinking core equations.

5. Capturing dynamic plant–soil-enhanced weathering interactions

Weathering depends on productivity, hydrology and nutrient cycling, but EW-type treatments have been shown to affect plant growth [52] and water cycling [53] over several years, and phosphorus availability within several months [54]. These feedbacks indicate that EW should be coupled to modules calculating crop growth, nutrient cycling and soil hydrology.

Models integrating EW with crop productivity and nutrient limitation [55] are especially desirable. Although some DGVMs have been adapted for food and biofuel crops [22], they represent nutrient cycling to a varying degree [56], limiting their ability to predict productivity under climate change [55]. Likewise, crop models explicitly representing nitrogen stress [57] suggest it will be significant for food crops as climate changes. Phosphorus limitation, a common problem in highly weathered tropical soils, is difficult to quantify but is being incorporated into DGVMs [58], albeit without weathering kinetics. However, P is derived from weathering, and P cycling deserves priority because basalt soil amendments increase available soil P with a likely feedback on crop productivity [54].

Separate plant and SOM pools for each nutrient along with root and nutrient uptake dynamics [59] would allow calculation of changing chemistry as a crop matures. Non-structural nutrients such as potassium may be released from the fastest-decomposing litter pools, while nitrogen may be retained in more recalcitrant SOM pools.

EW might also change the soil properties governing hydrology within DGVMs. Overall mineral composition and particle size distribution of the applied material change with time because of the different weathering rates of the constituent minerals. Soil porosity may change following application if individual or aggregated [60] particles change micropore characteristics; particles may be flushed to deeper layers or lost in run-off when they become smaller than micropores. Particle transport is, therefore, a priority for development and inclusion in EW models with associated hydrologic feedbacks on crop production.

6. Conclusion

EW models adopted for croplands need to capture the major water, nutrient and GHG fluxes to realistically simulate CO₂ capture by weathering. These fluxes do not occur in isolation; they interact and lead to feedbacks across a range of timescales. Critically, water flow and the concentration of weathering agents such as H⁺ in the soil solution are strongly controlled by crops. When combined with agricultural practices, these processes can lead to persistent soil acidity, suggesting a co-benefit for farmers because the base cations released by EW increase the alkalinity of the soil.

Existing crop models typically ignore weathering, but may simulate nitrogen cycling, proton balance and GHGs. On the other hand, weathering models driven by coupling with DGVMs may not capture enough cropland processes. The ICZ model couples a vegetation model with mineral solubility for carbonate phases and silicate mineral weathering kinetics. Nevertheless, a diverse set of modelling tools of varying complexity and taking a variety of approaches would likely be more useful for assessing the uncertainties and relative importance of EW parameters and processes than any one model.

One of the most important criteria for quantifying EW dynamics that emerges from this review of modelling approaches is representing the feedbacks between crop growth and mineral weathering kinetics. Understanding the mechanistic process linkages between EW and climate mitigation requires coupled calculations, on intra-annual time scales of relevance for agricultural practices and on inter-annual time scales for climate change mitigation. A priority for the development and assessment of EW across all scales as a possible climate mitigation option is to develop this coupled modelling capability and validate it with appropriate field and laboratory experiments.

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