Evaluating sensitivity of silicate mineral dissolution rates to physical weathering using a soil evolution model (SoilGen2.25)

E. Opolot and P. A. Finke

Department of Geology and Soil Science, Ghent University, Krijgslaan 281, 9000 Ghent, Belgium

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Correspondence to: E. Opolot (Emmanuel.Opolot@ugent.be) and P. A. Finke (peter.finke@ugent.be)

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Abstract

Silicate mineral dissolution rates depend on the interaction of a number of factors categorized either as intrinsic (e.g. mineral surface area, mineral composition) or extrinsic (e.g. climate, hydrology, biological factors, physical weathering). Estimating the integrated effect of these factors on the silicate mineral dissolution rates therefore necessitates the use of fully mechanistic soil evolution models. This study applies a mechanistic soil evolution model (SoilGen) to explore the sensitivity of silicate mineral dissolution rates to the integrated effect of other soil forming processes and factors. The SoilGen soil evolution model is a 1-D model developed to simulate the time-depth evolution of soil properties as a function of various soil forming processes (e.g. water, heat and solute transport, chemical and physical weathering, clay migration, nutrient cycling and bioturbation) driven by soil forming factors (i.e., climate, organisms, relief, parent material). Results from this study show that although soil solution chemistry (pH) plays a dominant role in determining the silicate mineral dissolution rates, all processes that directly or indirectly influence the soil solution composition equally play an important role in driving silicate mineral dissolution rates. Model results demonstrated a decrease of silicate mineral dissolution rates with time, an obvious effect of texture and an indirect but substantial effect of physical weathering on silicate mineral dissolution rates. Results further indicated that clay migration and plant nutrient recycling processes influence the pH and thus the silicate mineral dissolution rates. Our silicate mineral dissolution rates results fall between field and laboratory rates but were rather high and more close to the laboratory rates owing to the assumption of far from equilibrium reaction used in our dissolution rate mechanism. There is therefore need to include secondary mineral precipitation mechanism in our formulation. In addition, there is need for a more detailed study that is specific to field sites with detailed measurements of silicate mineral dissolution rates, climate, hydrology and mineralogy to enable the calibration and validation of the model. Nevertheless, this study is another important step to demonstrate the critical need to couple different soil forming processes with chem-
evaluated weathering in order to explain differences observed between laboratory and field measured silicate mineral dissolution rates.

1 Introduction

Silicate mineral weathering is the major source of most plant nutrients in soils (Carey et al., 2005; Hartmann et al., 2014) and it is probably the foremost process controlling soil production rates (Anderson et al., 2007; Dixon and von Blanckenburg, 2012). Silicate mineral dissolution rates also have implications on acidification in forest soils (Phelan et al., 2014) and carbon sequestration (Beaulieu et al., 2011; Goddéris et al., 2013; Pham et al., 2011). Quantifying the rates of silicate mineral dissolution is therefore of utmost importance to answer many environmental questions such as the surface and groundwater composition, the supply of macronutrients (e.g. K and Ca) in forests and the neutralization of acid precipitation (Ganor et al., 2007).

Indeed a lot of work has been devoted to quantifying silicate mineral dissolution rates using both laboratory experiments (Blum and Stillings, 1995; Chou and Wollast, 1985; Knauss and Wolrey, 1986; Lee et al., 1998; Stillings and Brantley, 1995; Zhu and Lu, 2009) and field experiments (Maher et al., 2009; Parry et al., 2015; White and Brantley, 2003; White et al., 1996; White, 2003, 2002). One common conclusion from most of these studies is that a discrepancy of up to 5 orders of magnitude (Oliva et al., 2003; Parry et al., 2015; White et al., 1996; Zhu, 2005) does exist between laboratory and field weathering rates. There seems to be a general consensus that these differences may be explained by (i) changes in fluid composition (ii) changes in primary mineral surfaces (reactive sites) (iii) the formation of secondary phases (iv) efficiency of solution/mineral contact and, (v) the composition of the soil solution in micro pores. White (2002) grouped these factors into two; intrinsic (e.g. mineral composition, surface area) and extrinsic factors (e.g. solution composition, climate, biological processes). All these five factors could slow field weathering rates compared to laboratory experiments where most of the physical, biological and chemical conditions can
be constrained (White and Brantley, 2003). In general the integrated effects of these intrinsic and extrinsic factors are complex and certainly difficult to capture both in the field and in the laboratory experiments. Moreover uncertainty in the extrinsic factors that occurred and varied in the past is difficult to constrain in experiments (Moore et al., 2012; White and Brantley, 2003).

Modelling approaches enhanced by an understanding of silicate kinetic rates and mechanisms from the experimental works are therefore essential to facilitate in the quantification of silicate dissolution rates (Beaulieu et al., 2011; Goddéris et al., 2006; Hellevang et al., 2013; Roelandt et al., 2010; Stendahl et al., 2013). However, in only a few of these modelling approaches (Goddéris et al., 2006; Maher et al., 2009; Moore et al., 2012) has the integrated effect of some intrinsic and extrinsic factors on silicate mineral dissolution rates been investigated. There is need for mechanistic models capable of simulating the integrated effect of physical, biological and chemical soil forming processes on chemical weathering rates. Such coupling will give the possibilities to determine the role played by intrinsic and extrinsic factors and explain the differences in dissolution rates observed in the laboratory and field experiments (Goddéris et al., 2006; Hartmann et al., 2014; Moore et al., 2012).

