# **1** The importance of mineral determinations to PROFILE base

## 2 cation weathering release rates: A case study

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## 17 Abstract

18 Accurate estimates of base cation weathering rates in forest soils are crucial for policy decisions on sustainable 19 biomass harvest levels and for calculations of critical loads of acidity. The PROFILE model is one of the most 20 frequently used methods to quantify weathering rates, where the quantitative mineralogical input has often been 21 calculated by the A2M ("Analysis to Mineralogy") program based solely on geochemical data. The aim of this 22 study was to investigate how uncertainties in quantitative mineralogy, originating from modeled mineral 23 abundance and assumed stoichiometry, influence PROFILE weathering estimate, by using measured quantitative 24 mineralogy by X-ray powder diffraction (XRPD) as a reference. Weathering rates were determined for two sites, 25 one in Northern (Flakaliden) and one in Southern (Asa) Sweden. At each site, 3-4 soil profiles were analyzed at 26 10 cm depth intervals. Normative quantitative mineralogy was calculated from geochemical data and qualitative 27 mineral data with the A2M program using two sets of qualitative mineralogical data inputs to A2M: 1) A site-28 specific mineralogy based on information about mineral identification and mineral chemical composition as 29 determined directly by XRPD and electron microprobe analyses (EMPA), and 2) regional mineralogy, representing 30 the assumed minerals present and assumed mineral chemical compositions for large geographical areas in Sweden, 31 as per previous published studies. Arithmetic means of the weathering rates determined from A2M inputs ( $W_{A2M}$ ) 32 were generally in relatively close agreement with those (W<sub>XRPD</sub>) determined by inputs based on direct XRPD and 33 EMPA measurements. The hypothesis that using site-specific instead of regional mineralogy will improve the 34 confidence in mineral data input to PROFILE was supported for Flakaliden. However, at Asa, site-specific 35 mineralogies reduced the discrepancy for Na between WA2M and WXRPD but produced larger and significant 36 discrepancies for K, Ca and Mg. For Ca and Mg the differences between weathering rates based on different 37 mineralogies could be explained by differences in the content of some specific Ca- and Mg-bearing minerals, in 38 particular amphibole, apatite, pyroxene and illite. Improving the accuracy in the determination of these minerals 39 would reduce weathering uncertainties. High uncertainties in mineralogy, due for example to different A2M 40 assumptions, had surprisingly little effect on the predicted weathering of Na- and K-bearing minerals. This can be 41 explained by the fact that the weathering rate constants for the minerals involved, e.g. K-feldspar and micas, are similar in PROFILE. Improving the description of the dissolution rate kinetics of the plagioclase mineral group as 42 43 well as major K-bearing minerals (K-feldspars and micas) should be a priority to help improve future weathering 44 estimates with the PROFILE model.

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## 47 **Definitions and abbreviations**

- 48
- 49 Mineralogy = the identity (specific mineral or mineral group) and stoichiometry (specific mineral chemical
- 50 composition) of minerals that are present at a certain geographic unit, a particular site (*site-specific mineralogy*)
- 51 or a larger geographic province (*regional mineralogy*)
- Quantitative mineralogy= the quantitative information (wt.%) on the abundance of specific minerals inthe soil.

### 54 Abbreviations:

- 55 M<sub>XRPD</sub> = quantitative mineralogy based on XRPD (amount) and electron microprobe analysis (composition)
- 56  $M_{A2M-reg}$  = quantitative mineralogy calculated with the A2M model and using regional mineralogy input
- 57  $M_{A2M-site}$  = quantitative mineralogy calculated with the A2M model and using site-specific mineralogy input
- 58  $W_{XRPD}$  = weathering rate based on quantitative mineralogy determined by direct XRPD and electron microprobe 59 analysis
- W<sub>A2M</sub> = weathering rate based on quantitative mineralogy determined by the A2M model (unspecific mineralogy
   input)
- W<sub>A2M-reg</sub> = weathering rate based on quantitative mineralogy determined by the A2M model, and assuming regional
   mineralogy input.
- 64 W<sub>A2M-site</sub> = weathering rate based on quantitative mineralogy determined by the A2M model and assuming site 65 specific mineralogy input.
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#### 69 1. Introduction

70 The dissolution of minerals in soils and rocks during weathering represents, together with deposition, the most 71 important long-term supply of base cations for plant growth as well as acting as a buffer against soil and water 72 acidification. Quantifying weathering rates is therefore of key importance to guide modern forestry demands on 73 biomass removal by helping to identify threshold levels that are sustainable for base cation removal from soils and 74 waters. With the introduction of the harvest of forest biomass for energy production that includes whole tree 75 harvest and stump extraction, about 2-3 times more nutrients are exported from the forest compared to stem-only 76 harvest. As a result, issues of acidification and base cation supply are exacerbated and the sustainability of this 77 practice is questioned (Röser, 2008; de Jong et al. 2017). Regional nutrient balance calculations for Sweden have 78 indicated that net losses of base cations from forest soils can occur in stem-only harvest scenarios, and this trend 79 would be substantially exacerbated and more frequent in whole-tree harvesting scenarios, largely due to low 80 weathering rates (Sverdrup and Rosén, 1998; Akselsson et al., 2007a,b). Furthermore, the same effect occurs both

under current and projected future climate conditions (Akselsson et al., 2016).

82 The weathering rates included in these nutrient balance calculations are in most cases based on the PROFILE 83 model. This is a process-oriented model calculating steady-state weathering rates using transition state theory and 84 physical and geochemical properties of the soil such as temperature, soil moisture, soil mineralogy and concentrations of base cation, hydrogen and organic acids. (Sverdrup, 1996). This model has been widely applied 85 86 in Europe, Canada and the US during the last several decades or more of weathering research (Olsson et al., 1993; 87 Langan et al., 1995; Kolka et al., 1996; Starr et al., 1998; Sverdrup and Rosén, 1998; Whitfield et al., 2006; 88 Akselsson et al., 2007a; Koseva et al., 2010; Stendahl et al., 2013). In some cases nutrient balance calculations 89 have also been based on the depletion method (Olsson et al., 1993).

90 Reliable weathering rate estimates are crucial for the accuracy of future nutrient budget calculations (Futter et al., 91 2012). Regarding the accuracy of the PROFILE model, the importance of high accuracy in physical input 92 parameters for the modelled weathering rate outputs has been highlighted by Hodson et al. (1996) and Jönsson et 93 al. (1995). Among the various parameters Hodson et al. (1996) noted that the weathering response of the entire 94 soil profile depends critically on its mineralogy and as such any choice of the model user about mineralogical input 95 data may affect the model outcome significantly (Hodson et al., 1997). In most cases the mineralogical input to 96 the PROFILE model is also derived by modelling yet little attention has been given to the influence of modelled 97 versus directly measured mineralogical input data on calculated base cation release rates.

The most widely used method for direct quantitative mineralogical analysis of soil samples is X-ray powder 98 99 diffraction, and the accuracy that can be achieved has been demonstrated in round robin tests most notably the 100 Reynolds Cup (McCarty, 2002; Kleeberg, 2005; Omotoso et al., 2006, Raven and Self, 2017). Casetou-Gustafson 101 et al. (2018) made some independent assessment of the accuracy of their own XRPD data by geochemical cross 102 validation (i.e. the mineral budgeting approach of Andrist-Rangel et al., 2006). Nonetheless, we should stress that 103 like all analytical methods the determined weight fractions of minerals identified in a soil sample by XRPD will 104 have an associated uncertainty. Additionally, minerals present in minor amounts, nominally < 1% by weight, may 105 fall below the lower limit of detection of the XPRD method.

