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# Inherited control of crystal surface reactivity

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1	Inherited control of crystal surface reactivity					
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10 Abstract: Material and environmental sciences have a keen interest in the correct prediction of material 11 release as a result of fluid-solid interaction. For crystalline materials, surface reactivity exerts 12 fundamental control on dissolution reactions; however, it is continuously changing during reactions and 13 governs the dynamics of porosity evolution. Thus, surface area and topography data are required as 14 input parameters in reactive transport models that deal with challenges such as corrosion,  $CO_2$ 15 sequestration, and extraction of thermal energy. Consequently, the analysis of surface reaction kinetics 16 and material release is a key to understanding the evolution of dissolution-driven surface roughness and 17 topography. Kinetic Monte Carlo (KMC) methods simulate such dynamic systems. Here we apply these 18 techniques to study the evolution of reaction rates and surface topography in crystalline materials. The 19 model system consists of domains with alternating reactivity, implemented by low vs. high defect 20 densities.

Our results indicate complex and dynamic feedbacks between domains of high versus low defect density, with the latter apparently limiting the overall dissolution rate of the former - a limitation that prevails even after their disappearance. We introduce the concept of "inherited" control, consistent with our observation that maximum dissolution rates in high defect density domains are lower than they would be in the absence of low defect density neighboring domains.

The controlling factor is the spatial pattern of surface accessibility of fluids. Thus, the distribution of large etch pits centers is inherited almost independently of spatial contrasts in crystal defect density during ongoing reactions. As a critical consequence, the prediction of both the material flux from the reacting surface and the evolution of topography patterns in crystalline material is constrained by the 30 reaction history. Important applications include the controlled inhibition of reactivity of crystalline 31 materials as well as the quantitative evaluation and prediction of material failure in corrosive 32 environments.

Keywords: Kinetic Monte Carlo simulation, rate spectra, crystal dissolution, surface reactivity, surface
 topography and roughness patterns

35

## 36 1 Introduction

37 Crystal defects cause the formation of etch pits during dissolution and corrosion processes. A large 38 body of literature exists about this process, ranging from BCF theory to the stepwave model (e.g., Casey 39 et al., 1988; Cheng, 1993; Dove et al., 2005; Lasaga and Luttge, 2001; Luttge et al., 2013; Tang et al., 40 2003). The growth and coalescence of etch pits controls the evolution of reacting rough surfaces and 41 pore walls and affects thus porosity in natural and technical materials. Examples include the formation 42 of secondary porosity in reservoirs (Taylor et al., 2010) or the impact of microstructures on corrosion 43 (Horlait et al., 2014). As a consequence, small-scale approaches for modeling pore-scale reaction 44 processes are gaining increased interest (Iliev et al., 2015; Steefel et al., 2015). The evolution of porosity 45 alters the fluid-solid interface and, thus, the effective reactivity of the dissolving material (Daval et al., 46 2013; Dele-Afolabi et al.; Dong et al., 2011; Fischer et al., 2014; Xu et al., 2016). By changing the 47 surface roughness and topography of the reacting pore walls and throats (Anbeek, 1992), the evolution 48 of new void volumes constrains via permeability alternations both the diffusive and advective transport 49 of dissolved material and thus impacts reactive flow conditions (Anovitz and Cole, 2015; Gao et al., 50 2017; Iliev et al., 2015; Mostaghimi et al., 2016). Moreover, coupled dissolution-precipitation reactions 51 rely on the formation of voids as initial pore volumes as this allows the fluid to maintain contact with 52 the reacting interfaces (Chen et al., 2014). Accordingly, the quantitative predictability of porosity 53 evolution during fluid-solid interaction is of great interest in multiple disciplines of material and 54 environmental sciences, and the development of appropriate tools is at the forefront of fundamental and 55 applied research (Steefel et al., 2015; Yang et al., 2015). The formation of void volumes significantly 56 influences the efficiency of fluid-solid interactions by affecting the rate of solute transport. Related 57 data are required as input parameters in reactive transport models dealing with questions such as  $CO_2$ 

sequestration (Hellevang et al., 2013; Monger et al., 2015), extraction of thermal energy (Dillinger et
al., 2014), as well as waste management in rock formations (Kim et al., 2011).