The objective of this work is to explore the integrated effect of other soil forming processes on the silicate mineral dissolution rates with particular emphasis on physical weathering. The relationship between particle size distribution and chemical mineral weathering is well known. Holding other factors constant (e.g. pH), the smaller the grain size the larger the surface area per unit mass and consequently the higher the rate of chemical weathering (Hartmann et al., 2014). In most cases, a constant grain size distribution has been assumed when estimating weathering rates, this assumption could be invalid especially when looking at longer time scales. This contribution applies a SoilGen model (a model that simulates evolution of soil properties as a function of several interactive soil forming processes including water flow, chemical weathering, physical weathering, carbon-cycling, cation exchange, clay migration, nutrient uptake by plants, bioturbation and leaching) to evaluate the sensitivity of silicate mineral dis-
solution rates to other soil forming processes. Specific objectives are to (i) assess the effects of parent material composition on the silicate mineral dissolution rates, (ii) to assess model sensitivity of chemical silicate mineral dissolution rates to change in soil texture, (iii) to assess the effect of physical weathering of primary minerals on their dissolution rates, (iv) to assess the effect of interactive soil forming processes on silicate mineral dissolution rates and (v) to compare our modelled silicate mineral dissolution rates to rates reported in literature.

2 Materials and methods

2.1 Study area

This is a sensitivity test study that is not specific to any location. However choice was made to do this study in the forested loess soils, in the Zonian forest, Belgium (50°46′31″N, 4°24′9″E) primarily because the soil forming processes (clay migration, physical weathering, decalcification, carbon-cycling) in the model have already been calibrated to this site (Finke and Hutson, 2008; Finke, 2012; Opolot et al., 2015; Yu et al., 2013). In addition, the measured soil data (Finke, 2012; van Ranst, 1981) and other reconstructed model input data (Finke and Hutson, 2008) were readily available for this site.

2.2 Research set up

As the objective (1 and 2) of this study is also to assess the sensitivity of silicate weathering rates to differences in parent material and soil texture, the research set up (Fig. 1) is such that different initial textures and mineralogy are captured. Therefore rather than using texture and soil mineralogy measurement from the study site, 6 different texture points were randomly selected from the USDA textural triangle to represent the initial soil texture. Three different parent materials (granite, basalt and peridotite) were selected in such a way that slow (felsic igneous rock), moderate (mafic igneous rock)
and fast weathering (ultramafic) rocks were taken into account. The geochemical data (oxide weight composition) typical of granite, basalt and peridotite was obtained from literature (Harris et al., 1967; Hartmann et al., 2013) and the mineralogical compositions were estimated from these geochemical data using the normative mineralogy calculation method (Cross et al., 1902; Kelsey, 1965). Only primary minerals were considered at this stage and their weight compositions were rescaled to sum up to one.

At this stage two scenarios (with physical weathering, PhyWE and without physical weathering, NoPhyWE) were defined but in two different model setups (Model A and Model B; Fig. 1). Model setup 1, hereafter referred to as Model A is intended to simulate majorly the effect of change in particle size (due to physical weathering process) on silicate weathering rates and therefore the effects of other processes (clay migration, carbon-cycling, bioturbation) on silicate weathering rates are minimized by deactivating these processes in this model set up. Model setup 2, hereafter referred to as Model B was intended to simulate the interactive effect of other soil forming processes (including physical weathering, clay migration, carbon-cycling, plant uptake, bioturbation) on silicate weathering rates and therefore all these processes are active. The soil forming processes included in the SoilGen and input data are briefly discussed in the subsequent sections. In total, 72 cases were run in the SoilGen model (i.e 2 set ups *2 scenarios *6 texture points *3 parent materials).

The output parameters from the model include soil texture (% mass fraction of clay, silt and sand), pH, base saturation, weathering indices, mass of each mineral remaining, etc. For this study the outputs of interest extracted included pH, clay mass fraction and mineral mass. The mass of each mineral remaining after the simulation period (15 000 years) was used to calculate the respective dissolution rates of each mineral and was the basis for the sensitivity analysis as will be explained in the subsequent sections.
2.3 The SoilGen model

SoilGen model is a 1-D model designed to simulate time – depth evolution of soil properties as a function of interactive soil forming processes (majorly driven by the soil forming factors (“CLORPT”): climate, living organisms, relief, parent material, time). The governing processes in the model include unsaturated water flow simulated using Richard’s equation, heat flow described following heat flow equation, and solute transport described by convection-dispersion equation (CDE). Other processes described in the model include carbon cycling (based on RothC 26.3 approach; Jenkinson and Coleman, 1994 but applied per soil compartment in SoilGen), clay migration, bioturbation, chemical and physical weathering, and biogeochemical recycling by plants (Finke and Hutson, 2008; Finke, 2012). The model has successfully been applied in several case studies e.g., to simulate clay migration in forest and agricultural land uses in northern France (Finke et al., 2015), to reconstruct soil properties (texture, bulk density, calcite content, pH and OC %) for archaeological land evaluation (Zwertvaegher et al., 2013), to estimate the effect of bioturbation (due to tree fall) on soil horizon thickness (Finke et al., 2013), to test how well soil development would be described by modelling (Sauer et al., 2012), to explain the effect of slope and exposition on soil properties and decalcification depth (Finke, 2012), and to assess the effect of varying climate on calcareous loess soils (Finke and Hutson, 2008). In general, the coupling of major soil interacting processes (biological, chemical and physical) in the SoilGen model makes its application domain versatile. State of art overview of the SoilGen model including simulated processes, data requirements, calibration and quality test studies have been presented in (Opolot et al., 2015) and are not discussed in details here. Physical and chemical weathering processes that form the basis of this study are however explained in details in the subsequent subsections.
2.3.1 Physical weathering

Physical weathering in SoilGen is defined as the stochastic process through which soil particles are split up into smaller particle sizes due to strain caused by temperature gradients. The process of physical weathering consequently leads to the reduction in grain size, producing material in the clay fraction that may be moved by clay migration (Finke, 2012).