106 Due mainly to the relative ease of measurement and consequent ready availability of total element geochemical 107 data on soils, indirect methods of determining quantitative soil mineralogy, such as so called 'normative' geochemical calculations have been widely used to generate mineralogical data for use in the PROFILE model. 108 109 One such method is the normative "Analysis to Mineralogy" (A2M) program (Posch and Kurz, 2007) that has 110 commonly been used in PROFILE applications (Stendahl et al., 2013; Zanchi et al., 2014; Yu et al., 2016; 2018; 111 Kronnäs et al., 2019). Based on a quantitative geochemical analysis of a soil sample, typically expressed in weight percent oxides, as well as on some assessment of the available minerals in the soil sample (minerals present) and 112 113 their stoichiometry (chemical compositions), A2M calculates all possible mineralogical compositions for the soil 114 sample. Thus the A2M output for a given soil sample input has multiple solutions and can be described as a 115 multidimensional mineralogical solution space. This necessitates a choice when using A2M output in applications 116 such as weathering rate studies, the convention for which has been to use the geometric mean mineralogical 117 compositions e.g. Stendahl et al. 2013. Casetou-Gustafson et al. (2018) compared the output of A2M with directly 118 determined XRPD mineralogies at two sites, applying A2M in two different ways. In the first case the information 119 on available minerals in the model input was obtained from direct XRPD mineral identifications and information 120 on mineral stoichiometry from direct microprobe analysis of the minerals at the specific site (hereafter denoted 121 "site-specific"). In the second case the mineral stoichiometry and mineral identity were both assumed based on an 122 expert assessment of the probable mineralogy at the regional scale as given by Warfvinge and Sverdrup (1995), 123 hereafter denoted "regional". Casetou-Gustafson et al. (2018) concluded that using A2M in combination with 124 regional input data yielded results with large deviations from directly (XRPD) measured quantitative mineralogy, 125 particularly for two of the major minerals, K-feldspar and dioctahedral mica. When site-specific mineralogical 126 input data was used, measured and modeled quantitative mineralogy showed a better correspondence for most 127 minerals. For a specific mineral and a specific site, however, the bias in determination of quantitative mineralogy 128 might be significant depending on the accuracy of input data to A2M, i.e. total geochemistry and/or mineral 129 stoichiometry (Casetou-Gustafson et al., 2018). Potential errors like these in mineralogical input data might be 130 assumed to affect the calculated weathering for different base cations significantly.

In the present study, we used the different mineralogical data from Casetou-Gustafson et al. (2018) to model weathering rates of soils with the PROFILE model. Rates calculated based on measured mineral abundances using quantitative XRPD in combination with measured mineral elemental compositions are taken as 'reference' weathering rates to which other rates are compared. Samples for this study were collected from podzolised till soils from 8 soil profiles at two forest sites in northern and southern Sweden, respectively.

136 The primary objective of this study was to describe and quantify the effect of differences in mineralogy input on 137 PROFILE weathering rates, leaving all other input parameters of the PROFILE model constant to isolate the effects 138 of variation in input of mineral stoichiometry and abundance. A first specific aim was to determine the 139 uncertainties in weathering rates caused by uncertainties in normative quantitative mineralogy. This was 140 approached by comparing PROFILE runs using modeled mineralogies based on the presence of minerals of a 141 specific site or a larger geographic region (i.e. site-specific and regional mineralogy) with PROFILE runs using 142 the directly measured mineralogy. The latter was assumed to represent the 'true' mineralogy at each site. The 143 comparison of PROFILE weathering rates, based on XRPD versus A2M mineralogy, was done using 1000 random 144 solutions per sample from the entire multidimensional A2M mineralogical solution space. In the following,

- 145 weathering rates calculated by PROFILE based on XRPD and A2M mineralogies are denoted W<sub>XRPD</sub> and W<sub>A2M</sub>, 146 respectively.
- 147 A second specific aim was to investigate how the over- or underestimation of WA2M in relation to WXRPD mirrors 148 the over- or underestimation of mineral contents estimated with A2M.

149 The following hypotheses were made:

150 (1) PROFILE weathering rates obtained with normative quantitative mineralogy calculated based on site-specific

151 mineralogical information about mineral identity and mineral stoichiometry, are more similar to the reference 152 weathering rates than PROFILE runs obtained with normative quantitative mineralogy calculated based on 153 regional information only.

- 154 (2) Over- and underestimations of weathering rates of different base cations by the PROFILE model can be 155 explained by over- or underestimations of mineral contents of a few specific minerals.
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#### 157 2. Materials and methods

#### 158 2.1 Study sites

- Two experimental forest sites, Asa in southern, and Flakaliden in northern Sweden, were used for the study (Table 159 160 1). Both sites have Norway spruce (Picea abies (L.) Karst) stands of uniform age, but differ in climate. Flakaliden 161 is located in the boreal zone with long cold winters, whereas Asa is located in the hemiboreal zone. The soils have 162 similar texture (Sandy loamy till), soil types (Spodosols) and moisture conditions. According to the geographical 163 distribution of mineralogy types in Sweden the sites belong to different regions (Warfvinge and Sverdrup, 1995). 164 The experiments, which started in 1986, aimed at investigating the effects of optimized water and nutrient supply 165 on tree growth and carbon cycling in Norway spruce forests (Linder 1995, Albaugh et al. 2009). The sites are 166
- incorporated in the Swedish Infrastructure for Ecosystem Science (SITES).

#### 167 2.2 Soil sampling and stoniness determination

168 Soil sampling was performed in October 2013 and March 2014 in the border zone of four plots each of the sites. 169 Plots selected for sampling were untreated control plots (K1 and K4 at Asa, 10B and 14B at Flakaliden) and 170 fertilized 'F' plots (F3, F4 at Asa, 15A, 11B at Flakaliden). A rotary drill was used in order to extract one intact 171 soil core per plot (17 cm inner diameter) expect for plot K4, F3 and F4 at the Asa site. A 1 x 1m soil pit was 172 excavated at each of the three latter plots due to inaccessible terrain for forest machinery. The maximum mineral 173 soil depth varied between 70-90 cm in Flakaliden and 90-100 cm in Asa.

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175 The volume of stones and boulders was determined with the penetration method by Viro (1952), and by applying

176 penetration data to the functions by Stendahl et al. (2009). A metal rod was penetrated at 16 points per plot into

- 177 the soil until the underground was not possible to penetrate any further, or to the depth 30 cm. There was a higher
- 178 average stoniness at Flakaliden than Asa (39 vol-% compared to 29 vol-% in Asa) that could partially explain the
- 179 lower maximum sampling depth at Flakaliden.

#### 180 2.3 Sample preparation

- 181 Soils samples for chemical analyses were taken at 10 cm depth intervals in the mineral soil. Prior to analysis all
- soil samples were dried at 30–40 °C and sieved at 2 mm mesh. Soil chemical analyses were performed on the fine
- 183 earth fraction (< 2mm).

#### 184 2.4 Analysis of geochemistry, total carbon and soil texture

185 Total carbon was determined using a LECO elemental analyzer according to ISO 10694. Analysis of total 186 geochemical composition, conducted by ALS Scandinavia AB, was made by inductively coupled plasma 187 spectrometry (ICP-MS). Prior to analyses, the samples were ignited at 1000° C to oxidize organic matter and 188 grinded with an agate mortar. Particle size distribution was analyzed by wet sieving and sedimentation (Pipette 189 method) in accordance with ISO 11277. More details about the analytical procedure was given by Casetou-190 Gustafson et al. (2018).