60 Crystal surface reactivity is an important but variable boundary condition that constrains the evolution 61 of porosity. The defect density variability in crystalline materials is a critical parameter that defines 62 spatial domains of contrasting reactivity (Hiemstra and van Riemsdijk, 1999). The accessibility of 63 domains with constant or contrasting surface reactivity does change continuously over reaction time. 64 Thus, changing surface-energetic complexity does superimpose the changing geometric complexity. As 65 a consequence, the boundary conditions of the reacting system change dynamically. The analysis of 66 such a system with multiple changing input parameters is a classic task for a modelling approach. 67 Kinetic Monte Carlo (KMC) simulation calculations offer a large enough length scale to address this 68 problem appropriately (Lasaga and Blum, 1986).

69 As a starting point, we study the evolution of reaction rates and surface topography pattern in a simple 70 Kossel crystal (Kossel, 1927; Stranski, 1928) by utilizing KMC simulation techniques. More 71 specifically, the model material is built up of domains of constant activation energies but having varying 72 defect densities in order to provide well-defined variations in the surface reactivity (Liu et al., 2017; 73 Liu et al., 2015; Luttge et al., 2013). We apply the rate spectra concept to KMC results in order to 74 analyze systematically the rate contributors in the frequency domain of rate datasets (Fischer et al., 75 2012; Fischer et al., 2014). Using this concept, the identification of both invariance or temporal 76 variability of such rate contributors over reaction time provides quantitative insight into the overall 77 reaction rates of crystalline materials (Fischer and Luttge, 2017; Michaelis et al., 2017; Saldi et al., 78 2017).

In detail this study addresses the following questions: (1) How variable are reaction rates during dissolution of multiple domains or grains of crystalline material with variable defect density? Do such domains provide constraints to the effective surface reactivity? (2) How variable are the resulting pore volume and etch pit distribution? What parameters govern the initial formation of topography patterns in crystalline material? Is there any predictive power associated with such a parameter? (3) To what extent is reactivity inherited in crystalline material? Are low-reactivity domains able to slow down the overall reaction rate significantly with consequences such as reduction in surface reactivity?

## 87 2 Methods

## 88 2.1 Kinetic Monte Carlo simulations

89 A Kinetic Monte Carlo (KMC) program simulates a series of single reactions at the atomic scale. Here, 90 we apply a KMC model for a *Kossel* crystal (Kossel, 1927). A Kossel crystal consists of a cubic lattice 91 composed of a single chemical species. The bond energies  $\Phi$  between all atoms are constant. The KMC 92 model is based on the Bortz-Kalos-Lebowitz (BKL) algorithm (Bortz et al., 1975). The algorithm allows 93 for a reactive event at each iteration step. The simulation provides the temporal evolution of a dissolving 94 surface, based on a series of single reaction events on the crystal surface. Lasaga and Luttge (2004) 95 provide detailed and general information about KMC models for mineral dissolution. In this study, the 96 model simulates dissolution reactions under conditions far from equilibrium.

97 KMC models are now routinely used for studying surface site-specific reaction kinetics (e.g., Cama et 98 al., 2010; Kerisit et al., 2008; Kurganskaya and Luttge, 2016; Meakin and Rosso, 2008). Implemented 99 parameters in KMC algorithms include the lattice resistance effect or the network of surface hydrogen 100 bonds. An increasing number of ab initio calculations provides the critical parameters for the probability 101 equations. Thus, the reactivity of solid materials without defects is properly implemented in KMC 102 approaches. In this study, we focus on the impact of varying defect densities in crystals. The reported 103 variability in natural and technical crystalline material is huge and includes several orders of magnitude 104 (Blum and Lasaga, 1987), similar to the ranges utilized in this study (Table 1).