As a starting point, the fine earth fraction is divided into particle size classes with boundaries at 2048-1024-512-256-128-64-32-16-8-4-2 µm. It is assumed that all particles are cubes and have an edge size halfway between the class limits: 1536, 768, 384, 192, 96, 48, 24, 12, 6, 3, and 1 µm. In principle, each particle has to be split in half up to 7 times to attain 8 equally sized particles in the next smaller particle size class. The splitting probability of each particle, $P_s$, is assumed to follow Bernoulli process and depends on the temperature gradient over a certain time interval, $dt$ (Finke, 2012):

$$P_s = \begin{cases} 
    P_{s,\text{max}} & \text{if } \frac{dT}{dt} > B \\
    P_{s,\text{max}} \times \frac{dT}{dt} & \text{if } \frac{dT}{dt} \leq B 
\end{cases}$$  \quad (1)

where, $B$ is a threshold temperature gradient over $dt$ where $P_{s,\text{max}}$ is maximal, $T$ is the temperature and $P_{s,\text{max}}$ is the maximal split probability. $P_{s,\text{max}}$ and $B$ are normally subjected to calibration in conjunction with parameters that describe clay migration (Finke et al., 2015).

The expected number of the potential splitting events, $E(N)$ that are needed to achieve successful splits, $m$ (i.e., $m = 7$) are assumed to follow the negative binomial distribution and are defined as:

$$E(N) = \frac{m}{P_s}$$  \quad (2)
Thus the number of grains, $S$ in any particle size class $i$ that is split in time $dt$ is obtained from Eq. (3):

$$S_{i,dt} = \min\left(k_{i,t-dt}k_{i,t-dt}/E(N)\right)$$  \hspace{1cm} (3)

where, $k_{i,t-dt}$ is the number of grains in particle size class at the start of $dt$ and $k_{i,t}$ is defined as:

$$k_{i,t} = k_{i,t-dt} - a \times S_{i,dt} + b \times 8 \times S_{i-1,dt}$$  \hspace{1cm} (4)

where, $a = 0$ for clay fraction ($i = 11$) and $a = 1$ else; $b = 0$ for the coarsest sand fraction ($i = 1$) and $b = 1$ else (Finke, 2012).

It has to be noted that physical weathering in SoilGen is assumed to be caused solely by temperature fluctuations and other mechanical processes that result into the breakup of bedrock (e.g. by plant roots) are not modelled. The splitting of gravel-sized particles is yet to be included in the description of physical weathering and this currently limits the use of the SoilGen model to unconsolidated, non-gravelly deposits (Opolot et al., 2015).

### 2.3.2 Chemical weathering

The weathering mechanism that is implemented here (i.e., in SoilGen 2.25) is different from the mechanism that has been used in the previous version of the SoilGen model (SoilGen 2.24). The previous chemical weathering module (unweathered phase) of SoilGen considers four most common primary minerals (Anorthite, Chlorite, Microcline, Albite) that respectively release Ca, Mg, K, and Na. Congruent weathering of Anorthite, Chlorite, Microcline and Albite release Al. Detailed mechanism is already presented in (Opolot et al., 2015; Sect. 2.1.2). In general the approach is based on the acidification models and takes only a few minerals into account. There was need to extend this module to allow simulation of chemical weathering of a wider range of primary minerals such that more chemical species may be simulated (Opolot et al., 2015).
extended chemical weathering system presented here is based on the transition state theory and similar to the approaches already presented in other weathering models e.g. Sverdrup and Warfringe (1995) and Goddéris et al. (2006).

The release rate of cation, $i$ from all the $k$ minerals, $r_{i,k}$ (mol m$^{-2}$ s$^{-1}$) can be computed as:

$$ r_{i,k} = \sum_{k=1}^{N} A_k v_{i,k} r_k m_k t $$  \hspace{1cm} (5)

where $A_k$ (m$^2$ mol$^{-1}$) is the specific surface area of the $k$th mineral, $v_{i,k}$ (−) is the stoichiometric number of the $i$th element in mineral $k$ and $r_k$ (mol m$^{-2}$ s$^{-1}$) is the dissolution rate constant of the $k$th mineral. $m_k$ is the amount of the $k$th mineral in the parent material expressed in (mol m$^{-3}$ soil) and $t$ (m) is the thickness of the soil compartment (= 0.05 m in SoilGen).

The total surface area of soil minerals, $A_j$ (m$^2$ g$^{-1}$) can be obtained based on the percentage fractions of sand, silt and clay (Eq. 6; Sverdrup and Warfvinge, 1993). The individual reactive area $A_k$ is obtained as a product of weight composition of $k$th ($k_{\text{comp}}$) mineral and $A_j$. $A_k$ is again multiplied by the relative formula mass of the mineral ($k_{\text{RFM}}$, g mol$^{-1}$) to give the mineral area, $A_k$ in m$^2$ mol$^{-1}$ (Eq. 7):

$$ A_j = 8x_{\text{clay}} + 2.2x_{\text{silt}} + 0.3x_{\text{sand}} + 0x_{\text{coarse}} $$  \hspace{1cm} (6)

Coefficients 8, 2.2 and 0.3 represent the specific surface areas (m$^2$ g$^{-1}$) of clay, silt and fine sand (< 256 µm) sized particles, respectively.

$$ A_k = A_j \times k_{\text{comp}} \times k_{\text{RFM}} $$  \hspace{1cm} (7)

The use of the above texture function (Eq. 6) is based on the assumptions that the particle size fractions of clay, silt, sand and coarse add up to 1 i.e 100% (Sverdrup and Warfvinge, 1995) and that the particle grains are of the same shape over time. It has to be noted here that the PROFILE model (Sverdrup and Warfvinge, 1995) has
considerably been criticized (e.g. Hodson et al., 1997), partly because of the use of this equation to estimate the mineral reactive area. The accurate estimation of reactive mineral surface area in natural environments is still a subject of considerable debate (e.g. Hodson et al., 1997; Brantley et al., 2008). Nevertheless the use of Eq. (6) along with experimental dissolution rates normalized to the BET surface area, allows for the good estimation of mineral surface area in natural environments (Sverdrup and Warfvinge, 1995). This function is therefore still widely used as a reasonable first estimate of mineral surface areas (Godéris et al., 2006; Gudbrandsson et al., 2011; Koptsik et al., 1999; Koseva et al., 2010; Phelan et al., 2014; Stendahl et al., 2013; Violette et al., 2010; Whitfield et al., 2010).