#### 191 2.5 Determination of quantitative mineralogy

A detailed description of methods used to quantify mineralogy of the samples was given by Casetou et al. (2018)and these are described in brief below.

#### 194 2.5.1 Measured mineralogy

Quantitative soil mineralogy was determined with the X-ray powder diffraction technique, XRPD ( $M_{XRPD}$ ) (Hillier 1969, 2003) (Table S1a-b). Preparation of samples for determination of XRPD patterns was made from spray drying slurries of micronized soil samples (<2 mm) in ethanol. Quantitative mineralogical analysis of the diffraction data was performed using a full pattern fitting approach (Omotoso et al., 2006). In the fitting process, the measured diffraction pattern is modelled as a weighted sum of previously recorded and carefully verified standard reference patterns of the prior identified mineral components. The chemical composition of the various minerals present in the soils was determined by electron microprobe analysis (EMPA) (Table S6).

#### 202 2.5.2 Calculated mineralogy

203 The A2M program (Posch and Kurz, 2007) was used to calculate quantitative mineralogical composition (MA2M) 204 from geochemical data. Based on a set of pre-determined data on mineral identity and stoichiometry, the model 205 outcome is a range of equally possible mineralogical compositions. The multidimensional structure of this 206 normative mineralogy model is a consequence of the number of minerals being larger than the number of analysed 207 elements, where a specific element can often be contained in several different minerals. A system of linear 208 equations is used to construct an M-N dimensional solution space (Dimension M= Number of minerals, Dimension 209 N=number of oxides). In this study we used one thousand solutions to cover the range of possible quantitative 210 mineralogies that may occur at a specific site.

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A2M was used to calculate 1000 quantitative mineralogies each for two different sets of mineral identity and

214 identity and stoichiometry for the four major mineralogical provinces in Sweden as suggested by Warfvinge and

215 Sverdrup (1995), of which Asa and Flakaliden belong to different regions (Table S5). Site-specific mineralogy

- 216 refers to the measured mineral identity and stoichiometry determined by the XRPD and electron microprobe
- analyses of the two sites (Table S6) (Casetou-Gustafson et al., 2018).

#### 218 2.6. Estimation of weathering rates with PROFILE

#### 219 2.6.1 PROFILE model description

220 The biogeochemical PROFILE model can be used to study the steady-state weathering (i.e. stoichiometric mineral 221 dissolution) of soil profiles, as weathering is known to be primarily determined by the physical soil properties at 222 the interface of wetted mineral surfaces and the soil solution. PROFILE is a multilayer model, thus, for each soil 223 layer, parameters are specified based on field measurements and estimation methods (Warfvinge and Sverdrup, 224 1995). Furthermore, isotropic, well mixed soil solution conditions are assumed to prevail in each layer as well as 225 surface limited dissolution in line with early views by Aagard and Helgeson (1982) and Cou and Wollast (1985) 226 (Sverdrup, 1996). Based on these major assumptions, PROFILE calculates chemical weathering rates from a series 227 of kinetic reactions that are described by laboratory determined dissolution rate coefficients and soil solution 228 equilibria (i.e. transition state theory) (Sverdrup and Warfvinge, 1993). The PROFILE version (September 2018) 229 that was used in this study is coded to produce information on the weathering contribution of specific minerals, 230 which allowed us to test our second hypothesis. This version is based on the weathering rates of 15 minerals. Of 231 these, apatite, pyroxene, illite, dolomite and calcite were not found at the two study sites according to XRPD data 232 (Table S1).

#### 233 2.6.2 PROFILE parameter estimation

The only parameter that was changed between different PROFILE runs was the quantitative mineralogy for each soil layer, as described above. Hence, PROFILE estimated weathering rates (W) based on measured mineralogy ( $W_{XRPD}$ ), and the two versions of A2M calculated mineralogy, regional ( $W_{A2M-reg}$ ), and site-specific ( $w_{A2M-site}$ ). In the regional mineralogy, plagioclase is assumed to occur as pure anorthite and pure albite for simplification as has been used in previous studies (Stendahl et al., 2013; Zanchi et al., 2014). This simplification was done in order to avoid having a number of minerals containing different amounts of Ca and Na, as a result of plagioclase forming a continuous solid solution series, since it would not affect the weathering rates.

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The physical soil layer specific parameters, that were kept constant between different profile runs, were exposed mineral surface area, stoniness, soil bulk density and soil moisture (Table 2). Exposed mineral surface area was estimated from soil bulk density and texture analyses in combination with an algorithm specified in Warfvinge and Sverdrup (1995) and critically discussed in Hodson et al. (1998). The volumetric field soil water content in Flakaliden and Asa was estimated to be 0.25 m<sup>3</sup> m<sup>-3</sup> according to the moisture classification scheme described in Warfvinge and Sverdrup (1995). It was used to describe the volumetric water content for each soil pit.

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Another group of parameters kept constant was chemical soil layer specific parameters. Aluminum solubility coefficient needed for solution equilibrium reactions, defined as log{Al<sup>3+</sup>}+3pH, was estimated applying a function developed from previously published data (Simonsson and Berggren, 1998) on our own total carbon and oxalate extractable aluminum measurements. The function is based on the finding that the Al solubility in the upper B-horizon of Podzols is closely related to the molar ratio of aluminum to carbon in pyrophosphate extracts,

- and that below the threshold value of 0.1, Al solubility increases with the Al<sub>p</sub>/C<sub>p</sub> ratio (Simonsson and Berggren,
- 255 1998). Thus, a function was developed for application to our own measurements of  $Al_{ox}$  and  $C_{tot}$  based on the
- assumption that it is possible to use the  $Al_{ox}/C_{tot}$  ratio instead of the  $Al_p/C_p$  ratio. Data on soil solution DOC were
- available from lysimeters installed at 50 cm depth for plot K4 and K1 in Asa and 10B and 14B in Flakaliden, and
- 258 these values were also applied to soil depths below 50 cm (H. Grip, unpublished data). The E-horizon (0 10 cm)
- at Flakaliden) and A-horizon (0 –10 cm at Asa) were characterized by higher DOC values based on previous
- 260 findings (Fröberg et al., 2013) and the classification scheme of DOC in Warfvinge and Sverdrup (1995). Partial
- 261 CO<sub>2</sub> pressure values in the soil were taken from the default estimate of Warfvinge and Sverdrup (1995).
- 262
- 263 Other site-specific parameters that were kept constant between PROFILE runs were evapotranspiration, 264 temperature, atmospheric deposition, precipitation, runoff and nutrient uptake. Temperature is one of the important 265 factors that regulate the weathering rate, and for steady-state calculations in PROFILE the mean annual 266 temperature is used. Kronnäs et al. (2019) demonstrates how weathering rates varies between seasons, due to e.g. 267 variations in temperature. Precipitation is used in PROFILE to calculate vertical water flow through the soil profile. 268 The main effect of precipitation on weathering rates is its impact on soil moisture, but in PROFILE soil moisture 269 is not internally modeled, but given as input. An estimate of the average evaporation per site was derived from 270 annual averages of precipitation and runoff data using a general water balance equation. Deposition data from two 271 sites of the Swedish ICP Integrated Monitoring catchments, Aneboda (for Asa) and Gammtratten (for Flakaliden) 272 (Löfgren et al., 2011) were used to calculate the total deposition. The canopy budget method of Staelens et al. (2008) was applied as in Zetterberg et al. (2014) for Ca<sup>2+</sup>, Mg<sup>2+</sup>, K<sup>+</sup>, Na<sup>+</sup>. The canopy budget model is commonly 273 used for elements that are prone to canopy leaching (Ca<sup>2+</sup>, Mg<sup>2+</sup>, K<sup>+</sup>, Na<sup>+</sup>, SO<sub>4</sub><sup>2-</sup>) or canopy uptake (NH<sub>4</sub><sup>+</sup>, NO<sub>3</sub><sup>-</sup>) 274 275 and calculates the total deposition (TD) as the sum of dry deposition (DD) and wet deposition (WD). Wet 276 deposition was estimated based on the contribution of dry deposition to bulk deposition, both for base cations and 277 anions, using dry deposition factors from Karlsson et al. (2012, 2013). Base cation and nitrogen accumulation rate 278 in above-ground tree biomass (i.e. bark, stemwood, living and dead branches, needles) was estimated as the 279 average accumulation rate over a 100 years rotation length in Flakaliden compared to a 73 years rotation length in 280 Asa. These calculations were based on Heureka simulations using the StandWise application (Wikström et al., 281 2011) for biomass estimates in combination with measured nutrient concentrations in above- ground biomass (S. 282 Linder unpubl. data).