105 We utilize periodic boundary conditions to eliminate the influence of model size in (x,y) direction. In 106 order to investigate material domains of varying reactivity, we apply a "layered cake" model of a solid, 107 composed of layers with variable thickness and defect densities. The type of crystal defects in this study 108 are screw dislocations. Screw dislocations form an initial structure during the dissolution process, i.e., 109 the hollow core. Ongoing dissolution results in the formation of etch pits along the hollow core. Our 110 model has well-defined input parameters, i.e., the number of layers, layer thickness, and defect density 111 (Table 1). Defects are randomly seeded in each layer by using uniformly distributed random numbers 112 for (x,y) core center positions. The defect length is identical to the domain thickness.

**Table 1:** KMC simulation parameters for the dissolution of KOSSEL crystals with a lattice constant of 0.2 nm. Several KMC runs utilized crystals that are built-up of several domains in z-direction (domain thickness) that are characterized by alternating defect densities. In this case, a structure starting with a high (h) reactivity domain followed by a low (l) reactivity domain was analyzed and vice versa, e.g., *hlhl & lhlh*, see **Fig. 2**.

Simulation results in	system size (area)	domain thickness	defects/ domain	number of domains	dissolved atoms	number of runs
Fig. 1	50×50 nm <sup>2</sup>	10 nm	50	1	1E+06	1
Figs. 2, 3	300×300 nm <sup>2</sup>	1 nm	1000 ( <i>h</i> ) 10 ( <i>l</i> )	4	1E+06	2
			1000	1	1E+06	1
Fig. 3C	300×300 nm <sup>2</sup>	10 nm	100	1	1E+06	1
			10	1	1E+06	1
Figs. 4, 5	300×300 nm <sup>2</sup>	10 nm	150 ( <i>h</i> ) 15 ( <i>l</i> )	2	10E+06	2
Fig. 6	300×300 nm <sup>2</sup>	2 nm	100 ( <i>h</i> ) 10 ( <i>l</i> )	10	6E+06	2
	300×300 nm <sup>2</sup>	20 nm	100 ( <i>l</i> )	1	6E+06	1
		2.0 nm	100 vs. 10	10		
Fig. 7		2.2 nm	110 vs. 11	10		
		4.0 nm	200 vs. 20	10		
		20 nm	1000(h)	1		

115 The simulation starts with the opening of hollow cores in the top layer. As soon as the dissolving layer 116 exposes a defect outcrop in the subjacent layer, a hollow core opens and a new etch pit evolves (Burton 117 et al., 1951; Lasaga and Luttge, 2001). KMC runs were performed under constant bond strength 118 conditions ( $\Phi = 4$ , normalized to *kT* units). The dissolution probability *P<sub>i</sub>* of an atom with *i* bonds is

$$P_i = e^{\left(-\frac{\Phi}{kT}\right)} \tag{1}$$

Multiple types of reactive domains were used in layered cake models. We investigated the impact of defect density and domain thickness in different KMC simulation calculations, details about the parameter variation of the crystals are summarized in **Table 1**. We varied the starting conditions of the model, i.e., the reactivity of the uppermost layer was either (h) or (l).