The dissolution rate of most silicate minerals, $r_k$ at far from equilibrium conditions is calculated as a function of pH (Eq. 8), based on laboratory kinetic laws derived from the concept of transition state theory (Eyring, 1935; Brantley et al., 2008).

$$r_k = k_H a_{H^+}^n + k_{H_2O} + k_{OH} a_{OH^-}^m$$

(8)

where $k_H$ and $k_{OH}$ are mineral dissolution rate constants at acidic and basic conditions, respectively. The parameters $k_H$ and $k_{OH}$ have to be corrected for temperature (Eqs. 9 and 10). $a_{H^+}$ and $a_{OH^-}$ are activities of $H^+$ and $OH^-$, respectively and superscripts $n$ and $m$ denote the reaction orders. $k_{H_2O}$ is a parameter describing the dissolution rate at neutral pH and was not considered in the implementation because at neutral conditions the dissolution rate of silicates is so slow that this term makes an insignificant contribution to the overall silicate dissolution rate (Brantley, 2003).

$$\frac{k_H}{k_{H25}} = \exp \left[ \frac{k_{E_a H}}{R} \left( \frac{1}{298.15} - \frac{1}{T} \right) \right]$$

(9)

$$\frac{k_{OH}}{k_{OH25}} = \exp \left[ \frac{k_{E_a OH}}{R} \left( \frac{1}{298.15} - \frac{1}{T} \right) \right]$$

(10)
where $k_{H25}$ and $k_{OH25}$ are measured dissolution rate constants at 25°C (298.15 K), $k_{EaOH}$, $k_{EaH}$ (KJ mol $^{-1}$ K$^{-1}$) are the activation energies of a $k$th mineral at acidic and basic conditions, respectively and $R$ is a gas constant (0.00831446 KJ mol $^{-1}$ K$^{-1}$). $T$ is absolute soil temperature (K) and it is simulated in the model.

2.4 Model input data

The SoilGen model was designed keeping in mind the generally accepted paradigm that soil is a function of soil forming factors; “CLORPT” (Jenny, 1941). Therefore the model uses these factors either as initial conditions (e.g mineralogy, texture) or boundary conditions (e.g. climate, vegetation, bioturbation, slope and exposition). The initial conditions specify to the model the initial soil properties at the beginning of the simulations and are usually assumed to be equal to the soil properties from the analysis of samples taken from the C – horizon. Initial texture and mineralogy used in this study are shown in Tables 1 and 2, respectively. Other initial soil properties (e.g. initial OC %, bulk density, solution and exchange surface chemistry), and boundary conditions (Fig. 2) were taken from Finke (2012).

2.5 Calculating average silicate dissolution rates

Silicate mineral dissolution rate usually reported in units of mol m$^{-2}$ s$^{-1}$ is defined as the amount of mineral (moles) that is released in form of constituent elements per unit area (e.g., cm$^2$, m$^2$ or ha) or volume (cm$^3$, m$^3$) over a given period time. Similar to the approach used in (White and Brantley, 2003), congruent weathering was assumed and the moles of each cation released during silicate mineral dissolution was based on the stoichiometric coefficient of that particular element in the mineral. To calculate the dissolution rate of a given mineral, the amount of mineral (mass per unit volume) remaining after defined simulation period was subtracted from the respective amount of each mineral initially present. This difference was then converted to mol m$^{-2}$ by multiplying with the respective compartment thickness ($t$) and dividing by the relative
formula mass (RFM). The resulting value was again divided by the simulation period to give dissolution rates in mol m\(^{-2}\) s\(^{-1}\) (Eq. 11).

\[
k_{\text{diss}} = \frac{(m_{k\text{init}} - m_{k\text{final}}) \times 1000 \times t}{\text{RFM}_k \times \text{SP}}
\] (11)

where \(k_{\text{diss}}\) (mol m\(^{-2}\) s\(^{-1}\)) is the dissolution rate of silicate mineral, \(k\), \(m_{k\text{init}}\) and \(m_{k\text{final}}\) are the initial and the final mass (kg m\(^{-3}\)) of silicate mineral \(k\), RFM\(_k\) is the relative formula mass (g mol\(^{-1}\)) of mineral \(k\) and SP is the simulation period (s). The number 1000 is the conversion factor from kg to g of mineral \(k\).

### 2.6 Sensitivity analysis

Morris’ sensitivity method (Morris, 1991) was used to assess the sensitivity of average silicate mineral dissolution rates to texture and physical weathering. It is one of the simplest and most widely used sensitivity analysis method (Saltelli et al., 2005). It is computationally cheaper than other sensitivity methods and therefore suitable for especially long run time models such as SoilGen (Finke et al., 2015; Yu et al., 2013). The method basically aims at quantifying the response of model output due to differences in the levels of input parameter (the so called elementary effects). In this study the levels include different textures and whether physical weathering is allowed or not. The output of interest in this case is the amount of mineral (Kg m\(^{-3}\)) lost over the simulation period due to chemical weathering which is itself influenced by differences in texture and physical weathering. The elementary effects of differences in texture \((u_i)\) on the amounts of mineral lost were calculated following Eq. (12) (Morris, 1991). Sensitivity of each silicate mineral was then evaluated by plotting the mean and the standard deviations of the elementary effects against each other (in the \(x\) and \(y\) axis, respectively) for both PhyWE and NoPhyWE scenarios and for each parent material.

\[
u_i = \frac{Y(x_1, x_2, x_3, \ldots x_i + \Delta x_i) - Y(x_1, x_2, x_3, \ldots x_i)}{\Delta x_i}
\] (12)
where $x_1, x_2, x_3, \ldots x_i$ are the different levels of input parameter (i.e., different textures, in this study), $\Delta x_i$ is the variation imposed on the input parameter measured as the Euclidean distance between two points in the textural triangle and $Y$ is the model output in response to each level of input parameter.