#### 283 2.7 A definition of significant discrepancies between WA2M and WXRPD

A consequence of the mathematical structure of the A2M program is that the final solution space of possible quantitative mineralogies produces an uncertainty range of weathering estimates, but in a different sense than the uncertainty caused by e.g. uncertainties in chemical analyses, because all mineralogies produced within this range are equally likely. Thus, here we define a significant discrepancy between  $W_{XRPD}$  and  $W_{A2M}$  to occur when the former is outside the range of the latter, as illustrated in Fig. 1a. The opposite case is a non-significant discrepancy,

- when the weathering rates based on XPRD are contained in the weathering range based on A2M (Figure 1b).
- 290

291 The uncertainty range of W<sub>A2M</sub> can potentially be reduced by reducing uncertainties in analyses of soil 292 geochemistry but most particularly by definitions of available minerals and their stoichiometry. Furthermore, some 293 discrepancies between  $W_{XRPD}$  and  $W_{A2M}$  might also arise due to limitations of the XRPD method, particularly 294 when minerals occur near or below the detection limit.

#### 295 2.8 Statistical analyses

296 In order to quantify the effect of mineralogy on PROFILE weathering rates two statistical measures were used to

297 describe the discrepancies between  $W_{XRPD}$  and  $W_{A2M}$ . Firstly, root mean square errors (RMSE) of the differences 298 between  $W_{XRPD}$  and the arithmetic mean of weathering rates based on regional and site-specific mineralogy, i.e.,

299  $W_{A2M-reg}$  and  $W_{A2M-site}$ , were calculated:

300

301 
$$RMSE = \sqrt{\frac{1}{n} \sum_{i=1}^{n} (WXRPD_i - WA2M_i)^2}$$
 Eq. (1)

302

303 RMSE's were calculated individually for each element, soil layer and soil profile for two data sets. An RMSE 304 expressing the error of the aggregated, total weathering rates in the 0–50 cm soil horizon was calculated to test our 305 first hypothesis (RMSE of total weathering). In addition, an RMSE expressing the errors originating from 306 discrepancies between  $W_{XRPD}$  and  $W_{A2M}$  for individual minerals was also calculated (RMSE of weathering by 307 mineral). In the latter case, sums of RMSE's by mineral were calculated for each element and soil profile by 308 analogy with the summing up of weathering rates for the whole 0–50 cm soil profile.

309

310 Secondly, relative discrepancies (i.e. average percentage of over- or underestimation of  $W_{A2M}$  compared to  $W_{XRPD}$ ) 311 were calculated as the absolute discrepancy divided by the measured value.

312

313 Relative error = 
$$\left(\frac{WA2M_i - WXRPD_i}{WXRPD_i}\right)$$
 100 Eq. (2)

314

Relative errors were calculated for each site by comparing the average sum of  $W_{A2M}$  in the upper mineral soil (0– 50 cm) with the sum of  $W_{XRPD}$  in the upper mineral soil.

Statistical plotting of results was performed using R (version 3.3.0) (R Core Team, 2016) and Excel 2016.

#### 318 **3. Results**

#### 319 3.1 Weathering rates based on XRPD mineralogy

soil layers (50-100 cm) is given in Table S2.

Weathering estimates with PROFILE are hereafter presented as the sum of weathering rates in the 0–50 cm soil horizon, since this soil depth is commonly used in weathering rate studies. Information on individual, and deeper

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Weathering rates of the base cations based on quantitative XRPD mineralogy (W<sub>XRPD</sub>), i.e. the reference weathering rates, were ranked in the same order at both sites, with Na>Ca>K>Mg (Table S2). On average, weathering rates of Na, Ca, K and Mg at Asa were 17.7, 8.4, 5.6 and 3.6 mmol<sub>c</sub> m<sup>-2</sup> yr<sup>-1</sup>, respectively.

- 327 Corresponding figures for Flakaliden were of similar magnitude, i.e., 14.8, 9.8, 5.7 and 5.6 mmol<sub>c</sub> m<sup>-2</sup> yr<sup>-1</sup>. The
- 328 variation in weathering rates between soil profiles was smaller at Asa than at Flakaliden, as the standard deviation

in relation to the means for different elements ranged between 0.2-2.3 at Asa, and 2.0 – 5.7 at Flakaliden (Table

330 S2).

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#### 332 3.2 Comparison between weathering rates based on XRPD and A2M mineralogy

333 At Flakaliden, WA2M-site was generally in closer agreement with WXRPD than WA2M-reg (Fig. 2b), in line with the first 334 hypothesis. The discrepancies between  $W_{XRPD}$  and  $W_{A2M}$  were small and non-significant for Mg regardless of the 335 mineralogy input used in A2M, although the estimated discrepancies were reduced when site-specific mineralogy 336 was used. The use of regional mineralogy in A2M underestimated K release rates compared to W<sub>XRPD</sub>, and the 337 discrepancy was significant. Using site-specific mineralogy resulted in smaller and non-significant discrepancy 338 for K release rates. A similar response to different mineralogies was revealed for Ca, although the result varied 339 more among soil profiles. In contrast to K and Ca, the release of Na was overestimated by both WA2M-site and WA2M-340 reg compared to W<sub>XRPD</sub>. The discrepancies were significant regardless of the mineralogy input used in A2M, 341 although using site-specific mineralogy slightly reduced the discrepancy. The generally closer agreement between 342 WA2M-site and WXRPD than WA2M-reg at Flakaliden was also indicated by the lower RMSEs of total weathering for all 343 base cations when site-specific mineralogy was used (Fig. 3a). Relative RMSE were below 20 % for W<sub>A2M-reg</sub>, but 344 below 10 % for WA2M-site. However, RMSE for Na was only slightly smaller for WA2M-site than WA2M-reg (16 % for 345 W A2M-site).