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1

## 125 **2.2 Reaction rate analysis: rate maps and rate spectra**

126 Rate maps are calculated using the formula:

- 127
- 128  $R(x, y, t_i) = \frac{h(x, y, t_{i-1}) h(x, y, t_i)}{t_i t_{i-1}} = \frac{\Delta h_i(x, y)}{\Delta t_i}$ (2)
- 129

130 R(x,y,t) is the local dissolution rate for a single node, defined by (x,y) coordinates, and i is the time 131 step. Such rates reflect the increase (growth) or decrease (dissolution) of the initial height h of a given 132 point (x, y) during a time interval  $(t_i - t_{i-1})$ . The frequency distribution of the R values forms a rate 133 spectrum (Fischer et al., 2012). The simulations output the surface height maps at fixed iteration steps, 134 the distributions of the reactive sites, and the overall material flux as a function of time. The iteration 135 steps are defined by the number of dissolved blocks. The reaction time increment,  $\Delta t_i$ , between the 136 events i - 1 and i, is calculated as a reciprocal sum of the number N of probabilities P of surface 137 reactions normalized by the reaction attempt frequency,  $\nu$  (Blum and Lasaga, 1987), i.e., the 138 fundamental frequency (Pelmenschikov et al., 2001), that we set to  $10^{12}$  [s<sup>-1</sup>]:

139 
$$\Delta t_i = \frac{1}{\nu \sum_{j=1}^N P_j} \tag{3}$$

140 This permits the tracking of the reaction progress and to analyze kinetic parameters such as overall rate, 141 kink and step site densities in time units (TU, see below). The applied fundamental frequency number 142 is in the order of magnitude of the intermolecular vibrational frequency of bulk water (Walrafen, 1972). 143 The frequency of the reaction attempts is similar to this frequency because the dissolution process is 144 water-catalyzed (Pelmenschikov et al., 2001). Consequently, bond strengths are utilized for probability 145 calculations.

#### 147 **3 Results and discussion**

## 148 **3.1 Quantitative analysis of surface reaction rates**

- 149 Material flux maps provide information about the spatial distribution of heterogeneous material flux.
- 150 Flux maps of KMC calculation results offer the opportunity to analyze the mechanistic connection
- between crystal surface reactions and material flux at any time step. Figure 1 exemplifies how this
- analysis reveals information about material flux characteristics.
- 153 Based on KMC results, a model for crystal dissolution via the stepwave movement mechanism has been
- 154 introduced (Lasaga and Luttge, 2001). According to this model, crystal dissolution is controlled by
- 155 stepwaves emanating from dislocation cores (**Fig. 1A; B**). The evolution of superimposing stepwaves
- and their efficiency in terms of material removal is quantitatively illustrated by using material flux maps
- 157 (Fig. 1C) and the corresponding rate components (Fig. 1D).



**Figure 1:** Visualization of results of a Kinetic Monte Carlo dissolution experiment using a *KOSSEL* crystal structure. (A, B) Topography with etch pits after removal of 100'000 and 200'000 blocks. (C) Material flux map shows inert (black) and reactive surface sections related to surface rates. Note the distribution and shape of reactive surface portions: a. highly reactive etch pit wall sections (yellow/red), b. single stepwave of an extinct screw dislocation, c. flux distribution related to complex stepwave structure with change of surface reactivity at an etch pit wall. (D) Rate spectrum of rate map (C) showing the dominant frequency (green color) of rate contributions owing to dissolution stepwaves.

158 As an example, large single stepwaves from inactive screw dislocations (at the height of screw bottoms)

159 represent low material fluxes. Superimposition of stepwaves from multiple generators results in flux

- 160 map sections that illustrate a complex pattern of high vs. low material flux distribution.
- 161



**Figure 2**: (A) Surface topography evolution of a KOSSEL crystal that consists of four layered domains having alternating defect densities, starting with a high defect density (1000 defects/domain), followed by a low defect density (10 defects/domain). (B) Topography evolution similar to (A), but starting with a low defect density (10 defects/domain), followed by a high defect density (1000 defects/domain). The column sketches in (A) and (B) illustrate the impact of the contrasting defect densities (low (*l*): blue; high (*h*): red) on etch pit size and number. (C) Evolution of RMS roughness Rq of both materials (A) and (B). Map size =  $300 \times 300 \text{ nm}^2$ . Note the difference in reaction time compared to the dissolution of an identical number of surface blocks, e.g. (1)-red vs. (1)-blue due to initial differences in surface reactivity (see A, B).