3 Results and discussion

3.1 pH evolution as a function of parent material

The evolution of pH as a function of parent material is shown in Fig. 3a (Model A) and Fig. 3b for Model B. There is erratic behaviour of pH in the beginning of the simulations (between 15,000 and 12,000 years BP) especially under granite. Generally, pH is increasing with depth and decreasing over time across the different parent materials as well as the two different model set ups (i.e., Model A and Model B). pH is generally higher in basalt and peridotite parent materials than granite but only in the first 5000 years of simulation (i.e., up to 10,000 years BP). The trends are however reversed in the subsequent years especially in Model A. There is generally a more gradual evolution of pH under Model B compared to Model A with a generally lower pH under Model A than Model B, when comparing respective parent materials.

The erratic behaviour of pH in the beginning of the simulations (between 15,000 and 12,000 years BP) especially under granite parent material (Fig. 3a, b) could be linked to the sensitivity of dissolution rates to dilution due to variation in precipitation. This period coincides with the drier periods (Fig. 2) with incidences of precipitation deficit in some years. Precipitation deficit means low dilution as well as limited mineral dissolution and release of cations, consequently keeping the pH low. At the current model version, the assumption is that dissolution occurs at far from equilibrium and thus the effect of the formation of secondary mineral on pH is not yet accounted for. This is certainly a limitation of this study and work is on-going to incorporate this mechanism into the model. A number of studies (Casey et al., 1993; Goddéris et al., 2006; Maher et
al., 2009; Moore et al., 2012; Zhu, 2005; Zhu et al., 2010) have already demonstrated that solute composition and secondary mineral precipitation controls the reaction affinity of primary minerals. The dissolution rates from this study are therefore expected to be faster than they would if secondary mineral precipitation were to be taken into account. The plunge in pH after 10 000 years BP for basalt and peridotite (Fig. 3a and b) could be linked to the depletion of forsterite at that time and thus less release of Mg\(^{2+}\). Comparing Fig. 3a and b, the effect of mineralogical composition on pH appears to become less in Fig. 3b (particularly after year 10 000 BP; between 500–1400 mm) than in Fig. 3a. This trend is likely due to the cation exchange capacity (CEC) buffering effect on pH in the zone of clay accumulation (Finke, 2012).

### 3.2 Evolution of clay mass fraction

Figure 4a shows the depth distribution of clay mass fraction taken at the final year of simulation (present situation). There is a clear difference between Model A and Model B, with a clear effect of physical weathering (PhyWE; dashed lines) on the amount of clay mass fraction in Model A (particularly in the top 0.3 m depth) where up to 8% of clay mass is produced due to physical weathering (Fig. 4; texture number 3). The effects of other processes notably clay migration on clay mass fraction is clearly visible in Model B with likely formation of an illuvial horizon (Bt horizon). Physical weathering and clay migration processes in the SoilGen model have been calibrated and their effect on the formation of eluvial (E) and Bt horizons was demonstrated in a modeling approach by Finke (2012). Although the clay contents were generally underestimated, Finke (2012) was able to reproduce the measurements of E and Bt horizon thicknesses by van Ranst (1981) in all the three loess profiles in the Zonian forest (Fig. 4b). Since the chemical and mineralogical analyses presented by van Ranst (1981) did not support any clay new formation, the increase in clay content with time was mainly attributed to physical weathering process. The right part of Fig. 4a shows the clear effect of clay migration process on clay mass fraction. The clay mass that is produced by physical weathering (Fig. 4a, left panel), is subsequently transported from the top compart-
ments into the lower compartments (through clay migration), forming E and Bt hori-
zens (Fig. 4a, right panel), respectively (Finke, 2012). The complete Bt belly could not
be shown by our results probably because of our shallow profile which was considered
to reduce the run-time of the model.

3.3 Mineral dissolution rates

3.3.1 Effect of parent material composition on dissolution rates

Figure 5 shows the effect of parent material composition on the average dissolution
rates of K-feldspar, albite, quartz and forsterite over successive time intervals of 500
years. With exception of quartz whose rates were increasing with time, the dissolution
rates across the 3 parent materials decrease with time. The dissolution rates of albite
and K-feldspar are higher (especially at the beginning of the simulation) under the
granite parent material than in basalt and peridotite. Model A dissolution rates across
all the minerals are generally higher than the rates from Model B. In granite however,
dissolution rates of albite and K-feldspar in Model A between 15 000 and 13 000 years
BP are lower than the respective dissolution rates in Model B. From 13 000 years BP
until 9000 years the dissolution rates are similar between the two models. In Basalt and
peridotite, the dissolution rates of albite and K-feldspar between 13 000 and 9000 years
BP are higher in Model B than in Model A. From 9000 until 0 years BP, the dissolution
rates of all minerals (except for quartz) across 3 parent materials are generally higher
in Model A than in Model B.