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347 PROFILE weathering rates for Asa revealed a different pattern compared to Flakaliden, and the results for Ca, Mg 348 and K were contradictory to hypothesis one. WA2M-reg was in close agreement with WXRPD for K, Ca and Mg, and 349 the small discrepancies were non-significant (Fig. 2a). However, WA2M-reg for Na was consistently overestimated 350 compared to W<sub>XRPD</sub> and the discrepancies were significant. Using site-specific mineralogy improved the fit 351 between W<sub>XRPD</sub> and W<sub>A2M</sub> for Na but had rather the opposite effect on the other base cations at this site. For K, Ca 352 and Mg, WA2M-site overestimated weathering rates, and resulted in significant discrepancies, and larger RMSE, 353 whereas the discrepancies for Na were reduced and non-significant (Fig. 3a). At Asa, the highest relative RMSEs of total weathering occurred for Ca and Mg with WA2M-site (> 30 %) (Fig. 3a). Large standard deviations were due 354 355 to a single soil profile, F4. The better consistency with WA2M-reg was indicated by RMSE below 10 % for Ca and Mg, and that RMSE for Mg was half of the error with WA2M-site. Only for Na, RMSE was lower for WA2M-site than 356 357 with WA2M-reg.

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A complementary illustration of the relationships between weathering rates based on XRPD and A2M is shown in Fig. 4 and provided as Tables S3 and S4, which includes all data from individual soil layers 0–50 cm. A general picture is that  $W_{A2M-site}$  was less dispersed along the 1:1-line than  $W_{A2M-reg}$ , in particular for Flakaliden. On the other hand, for weathering rates in the lower range (< 5 mmol<sub>c</sub> m<sup>-2</sup> yr<sup>-1</sup>) site-specific mineralogy tended to generate both over- and underestimated weathering rates. In most soil profiles, deviations from the 1:1-line were frequent

in soil layers below 20 cm. For Na under- and overestimations occurred in the whole range of weathering

365 estimates,

#### **366 3.3 Mineral-specific contribution to weathering rates**

367 In spite of its intermediate dissolution rate, plagioclase was, due to its abundance, the most important Na-bearing 368 mineral determined in this study (Table 3 and Fig. 5). Plagioclase is a variable group of minerals with different 369 stoichiometric proportions of Ca and Na, from the purely sodic albite on the one hand to the purely calcic anorthite 370 on the other hand (Table S5) as well as with intermediate compositions (Table S6). For simplicity, they will be 371 referred to in this study as sodic and calcic plagioclase. Based on the same quantitative mineralogy (i.e. same 372 elemental compositions and identity of minerals), W<sub>XRPD</sub> and W<sub>A2M-site</sub> gave strong weight to both calcic and sodic 373 plagioclase in estimating Na release rates, but WA2M-site gave stronger weight to calcic versus sodic plagioclase at 374 Asa, and vice-versa at Flakaliden (Fig. 5). In spite of these differences, the resultant release rates of Na according 375 to  $W_{A2M-site}$  and  $W_{XRPD}$  were rather similar (Fig. 5).

376

Total Na release rates of  $W_{A2M-reg}$  compared to  $W_{XRPD}$  were moderately overestimated. The relative RMSE of weathering by specific Na-containing minerals were of more similar magnitude for Na at Flakaliden compared to Asa (Fig. 3b). However, the standard deviations of RMSE were relatively large at Flakaliden, due to large RMSE for albite in one specific soil profile (11B) (Table S7). Contrary to relative RMSE of total weathering, the relative

RMSE of weathering by specific minerals was lower for Na at Asa with regional than site-specific mineralogy.

381 382

According to  $W_{XRPD}$ , calcic plagioclase weathering was the most important source to Ca release at Flakaliden, and the second most important source at Asa after epidote (Fig. 5). As for Na,  $W_{A2M-site}$  gave stronger weight to calcic plagioclase than  $W_{XRPD}$  at Asa. It was the other way around for  $W_{A2M-site}$  at Flakaliden and the regional mineralogy (i.e.  $W_{XRPD}$  gave stronger weight to calcic plagioclase than  $W_{A2M-site}$ ). Another important Ca source in weathering estimates based on A2M was apatite. This mineral was not detected in the XRPD analyses but was included in both A2M mineralogies as a necessary means to allocate measured total phosphorus content to a specific mineral (Casetou-Gustafson et al. 2018).

390

Similar to Na, relative RMSE of weathering by Ca-containing minerals were several magnitudes larger than RMSE of the total weathering of Ca. In other words, although an overall similar weathering rates might be generated by the PROFILE model based on different quantitative mineralogies, the underlying modelled contributions from different minerals can be markedly different. At Flakaliden, the mean relative RMSE by specific minerals were larger for regional than site-specific mineralogy at Flakaliden (Fig. 3b). However, the difference was not significant since the standard deviations were high, probably due to larger RMSE for Ca-bearing minerals in soil profile 11B (Table S7).

398

A general picture of the mineral contribution to Mg release is that  $W_{XRPD}$  placed most weight to amphibole whereas in  $W_{A2M}$ , Mg release was more equally distributed among other minerals, notably hydrobiotite, trioctahedral mica and vermiculite. At Asa, and to an even larger extent at Flakaliden, Mg release by A2M mineralogies was determined by a higher contribution of minerals with high dissolution rates (Fig. 5 and Table 3) (i.e. In  $W_{A2M-site}$ , hydrobiotite and trioctahedral mica; In  $W_{A2M-reg}$ , muscovite and vermiculite at Asa and biotite and illite at Flakaliden). At Asa, less weight was given to amphibole by  $W_{A2M-site}$  compared to  $W_{XRPD}$ . At Flakaliden, the  $W_{A2M}$ site was close  $W_{XRPD}$  in spite of the very different allocations of weathering rates to different minerals. The

- 406 underestimation of Mg release by  $W_{A2M-reg}$  was largely explained by the lower weight given to amphibole in both 407 A2M scenarios (Fig. 5). However, A2M gave larger weight to other minerals. The sums of RMSEs of weathering 408 from specific Mg-bearing minerals were much larger for regional than site-specific mineralogy at Flakaliden and 409 reached a maximum value of 156 %. A contributing factor were generally larger RMSE for the mineral 410 contribution of amphibole to Mg weathering and the fact that pyroxene contributed to the RMSEs of the total 411 weathering of Mg. Furthermore, a large standard deviation for the sum of RMSE of specific minerals (Fig. 3b)
- 412 was caused by soil profile 11B where more weight was placed on amphibole and biotite in contributing to Mg
- 413 weathering (Table S7). The two A2M mineralogies resulted in the same RMSEs for Mg-bearing minerals at Asa
- 414 (Fig. 3b).
- 415

416 Potassium release rates were largely dominated by K-Feldspar weathering in both W<sub>XRPD</sub> and W<sub>A2M-site</sub>. However, 417 K release by W<sub>A2M-reg</sub> (Fig. 5) were largely determined by micas at both sites. Together with Mg, these elements had also the lowest weathering rates, indicating that differences between WA2M-reg and WXRPD in relative terms were 418 419 not correlated with the magnitude of weathering. Unlike the other base cations, relative RMSE of K-bearing 420 minerals were lower at both sites when site-specific mineralogy was used instead of regional (Fig. 3b), and the 421 mineral specific RMSEs were also of similar magnitude as the RMSE of the total weathering (Fig.3a). WA2M-site of 422 K (Fig. 3b), were not several magnitudes larger than RMSE of the total weathering (Fig. 3a). The largest relative RMSEs of K-containing minerals were reached by WA2M-reg at Flakaliden in soil profile 11B, indicated by the high 423 424 standard deviation.