162

## 163 **3.2 Reactivity of layered crystals with domains having varying defect densities**

164 We compare the dissolution rates of KOSSEL crystals that are composed of domains of contrasting

- 165 screw defect densities. We vary the initial conditions with respect to the defect density of the uppermost
- 166 crystal domain, i.e., high (*h*) vs. low (*l*) reactivity (**Table 1**). Figure 2 illustrates the evolution of the
- 167 surface topography and roughness (**Fig. 2 A**: *h*, **B**: *l*). As expected, the initial topographic contrast of
- 168 the surface maps of both crystals is caused by the defect density (Anbeek et al., 1994). A high defect

169 density results in the formation of multiple etch pits having a small lateral extension (Fig. 2A, 1-2), a 170 low defect density results in the formation of a few pits with large lateral size (Fig. 2B, 1). Subsequently, 171 the dissolution of the subjacent crystal domain leads to the superimposition of small and large etch pits, 172 according to the defect density of the newly-exposed crystal domains. However, the roughness 173 evolution (Fig. 2C) is dominated by the topography of large and deep etch pits. The change in-between 174 low- vs. high-reactivity has no clear equivalent in the topography evolution. The reason is that the 175 evolution and growth of a few large pits is a dominant factor of the overall evolution of surface 176 topography. Consequently, the RMS roughness (Thomas, 1999) increases over the total reaction time. 177 Figure 3 shows the rate maps of the dissolving crystalline material. Figure 3A illustrates the locally 178 low rates associated with the dissolution of material having a high defect density (Fig. 3A, [1-2], blue 179 color). The growth of etch pits within crystal domains with low defect densities results in locally 180 enhanced reaction rates (Fig. 3A, [4], green color). The existence of steep etch pit walls (Fig. 2) dictates the further evolution of surface rates and topography, i.e., the increase in spatial heterogeneity of 181 182 material flux from the surface. A similar evolution is observed for crystal domains with an inverse order 183 of defect density (Fig. 3B). The initial contrast in surface rates of high vs. low defect density domains 184 is clearly visible in Fig. 3C, sections [1-2] of both h and l domains. The large initial contrast of the 185 material flux is decreased irreversibly during the ongoing reaction, see Fig. 3C, all map sections later 186 than [2]. For comparison, Fig. 3C shows the rate evolution during dissolution of KOSSEL crystals with 187 constant defect densities (10, 100, 1000 defects/unit volume). The initial results of the material release 188 s are similar to the above investigations, see Fig. 3C, sections 1-2 of both h and l domains and 189 homogeneous crystals with high (1000) and low (10) defect densities. However, the subsequent rate 190 evolution of both layered crystals is similar to those of the homogenous crystal with the lowest defect 191 density (blue dotted line in Fig. 3C). Thus, we hypothesize that crystal domains with low defect 192 densities act as inhibitors. Below, we investigate this point in greater detail.



**Figure 3:** Surface rate evolution of the KOSSEL crystal shown in **Fig. 2**. (A) Rate maps resulting from the dissolution of four layered domains having alternating defect densities, starting with a high defect density (1000 defects/domain), followed by a low defect density (10 defects/domain). (B) Surface rate evolution similar to (A), but starting with a low defect density (10 defects/domain), followed by a high defect density (1000 defects/domain). (C) Evolution of the rate [expressed by total number of removed blocks] of both materials (A) and (B). For comparison, simulation results are shown of single domains having 1000, 100, and 10 defects, respectively. Map size =  $300 \times 300$  nm<sup>2</sup>.