The properties of the parent material very much influence the chemical weathering
rates (Hartmann and Moosdorf, 2011; Navarre-Sitchler and Brantley, 2007; Oliva et al.,
2003). Results from this study indicate that the composition of the parent material in-
fluences directly the pH of the soil solution in two different ways (i) by the type of cation
it releases into the solution (i.e, monovalent, divalent, trivalent) and (ii) by the amount
of cations released which is directly related to the amount of mineral that is reacting.
Therefore all the trends pointed above and shown in Fig. 5 can be explained by the
influence that the parent material has on pH (e.g., interpreting Fig. 3a for Model A and 3b for Model B). The higher dissolution rates (especially in the beginning) of albite and K-feldspar observed in granite compared to basalt and peridotite could therefore be due to lower pH observed in granite than in Basalt and Peridotite at that point in time. The Mg$^{2+}$ released from forsterite (which is absent in granite) keeps the pH in the soil solution higher in basalt and peridotite than in granite and thus the lower dissolution rates of albite and K-feldspar in basalt and peridotite. The differences in Models A and B across the parent materials also follow the pH trends. For example in granite, the average pH (at 0.5 m depth) in Model B is generally lower than the pH in Model A between 15 000 and 13 000 years BP and therefore higher albite and K-feldspar dissolution rates and lower quartz dissolution rates in Model B. In basalt and peridotite, the average pH at this period (between 15 000 and 13 000 years BP) is more less the same and therefore the same dissolution rates of albite and K-feldspar between for both Model A and Model B. However between 12 000 and 9000 years BP, the average pH in basalt and peridotite is lower in Model B than in Model A, thus explaining the observed rise in the dissolution rates of albite and K-feldspar in Model B that are not observed in Model A. From 9000 years BP until 0 years BP, Model A dissolution rates of albite, K-feldspar and forsterite are higher than respective rates in Model B owing to the lower pH in Model A (averaged over 0.5 m) than in Model B (Fig. 3a and b). Quartz is less sensitive to pH less than 6 (Knauss and Wolery, 1988) and thus it's dissolution rates in Model A and Model B were not any different and did not seem to change from 10 000 until 0 years BP.

### 3.3.2 Effect of initial texture

The effect of initial texture on silicate mineral dissolution rates for Model A and Model B is presented in Fig. 6. As would be expected and consistent with previous studies (e.g. Hartmann et al., 2014; Phelan et al., 2014), the mineral dissolution rates are higher for finer textures than for coarse textures because of higher mineral surface area of clay and silt sized particles compared to the sand sized particles. In Model A,
 albite and K-feldspar dissolution rates across all initial textures, generally decrease with depth while dissolution rates of quartz generally increase with depth. In Model B, albite and K-feldspar dissolution rates across all initial textures, are generally constant with depth (except for texture number 4) while dissolution rates of quartz generally follow the same trend as in Model A and increase with depth. These dissolution rate-depth trends are related to pH which is generally increasing with depth. High pH favours quartz dissolution rates and slows down albite and K-feldspar dissolution rates.

3.3.3 Effect of physical weathering

The effect of only physical weathering (Model A) and the integrated effect of all soil forming processes (Model B) on the average silicate dissolution rates are shown in Fig. 7a and b, respectively. The rates are presented as a ratio of physical weathering to no physical weathering (i.e., PhyWE/NoPhyWE) where a value greater than 1 implies higher dissolution rate due to physical weathering. The results (both in Models A and B) indicate that the dissolution rates are generally higher in the top of the profile and decrease down the soil profile. Except for Forsterite, results in Model A indicate a positive effect of physical weathering on silicate dissolution rates (i.e., PhyWE/NoPhyWE > 1). Dissolution rates due to physical weathering are particularly higher in texture number 4 (solid black line) across all the minerals and parent materials with exception of Quartz mineral (in Model A) where dissolution rate due to physical weathering is highest under texture number 1. In Model B however, the effect of physical weathering is almost not visible (except for the texture number 4; solid line) as indicated with unity PhyWE/NoPhyWe ratios of all minerals across the different textures. Higher dissolution rates with no physical weathering compared to with physical weathering (i.e., PhyWE/NoPhyWE < 1) were only observed for Albite and K-feldspar below 0.75 m under Model B (see Fig. 5b: granite; texture number 4).

As already mentioned in the previous section, the direct effect of texture on chemical weathering is through it's influence on the mineral surface area. Based on Eq. (6), this would imply that the higher the number of particles moved from coarse to fine classes,
the higher the mineral surface area and thus the higher the mineral dissolution rate. This seems to be the case especially for coarse texture (texture number 4) where the dissolution rates of albite and K-feldspar in basalt and peridotite (Fig. 7a) are up to 1.4 times higher with PhyWE compared to NoPhyWE. The effect of physical weathering on the dissolution rates of albite and K-feldspar seems to be more pronounced in basalt and peridotite where pH is relatively higher and thus imposes less dominance on the chemical dissolution rates of albite and K-feldspar. The generally lower pH under granite could explain the higher effect of physical weathering on quartz dissolution rates under coarse textures 1, 4 and 6 (Fig. 7a).

The effect of texture on the dissolution rates could also be indirect through its relationship with hydrology. Our results imply that, although the physical weathering process produced more clay sized particles (Fig. 4, left panel) from already fine textures (i.e., 3 and 5; Table 1), the slowing down of water flow by this fine texture resulted into reduced leaching and higher pH, consequently cancelling out the added effect of physical weathering. Hydrology (water flow) and fluid residence time influence leaching and saturation levels of the soil solution (Moore et al., 2012; Velbel, 1993). According to Moore et al. (2012), hydrology is a key physical extrinsic factor and perhaps one of the most important factors that could explain observed differences between laboratory and field measured rates.

3.3.4 Interactive effects of selected soil forming processes on chemical weathering rates

The interacting soil forming processes that affect chemical weathering and are discussed this study include clay migration, plant uptake, carbon cycling and physical weathering. The results of these processes are presented under Model B (in Figs. 3b, 4 (right panel), 5, 6, 7b, 8 and 9). These processes have both direct and indirect effects on chemical weathering rates (White, 2002) through their influence on texture (e.g. clay migration, physical weathering, bioturbation) and on pH (e.g. clay migration, plant uptake, CO₂ production by mineralisation of organic matter). As discussed in
Finke (2012), clay migration process moves clay mass from the top part of the profile into the lower part of the profile (Fig. 4, right panel) leading to the formation an argillic (Bt) horizon which slows down water flow thus increasing solute concentration and lowering reaction affinity (Smeck and Ciolkosz, 1989; White and Brantley, 2003). Clay migration processes also has a pH buffering effect (Fig. 3b) through its influence on cation exchange capacity (Finke, 2012). Element cycling through plant uptake and release (through organic matter decomposition) influences the pH and consequently mineral weathering rates (Brady et al., 1999; Drever, 1994; Moulton et al., 2000; Stiillings et al., 1996). Higher pH that is visible in the top 0.25 m (Fig. 3b) can therefore be attributed to plant nutrient cycling process.