#### 425 4. Discussion

#### 426 4.1 General range of weathering rates in relation to expectations from other sensitivity studies, and the 427 range of discrepancies between W<sub>XRPD</sub> and WA2M

428 To our knowledge, the present study is the first to have examined the sensitivity of the PROFILE model on real 429 case study differences of directly measured mineralogy versus indirectly determined normative mineralogy. 430 However, a few systematic studies have been made previously to test the influence of mineralogy inputs, amongst 431 other input parameters, to PROFILE weathering rates. Jönsson et al. (1995) concluded that uncertainty in 432 quantitative mineralogy could account for a variation from the best weathering estimate of about 20 %, and that 433 variations in soil physical and chemical parameters could be more important. The sensitivity analysis of Jönsson 434 et al. (1995) was made by a Monte Carlo simulation where mineralogical inputs were varied by  $\pm 20$  % of abundant minerals, and up to  $\pm$  100 % of minor minerals. Shortly after, Hodson et al. (1996) examined the sensitivity of the 435 436 PROFILE model with respect to the sensitivity of weathering of specific minerals and concluded that large 437 uncertainties especially in soil mineralogy, moisture, bulk density, temperature and surface area determinations 438 will have a larger effect on weathering rates than was reported by Jönsson et al. (1995).

- 439 Compared with the sensitivity analyses by Jönsson et al. (1995), the range of uncertainty in dominating mineral
- 440 inputs used in the present study was of similar order of magnitude. For this study we used the XRPD measured
- 441 (M<sub>XRPD</sub>) and A2M estimated mineralogies (M<sub>A2M</sub>) determined by Casetou-Gustafson et al. (2018). For example,
- 442 they concluded that  $M_{A2M-reg}$  produced a low relative RMSE of total plagioclase (7 11 %) but higher relative
- 443 RMSE for less abundant minerals, such as dioctahedral mica (90 106 %). They also showed that when regional

- 444 mineral identity and assumed stoichiometry was replaced by site-specific mineralogy ( $M_{A2M-site}$ ), the bias in 445 quantitative mineralogy was reduced.
- 446 Thus, given this bias in quantitative mineralogy input to PROFILE, discrepancies of  $W_{A2M}$  from  $W_{XRPD}$  at our
- 447 study sites should have been on the order of 20 % or less, and site-specific mineralogy inputs should produce
- 448 weathering rates with lower discrepancies than regional mineralogy. The result of this study was in agreement
- 449 with this expectation for all elements at Flakaliden but only for Na at Asa. The different quantitative mineralogies
- 450 resulted in discrepancies between  $W_{A2M}$  and  $W_{XRPD}$  that differed with site (Fig. 3a, 5).

#### 451 **4.2** Is W<sub>A2M-site</sub> more consistent than W<sub>A2M-reg</sub>?

- 452 Our first hypothesis, that using site-specific mineralogy in the PROFILE model compared to regional mineralogy,
- 453 should result in weathering rates closer to XRPD-based mineralogy, and thus be more consistent, was generally
- 454 supported for Flakaliden, but only for Na at Asa. This result was revealed from both the occurrence of significant
- 455 discrepancies as well as the RMSE of the total weathering rates. Thus, the results did not support our first
- 456 hypothesis in a consistent way. The possible reasons for this outcome are discussed below, based on the analysis
- 457 of how different minerals contributed to the overall weathering rates.

# 4.3 How are discrepancies between W<sub>A2M</sub> and W<sub>XRPD</sub> correlated to bias in determinations of quantitative mineralogy

- 460 The version of the PROFILE model used in this study allowed a close examination of the per element weathering
- 461 rate contributions obtained from different minerals that provide some insight into the causes to the total  $W_{A2M}$ 462 discrepancies.

#### 463 4.3.1 Sodium release rates

464 A biased determination of mineralogy may not necessarily result in a corresponding bias of PROFILE weathering 465 estimates if the discrepancies are cancelling each other out, and if dissolution rates of the different minerals are rather similar. This was probably the case for Na. At both study sites and for both W<sub>XRPD</sub> and W<sub>A2M</sub>, Na release 466 rates were largest for plagioclase minerals. The Na release from WA2M-site and WA2M-reg were close to WXRPD at both 467 468 study sites (i.e. all weathering rates were in the range of 17-19 mmol<sub>c</sub> m<sup>-2</sup> yr<sup>-1</sup>) nonetheless W<sub>A2M-site</sub> placed more 469 weight to calcic plagioclase and WA2M-reg more weight to albitic plagioclase (Fig.5). Contrary to our second 470 hypothesis, the relatively high precision in total release rates (i.e.<10%; Fig. 3a) of Na was not correlated to the 471 actual low precision in mineral contribution to the total Na release rates (i.e. >30 %; Fig. 3b). The latter can be 472 explained by the fact that in PROFILE all types of plagioclase have the same dissolution rate coefficients (Table 473 3). Due to this, and in combination with the fact that plagioclase type minerals are a major source for Na, the 474 mineralogical uncertainty in estimating Na release rates with PROFILE was relatively low in this study (i.e. <20 475 %). In context, however, we note that it is generally accepted that under natural conditions different plagioclase 476 minerals weather at different rates, (Allen and Hajek, 1989, Blum and Stillings, 1995).

#### 477 4.3.2 Calcium release rates

- 478 According to  $W_{XRPD}$  and  $W_{A2M}$ , a key mineral for Ca release rates was calcic plagioclase at Flakaliden and epidote

- 480 of epidote and amphibole (Casetou-Gustafson et al., 2018) was directly reflected in the significant discrepancy 481 and overestimated weathering rates of Ca by  $W_{A2M-site}$  compared to  $W_{XRPD}$  (Fig. 5, and 1a). This discrepancy was 482 due to differences between  $W_{A2M-site}$  and  $W_{XRPD}$  in the mineral stoichiometry of calcic plagioclases, and not in 483 geochemistry, as the same geochemical analyses were also used for  $W_{A2M-reg}$ .
- 484

485 At Flakaliden, A2M based on site-specific mineralogy overestimated epidote at the expense of amphibole 486 (Casetou-Gustafson et al., 2018), leading to an underestimation of Ca weathering rates from amphibole compared 487 to epidote (Fig. 5). On the other hand, at Asa, it was the regional mineralogy input to A2M that resulted in 488 overestimated amounts of epidote at the expense of dioctahedral vermiculite and amphibole, and this bias was 489 directly reflected in the underestimated release of Ca from amphibole in WA2M-reg. Conversely, the relatively small and non-significant discrepancies of Ca release by WA2M-site at Flakaliden and by WA2M-reg at Asa did not depend 490 491 on a high precision in estimating the contribution from different minerals, since the precision was actually low. In 492 these cases, the good fits seem to be simply coincidental. Owing to differences in dissolution rates, Ca-bearing 493 minerals tend to compensate each other in terms of the total weathering rate that is calculated. This compensatory 494 effect is perhaps the reason why by coincidence, both WA2M-reg and WA2M-site discrepancies for Ca diverge in 495 different directions at Asa compared to Flakaliden.