## 193 **3.3 Rate spectra analysis of reaction inhibition**

194 We analyzed rate spectra in order to investigate the relationship between reaction kinetics, surface 195 topography, and potential inhibition of surface reactivity. Figure 4 illustrates the surface evolution (Fig. 196 4A-D) and the respective rate spectra (Fig. 4A'-D'). During dissolution of the upper crystal domain, 197 the evolution of both surface topography and rate spectra reflects the situations of low defect density 198 (Fig. 4A, A') vs. high defect density (Fig. 4C, C'), cf. Table 1. Both rate spectra show an asymmetric 199 shape. The less reactive crystal domain is characterized by a spectrum with a tail towards lower rate 200 components. Accordingly, the more reactive crystal domain shows the opposite behavior. Then, the 201 underlying domain gets partly and later completely exposed to dissolution during the ongoing removal 202 of the upper domain. Figures 4B, D illustrate the crystal surface topography after complete removal of

- the uppermost domain. At this time, both surfaces are dominated by large etch pit structures. It is a surprising result that the high defect density of the newly exposed material in **Figure 4 B** does not
- 205 impact significantly the reaction rate.



**Figure 4:** Surface topography and rate spectra evolution of two different layered cake model structures. Left (A,B) with low (blue, upper part) and high (red, lower part) reactivity domains (dt =  $5 \times 10^9$  s). Right (C,D) shows inverse order of domain reactivity (dt =  $6 \times 10^9$  s). Corresponding rate spectra (A', B') and (C', D') are shown below, time unit (TU) =  $6 \times 10^9$  s. System size =  $300 \times 300$  nm<sup>2</sup>.

- 206 Etch pits related to the high defect densities do open and their frequent occurrence is visible at the walls
- 207 of the large pit structure. Their spatial evolution is geometrically constrained and hindered by the large
- 208 pit structure which produces surface steps much faster than the newly-formed pits. Both the lifetime of
- 209 pits and their impact on the overall rate are thus restricted by the final stepwave velocity dictated by the

210 large pit (Lasaga and Luttge, 2001). This result is illustrated by the characteristic shape of the rate 211 spectrum (Fig. 4B'). Here, the expected asymmetric shape towards higher rates (cf. Fig. 4 C') is not 212 observed. Instead, the rate spectra shapes of contrasting domains are fairly similar (Figs. 4 B' and D'). 213 This result shows that the initial low surface reactivity and the corresponding surface evolution are able 214 to constrain the effective reactivity of a material at later development stages, regardless of its defect 215 density. Thus, a proxy such as defect density is not in general able to serve as an input value for the 216 prediction of reactivity and material flux. Based on defect density we would expect a change to higher 217 material flux and thus a higher reaction rate (Fig. 4 B, B'). The rate spectra analysis, however, does 218 show that the expected elevated reactivity of the material is limited due to the previous reaction steps.

219

## 220 **3.4 Evolution of material flux and surface topography**

We investigate the evolution of an important reactive crystal surface site density, i.e., kink sites, and compare their frequency with the evolution of the cumulative material flux (overall void volume) and the rough topography of the reacting surface. In **Figure 5**, the transition between domains of contrasting reactivity is indicated by pairs of dashed lines.



**Figure 5**: (A) evolution of kink site number, (B) evolution of total dissolved volume, (C) evolution of local void volume (related to rough surface topography) in two different settings (black line, initially high reactivity, gray line: initially low reactivity). Blue and red letters refer to the reactivity of crystal domains, see Fig. 2. Pairs of dashed lines refer to the interface between domains of contrasting reactivity, i.e., initial exposition of underlying domain and complete removal of upper domain.