3.4 Sensitivity of mineral dissolution rates to physical weathering

Sensitivity of silicate mineral dissolution rates to texture and physical weathering are shown in Fig. 8. Results show low sensitivity (Model A) to no sensitivity of dissolution rates (Model B) due to differences in texture and physical weathering, across different minerals and parent materials. The pH of the soil solution seems to be a dominant factor to the chemical weathering of silicate minerals. In addition, the indirect effect of physical weathering on water flow and thus soil pH seem to oppose and cancel out the direct effects of physical weathering on the mineral surface area (as shown in Fig. 7b) and consequently the mineral dissolution rates.

3.5 Comparison between SoilGen modelled average mineral dissolution rates with Laboratory and field measured rates

SoilGen modelled silicate dissolution rates (Models A and B) were compared with rates obtained from field and laboratory experiments (Fig. 9). Rates plotted are for the whole profile depth (1.5 m) and for all the 6 different texture positions shown in Table 1. These rates are averaged for 15 000 years BP and they generally fall between what is reported from field measurements and what is reported from the laboratory studies. Our results...
are however generally closer to the laboratory rates than the field measured rates most likely because we assumed far from equilibrium reactions.

There seems to be no difference between dissolution rates from Model A and Model B across different parent materials when looking at the average rates over the whole simulation period of 15,000 years (Fig. 9). However, when the rates are calculated over short time intervals, e.g. 500 years, there is a clear difference at some points in time between the two models and even across different parent materials (see Fig. 5 for example). The evolution of silicate dissolution rates with time (Fig. 5) is not linear and this is in line with other previous studies (e.g. Hodson and Langan, 1999; White and Brantley, 2003). Generally silicate dissolution rates decrease with time due to depletion of reactive surfaces and, the formation of leached layers and secondary minerals (Hodson and Langan, 1999; White and Brantley, 2003). The comparison of dissolution rates obtained at different time scales therefore remains a challenge and could partly explain the significant differences in silicate dissolution rates reported in literature (White and Brantley, 2003). In addition, the various definitions of chemical weathering rates used in different studies e.g. cation chemical weathering rates (CCWR), chemical silicate rock weathering rates (CSRWR) and total chemical weathering rates (TCWR) make it difficult to compare results between studies (Hartmann and Moosdorf, 2011). Interpreting and comparing results from different studies should therefore be done with utmost care.

In our comparisons (Fig. 9) we chose to use only field and laboratory dissolution rates normalized to BET surface areas because the texture equation (Eq. 6) used to model mineral surface area was based on the calibration study with measured BET surface area (Sverdrup and Warfvinge, 1995). The field and laboratory rates were however not tied to the time scales or parent materials which could also limit our comparisons to some extent. There are also other questions that could be raised and that could potentially limit our comparisons with field and laboratory measurements. For example if the calibrations already done for clay migration, physical weathering and carbon cycling processes in the quartz-dominated loess sediment (Zonian forest; Finke, 2012) hold
for other sites with mafic and ultramafic parent materials? and whether ignoring the differences in climate and the time scales would invalidate the comparison between the dissolution rates from this study and previous studies? To answer such questions requires a more detailed study that is specific to field sites with field data of soil age, silicate dissolution rates, climate, hydrology, mineralogy and any other important factors to enable the calibration and validation of the model. Although still challenging, such studies are already feasible on well-studied chronosequences (e.g. Moore et al., 2012). However the intention of these comparisons (Fig. 9) is to show the general trends of our simulations rather than the absolute values.

4 Conclusions and outlook

We have used a fully mechanistic soil evolution model (SoilGen) to explore the sensitivity of silicate dissolution rates to the interaction between intrinsic (mineral composition, mineral surface area) and extrinsic factors (climate, physical weathering, clay migration, plant uptake, hydrology). Results from this study have shown consistency with both theoretical understanding of the effects of these factors on chemical weathering, and with observations from experiments and some modelling studies. Our results have demonstrated that although soil solution chemistry (pH) plays a dominant role in determining the silicate dissolution rates, all processes that directly or indirectly influence the soil solution composition play a major role in driving silicate dissolution rates. For example, although the sensitivity results did not confirm sensitivity of dissolution rates to physical weathering, the effect of texture (as influenced by physical weathering) on hydrology could have a substantial effect on the water flow, element leaching and consequently the pH and silicate dissolution rates.

Our dissolution rates results were in between field and laboratory rates, however they were rather high and closer to the laboratory rates owing to the assumption of far from equilibrium reaction. This remains a limitation of this study since near-to-equilibrium conditions have mainly been reported from the field experiments. However these find-
ings are important and challenge us to include secondary mineral precipitation mechanism in the model and perform comparative study to quantify these effects. Furthermore, calibration and validation of the model to the sites with detailed chronosequence data (soil age, silicate dissolution rates, climate, hydrology, mineralogy) is needed.

Despite the limitations identified, this study is another important step to demonstrate the critical need to couple different soil forming processes with chemical weathering in order to explain differences between silicate dissolution rates measured in the laboratory and in the field. In summary, results showed an inverse relationship of silicate mineral dissolution rates with time, an obvious effect of texture and, an indirect but substantial effect of physical weathering on silicate dissolution rates. Additionally, results have shown that clay migration and plant nutrient recycling processes influence the pH and thus the silicate dissolution rates.

**Code availability**

The SoilGen model is freely available. The manual and the programs for previous versions can be downloaded at: http://users.ugent.be/~pfinke/index_bestanden/Page1167.htm. SoilGen2.25 version is not yet available on the website but can be obtained on request (by sending an email to peter.finke@ugent.be).