496

497 Another source of uncertainty associated with the release of Ca is the role of minerals with high dissolution rates 498 that occur in low abundance, for example apatite, pyroxene and calcite. Apatite was included in M<sub>A2M</sub>, but if present 499 in the soils studied was below the detection limit of 1 wt.% in the XRPD analyses as were pyroxene and calcite 500 (Casetou-Gustafson et al., 2018). Additionally, the assumption made in the A2M calculations that all P determined 501 in the geochemical analyses is allocated to apatite will likely overestimate the abundance of this mineral since soil 502 P can also occur bound to Fe and Al oxides and soil organic matter in acidic mineral soils (Weil and Brady, 2016). 503 The relatively high abundance of paracrystalline Fe-oxyhydroxide and Al-containing allophane and imogolite at

- 504 Flakaliden indicates that this could be the case, at least at Flakaliden.
- 505

506 Regarding pyroxene, XRPD might also have failed to detect and quantify pyroxene due to low abundance at 507 Flakaliden (Casetou-Gustafson et al., 2018). Analytical limitations of XRPD would thus imply that W<sub>XRPD</sub> of Ca 508 might be underestimated at Flakaliden and Asa. However, in the absence of XRPD detection it is also possible that 509 M<sub>A2M-reg</sub> overestimated the pyroxene contents at Flakaliden. Thus, apatite and pyroxene added relatively large 510 uncertainties to the weathering estimates of Ca at Flakaliden due to the fact that they have a low abundance in 511 combination with very high dissolution rates. In terms of other reactive trace mineral phases, White et al. (1996, 512 2017) has highlighted the importance of small amounts of calcite in intact granitoid rocks and its significance for 513 Ca found in watershed studies. They also noted that in laboratory leaching experiments on the rocks they studied, 514 reactive calcite became exhausted after just 1.5 years. Given the trace concentrations involved and the high 515 solubility of calcite, it is doubtful that calcite is or has been of any long-lived significance in the soil profiles 516 studied, even though they are derived largely from rocks of granitic composition. Although, the results of White 517 et al. (1996, 2017) do suggest that calcite present in the in-situ granitoid rocks underlying the soils may well contribute to Ca export from the catchment. Additionally, the overestimation of the slowly weatherable mineral 518 519 illite by MA2M-reg (Casetou-Gustafson et al., 2018) resulted in an underestimation of Ca release by WA2M-reg at

- 520 Flakaliden, since less Ca was allocated to the more weatherable minerals. Although, it should also be noted
- 521 parenthetically that Ca can only occur as an exchangeable cation in illite, it is not an element that occurs as part of
- 522 the illite crystal structure, such that the 'illite' composition used in PROFILE is in need of some revision.

#### 523 4.3.3 Magnesium release rates

524 At both study sites, a large number of Mg-containing minerals contributed to the release of Mg, but amphibole 525 was the predominant mineral according to W<sub>XRPD</sub> and W<sub>A2M-site</sub>. The only significant discrepancy in Mg release 526 rates was revealed for W<sub>A2M-site</sub> at Asa, which resulted in an overestimation by 41 %. This overestimation was an 527 effect of underestimated contribution from amphibole in combination with overestimated contributions from 528 hydrobiotite and trioctahedral mica. This result for Asa supported our second hypothesis. At Flakaliden, WA2M-site 529 produced the same shift in the contribution of Mg by minerals, but the net effect was a very small and non-530 significant discrepancy to W<sub>XRPD</sub>. As was noted for Ca, the different outcomes of using site-specific mineralogies 531 at Asa and Flakaliden has no systematic underlying pattern.

Using PROFILE based on regional mineralogy resulted in surprisingly low and non-significant discrepancies in
Mg release rate, despite both the qualitative and quantitative mineralogies being very different from XRPD,
particularly at Flakaliden. For example, both pyroxene and illite were included in M<sub>A2M-reg</sub>, but not in M<sub>XRPD</sub>. Thus,

- at Flakaliden, the overestimation of illite in M<sub>A2M-reg</sub> caused an underestimation of Mg release rates comparable to
- the underestimation of Ca release rates.

#### 537 4.3.4 Potassium release rates

538 Weathering of K-feldspar was the most important source of K release by PROFILE regardless of the different 539 types of mineralogy input. Casetou-Gustafson et al. (2018) showed a strong negative correlation between MA2M-reg 540 and M<sub>XRPD</sub> for two of the major K-bearing minerals observed at both study sites, i.e., illite (or dioctahedral mica, 541 muscovite) and K-feldspar. Contrary to our second hypothesis, the results of the present study demonstrate that 542 over-or underestimation of WA2M-reg compared to WXRPD cannot be explained by significant negative correlation of 543 illite and K-feldspar in  $M_{A2M-reg}$ . However, this is likely related to the fact that illite and K-feldspar have the lowest 544 and also quite similar dissolution rates among minerals included in PROFILE (i.e. the highest dissolution 545 coefficients, Table 3). Although very different inputs in relation to K bearing minerals produced very similar 546 outputs, we note that this appears contradictory to differences in the behaviour of K-feldspars and K-micas as 547 sources of K via weathering to plants as reviewed for example by Thompson and Ukrainczyk (2002). Additionally 548 we note that Hodson et al. (1997) compared reaction rate constants for different minerals from Sverdrup et al. 549 (1990) with their own calculations and the discrepancies were relatively large for some minerals, e.g. muscovite.

#### 550 5. Concluding remarks

Based on comparing the full solution span of normative mineralogy from the A2M program to measured reference mineralogy from XRPD overall similar weathering rates were generated by the different mineralogical inputs to the PROFILE model. However, the underlying contributions from different minerals to the overall rates differed markedly. Although the similarity of overall rates lends some support to the use of normative mineralogy as input to weathering models, the details of the comparison reveal potential short-comings and room for improvements in the use of normative mineralogies.

- Compared with regional mineralogy, weathering rates based on site-specific mineralogy were more comparable to the reference rates generated from XRPD mineralogy, in line with hypothesis 1, at one of the study sites (Flakaliden), but not at the other (Asa). Thus, although intuitively the more detailed site-specific quantitative mineralogy data might be expected to give more comparable results, this is not supported by this study.
- For Ca and Mg the differences between weathering rates based on different mineralogies could be explained by differences in the content (modelled or actual) of some specific Ca- and Mg-bearing minerals, e.g. amphibole, apatite, pyroxene, calcite and illite. Improving certainty in relation to presence versus absence of some of these minerals and if present accurate quantification at low levels would reduce weathering rate calculation uncertainties.
- High uncertainties in mineralogy, due for example to different A2M assumptions, had surprisingly low effect on the weathering from Na- and K-bearing minerals. This can be explained by the fact that the weathering rate constants for the minerals involved, e.g. the plagioclase feldspars and K-feldspar and dioctahedral micas, are similar in PROFILE such that they compensate each other in the overall weathering rate outputs for these elements, a situation that is unlikely to reflect reality.
- For more in-depth analysis of the uncertainties in weathering rates caused by mineralogy, the rate
   coefficients of minerals should be revisited and their uncertainties assessed. A revision of rate constants
   could lead to results more in line with hypothesis 1.
- 575

#### 576 6. Authors contribution

Authors contributed to the study as in the following: S. Casetou-Gustafson: study design, data treatment, PROFILE
model analyses, interpretation and writing. C. Akselsson: study design, PROFILE model development,
interpretation and writing. B.A. Olsson: study design, interpretation and writing. S. Hillier: interpretation and
writing.

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721 **Table 1**. Characteristics of the study sites.