**Author contributions.** Peter Finke developed the model code and designed the research. Emmanuel Opolot contributed to the model code development (weathering module), performed the simulations and prepared the manuscript with continuous and valuable contribution from Peter Finke.

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Evaluating sensitivity of silicate mineral dissolution rates

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References


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**Table 1.** Texture points randomly selected from the textural triangle and used as initial soil texture in all the model runs.

<table>
<thead>
<tr>
<th>Texture Number</th>
<th>Sand (%)</th>
<th>Clay (%)</th>
<th>Silt (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>63.3</td>
<td>12.0</td>
<td>24.7</td>
</tr>
<tr>
<td>2</td>
<td>41.6</td>
<td>18.7</td>
<td>39.8</td>
</tr>
<tr>
<td>3</td>
<td>5.5</td>
<td>27.4</td>
<td>67.1</td>
</tr>
<tr>
<td>4</td>
<td>86.8</td>
<td>6.1</td>
<td>7.0</td>
</tr>
<tr>
<td>5</td>
<td>8.7</td>
<td>10.7</td>
<td>80.6</td>
</tr>
<tr>
<td>6</td>
<td>51</td>
<td>4.1</td>
<td>44.9</td>
</tr>
</tbody>
</table>
**Table 2.** Primary minerals and their relative weight composition.

<table>
<thead>
<tr>
<th>Parent material type</th>
<th>Primary Silicate Mineral (wt %)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Albite</td>
</tr>
<tr>
<td>Granite</td>
<td>42.3</td>
</tr>
<tr>
<td>Basalt</td>
<td>32.1</td>
</tr>
<tr>
<td>Peridotite</td>
<td>10.9</td>
</tr>
</tbody>
</table>
Figure 1. Research set up.
Figure 2. Boundary conditions (Temperature, evapotranspiration, vegetation, bioturbation) used in the model. G = Grassland, C = Coniferous forest, D = Deciduous forest (Source: Finke, 2012).
Figure 3. (a) Time – depth evolution of pH with physical weathering (Model A) for 3 parent materials with texture number 5 (Table 1).
Figure 3. (b) Time–depth evolution of pH with interactive soil forming processes (Model B) for 3 parent materials with texture number 5 (Table 1).
Figure 4. (a) Clay mass fraction (%) evolution as a function of physical weathering (Model A, left) and as a function of interactive soil forming processes (Model B, right). Roman numerals 1–6 represent texture numbers presented in Table 1. Lines represent initial textures while broken lines represent evolution of texture as affected by other soil processes i.e., only physical weathering (left hand side) and a combination of mainly physical weathering and clay migration (right hand side). Notice that with only physical weathering allowed (Model A), the initial textures (lines) do not change whereas in Model B, even the initial textures change due to other processes notably clay migration. (b) Simulated clay (solid black line) and measured clay (gray line with triangles) showing increase in clay content in the sub layer (formation of Bt) due to clay migration and physical weathering processes (Source: Finke, 2012).
Figure 5. Example (based on soil texture number 5; Table 1) of time evolution of silicate dissolution rates in different parent materials (Granite, Basalt and Peridotite). The modelled rates are calculated for a depth of 0.5 m for every 500 years. Model A: Circles; solid (no physical weathering) open (physical weathering allowed). Model B: Triangles; solid (no physical weathering) open (physical weathering allowed).
Figure 6. Effect of initial texture (shown in Table 1) on the depth distribution of silicate dissolution rates. The rates shown are taken from granite parent material and are averaged over 15,000 years simulation period.
**Figure 7a.** Effect physical weathering on silicate mineral dissolution rates (Model A). Dissolution rates are presented here as a ratio of physical weathering to no physical weathering (i.e., PhyWE/NoPhyWE). Values greater than one imply that the rates are higher when physical weathering is allowed. Roman numerals 1–6 represent texture numbers presented in Table 1.
Figure 7b. Integrated effect of soil forming processes on silicate mineral dissolution rates (Model B). Dissolution rates are presented here as a ratio of physical weathering to no physical weathering (i.e., PhyWE/NoPhyWE). Values greater than one imply that the rates are higher when physical weathering is allowed. Roman numerals 1–6 represent texture numbers presented in Table 1.
Increasing variability in dissolution rates due to different elementary effects

Increasing mean differences in mineral dissolution rates due to elementary effects

Model A

Model B

Granite

Basalt

Peridotite

AlbiteNoPhyWE

AlbitePhyWE

KfeldNoPhyWE

KfeldPhyWE

QtzNoPhyWE

QtzPhyWE

AlbiteNoPhyWE

AlbitePhyWE

KfeldNoPhyWE

KfeldPhyWE

ForsteriteNoPhyWE

ForsteritePhyWE

Increasing mean differences in mineral dissolution rates due to elementary effects
Figure 8. Sensitivity of mineral dissolution rates to physical weathering (Model A) and to other interactive soil forming processes (Model B). The data used in this sensitivity analysis was extracted from the top soil compartment (0.05 m depth). \( \mu \) and \( \delta \) are the mean and standard deviation, respectively of elementary effects \( (u_i) \) calculated from Eq. (12).
Figure 9. Comparison of SoilGen average mineral dissolution rates (Models A and B) to laboratory and field determined dissolution rates. Field rates were taken from Parry et al. (2015) and White (2009) (Table 2) while lab rates were taken from: Holdren and Speyer (1987), Siegal and Pfannkuch (1984), Swoboda-Colberg and Drever (1993), Blum and Stillings (1995), Lee et al. (1998) (K-feldspar), Stillings et al. (1996), Welch and Ullman (1996), Oxburgh et al. (1994), Blum and Stillings (1995), Chou and Wollast (1985), Knauss and Wolrey (1986), Hamilton et al. (2000) (Albite), Brady and Walther (1990), and Dove (1994) (Quartz). The same field and laboratory rates are repeated for different parent materials (Granite, Basalt and Peridotite). Laboratory and field dissolution rates for Forsterite are not shown.