Asa	Flakaliden
57° 08' N; 14° 45 Έ	64° 07'N; 19° 27'E
225-250	310-320
688	523
5.5	1.2
Acidic intrusive rock	Quartz-feldspar-rich
	sedimentary rock
Sandy loam	Sandy loam
Sandy loamy till	Sandy loamy till
Udic	Udic
Spodosols	Spodosols
3	1
	Asa 57° 08' N; 14° 45 E 225-250 688 5.5 Acidic intrusive rock Sandy loam Sandy loamy till Udic Spodosols 3

<sup>a</sup> Bergh et al. 2005

<sup>b</sup>Long-term averages of annual precipitation and temperature data (1961-1990) from nearest SMHI meteorolgical stations (Asa: Berg; Flakaliden: Kulbäcksliden)

<sup>c</sup>SGU bedrock map (1:50000)

<sup>d</sup>Soil texture based on own particle size distribution analysis by wet sieving according to ISO 11277

<sup>e</sup>USDA Soil Conservation service, 2014

<sup>f</sup> Warfvinge and Sverdrup (1995)

Parameter	Description	Unit	Source	
Temperature	Site	°C	Measurements from nearby SMHI stations	
Precipitation	Site	m yr	Measurements from nearby SMHI stations	
Total deposition	Site	mmol <sub>c</sub> m <sup>-2</sup> yr <sup>-1</sup>	Measurements of open field and throughfall	
			deposition available from nearby Swedish ICP	
			Integrated Monitoring Sites	
BC net uptake	Site	mmol <sub>c</sub> m <sup>-2</sup> yr <sup>-1</sup>	Previously measured data from Asa and Flakaliden:	
			Element concentration in biomass from Linder	
			(unpublished data). Biomass data from Heureka	
			simulations.	
N net uptake	Site	mmol <sub>c</sub> m <sup>-2</sup> yr <sup>-1</sup>	Previously measured data from Asa and Flakaliden:	
			Element concentration in biomass from Linder	
			(unpublished data). Biomass data from Heureka	
			simulations.	
BC in litterfall	Site	mmol <sub>c</sub> m <sup>-2</sup> yr <sup>-1</sup>	Literature data for element concentrations from	
	~.		Hellsten et al. 2013	
N in litterfall	Site	mmol <sub>c</sub> m <sup>-2</sup> yr <sup>-1</sup>	Literature data for element concentrations from	
	<b>a</b> .	<b>D</b>	Hellsten et al. 2013	
Evapofraction	Site	Fraction	Own measurements and measurements from nearby	
	0.1		Swedish Integrated Monitoring Sites	
Mineral surface area	5011	m <sup>2</sup> m <sup>3</sup>	Warfwings and Swordman (1005)	
Soil bulk donsity	Soil	ka m <sup>-3</sup>	Own moosurmonts	
Soil moisture	Soil	$m^3 m^{-3}$	Based on paragraph 5.9.5 in Warfwinge and Sverdrup	
Son moisture	5011	111 111	(1995)	
Mineral composition	Soil	Weight fraction	Own measurments	
Dissolved organic	Soil	mg $l^{-1}$	Previously measured data from Asa and Flakaliden:	
carbon		C	Measurements for B-horizon from Harald Grip and	
			previously measured data from Fröberg et al. 2013	
Aluminium solubility	Soil	kmol m <sup>-3</sup>	Own measurements for total organic carbon and	
coefficient			oxalate extractable aluminium together with function	
			developed from previously published data	
			(Simonsson and Berggren, 1998)	
Soil solution CO2	Soil	atm.	Base on paragraph 5.10.2 in Warfvinge and Sverdrup	
partial pressure			(1995)	

**Table 2.** PROFILE parameter description.

- **Table 3** Mineral dissolution rate coefficients (kmol<sub>c</sub>  $m^{-2} s^{-1}$ ) used in PROFILE for the reactions with H<sup>+</sup>, H<sub>2</sub>O,
- 727 CO<sub>2</sub> and organic ligands (R<sup>-</sup>) (Warfvinge and Sverdrup, 1995).

Mineral	pkH	pkH2O	pkCO2	pKR
Pyroxene	12.3	17.5	15.8	14.4
Apatite	12.8	15.8	15.8	19.5
Hornblende	13.3	16.3	15.9	14.4
Epidote	14	17.2	16.2	14.4
Plagioclase	14.6	16.8	15.9	14.7
K-Feldspar	14.7	17.2	16.8	15
Biotite	14.8	16.7	15.8	14.8
Chlorite	14.8	17	16.2	15
Vermiculite	14.8	17.2	16.2	15.2
Muscovite and Illite	15.2	17.5	16.5	15.3

### 745 Figure captions

- **Figure 1.** The first scenario for describing the effect of mineralogy on weathering rates in the upper mineral soil
- for a specific soil profile (a) happens when the PROFILE weathering rate based on XRPD (reference weathering
- rates) is not contained in the weathering range produced using PROFILE in combination with the full A2M solution
   space. There are two possible explanations of why a significant discrepancy introduces an uncertainty range, i.e.
- (1) due to uncertainties related to the mineralogical A2M input and (2) due to uncertainties related to the limitation
- of the XRPD method itself (i.e. detection limit). The second scenario (b) occurs when the reference weathering
- rate is contained in the full A2M weathering span. In this case we speak of 'non-significant discrepancies'.
- **Figure 2.** Comparison of PROFILE weathering rates of base cations (mmol<sub>c</sub> m<sup>-2</sup> yr<sup>-1</sup>) at Asa (a) and Flakaliden (b)
- sites in the 0–50 cm horizon based on XRPD mineralogy (vertical dashed lines) with PROFILE weathering rates
- based on one thousand random regional A2M mineralogies versus one thousand random site-specific A2M
- mineralogies. Data presented are from four different soil profiles per site. Regional graph for soil profile 10B at
  Flakaliden is missing since A2M did not calculate 1000 solutions for soil layer 20-30, due to "Non-positive
  solution".
- Figure 3. Root-mean square error (RMSE) of average PROFILE weathering rates (mmol<sub>c</sub> m<sup>-2</sup> yr<sup>-1</sup>) of one thousand
   A2M mineralogies per soil layer, compared to weathering rates based on XRPD mineralogy per soil layer.
   Comparisons are based on the total weathering per element (A) and on the sum of mineral contributions to total
- 762 weathering per element (B). RMSE describes the prediction accuracy for a single soil layer.
- Figure 4. Comparison of PROFILE weathering rates based on XRPD mineralogy (mmol<sub>c</sub> m<sup>-2</sup> yr<sup>-1</sup>) with PROFILE
   weathering rates based on regional A2M mineralogy (upper figures) versus site-specific mineralogy (lower
   figures). Each data point represents a mean of one thousand PROFILE weathering rates for a specific soil depth of
   one of 4 soil profiles per site.
- 767 Figure 5. Comparison of sums of PROFILE base cation weathering rates for different minerals in the upper 768 mineral soil (0-50 cm) based on XRPD mineralogy and the average PROFILE base cation weathering rate (i.e. 769 based on one thousands input A2M mineralogies per mineral) according to the two normative mineralogical 770 methods and for each study site (i.e. As a site-sepcific, Flakaliden site-specific, As a regional, Flakaliden regional). 771 For W<sub>A2M</sub>, relative error (% of W<sub>XRPD</sub> estimate) are given at the end of each bar to illustrate the average deviation 772 of  $W_{A2M}$  and  $W_{XRPD}$  in the upper mineral soil. \*=significant discrepancy as defined in section 2.7. 773 Vrm1=Trioctahedral vermiculite; Vrm2=Dioctahdreal vermiculite. Information on chemical compositions of 774 minerals are given in Table S5 and S6.
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806 Figure 2a





813 Figure 3, A, B













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Pyroxene

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Figure 5