This transition represents the reaction period that starts with the initial exposure of material from the underlying domain and ends with the complete dissolution of material from the upper domain. The contrast in reactivity of the upper domains in both systems (low – i vs. high – iii) is illustrated by the contrast of surface kink site concentration during the initial reaction period (**Fig. 5 A**). Thus, the material flux from the more reactive domain with higher defect density is higher (red vs. blue circles). While the void volume formation and the kink site density during this reaction period are not identical, the void volumes of the rough topographies are similar (**Fig. 5**). 232 This picture changes during the reaction period that is required to remove the material of the 233 upper domain completely. The concentration of kink sites of the highly-reactive domain drops 234 significantly (red  $\rightarrow$  black circles; iii $\rightarrow$ iv). This is caused by the high consumption rate of the 235 material of the highly-reactive domain and synchronously the low rate of etch pit and kink site 236 formation in the underlying low-reactivity domain. Consequently, the cumulative material flux 237 decelerates (Fig. 5 B) and the corresponding surface pore volume (Fig. 5 C) shows a minimum 238 value. One would expect that this picture does reverse after the complete removal of the upper domain 239 of the initially low-reactive crystal (blue  $\rightarrow$  open circles;  $i \rightarrow ii$ ). However, the interesting result is that 240 both the kink site density and the overall material flux of the highly reactive domain (Figure 5: open 241 symbols) do not mirror the picture that we observed initially. Instead, the kink site density of both 242 experiments (Fig. 5 A, right section) fluctuates around a mean value similar for both systems. Some fluctuations during the dissolution period of the crystal domain with higher defect density (see section 243 244 (ii)) are caused by an undisturbed growth of newly-formed etch pits. A similar behavior is observed for 245 the void volumes (Fig. 5 B,C). These results explain the observations of the limited reaction rates that 246 are illustrated in Figure 4 B & 4 B', see above. Our interpretation is as follows: While the defect density 247 of the highly-reactive domain supports the frequent opening of pits, the life-time of their stepwaves is 248 comparatively short. Consequently, the reactivity of the system is no longer defined by its defect density 249 but rather by the existence of inherited etch pit positions and their large size compared to newly-formed 250 pits (Fig. 4B). The position of such pits has been inherited from the previously dissolving domain 251 because the bottom of such deep pits provide the initial exposure to the subjacent crystal domain. As 252 soon as a surface section of the latter had been exposed, the seeding of a new pit started because of the 253 high probability of exposing a defect and opening of a hollow core.



**Figure 6:** (A) Temporal variability of the material flux of dissolving KOSSEL crystals with 10 domains having alternating reactivity, implemented by the number of defects per domain (10 vs. 1000). The blue-colored curve indicates results of a crystal starting with a low-reactivity domain, the red-colored curve represents a crystals having a high-reactivity starting domain. Topography maps (amber-colored) illustrate the respective topographies. (B) Rate map (topography difference map based on topography maps (38 - 33) identifies large etch pit walls as the source of highest rate portions.

#### 256 **3.5 Analysis of larger systems**

257 The results in Figs. 3 and 4 suggest the inhibition of surface reactivity by crystal domains with low 258 reactivity. In this subchapter we ask the question whether this result is valid during the long-term 259 evolution of dissolution of multiple crystal domains (e.g., Taleb and Stafiej, 2011). We investigate such 260 behavior by using KOSSEL crystals that consist of a larger number layered domains (Table 1). Figure 261 **6** shows the results of the long-term simulations using a system that consists of 10 layers. Here, the 262 initial sections of the rate curves show the expected contrast, i.e., a higher number of removed blocks 263 of the highly-reactive surface portion compared to the low reactivity material at the identical time step, 264 see also **Figure 3C**. Subsequently, both rate curves indicate results that are typical for systems with low 265 surface reactivity and, thus, low reaction rates. Similar to the shorter simulation runs (Figs. 2-5), the 266 reactivity of the dissolving crystal is dictated by the low-reactivity domains. The surface maps of both 267 systems explain such behavior (Fig. 6). The long-term evolution of the reacting crystal surfaces is 268 characterized by the evolution of large single pits. This typical result illustrates the dominant growth of 269 large pits on the expense of small pits. Regardless of the opening of new etch pits, the further growth 270 of existing large pits is the dominating process (Fig. 6B).



**Figure 7:** (A) Long-term rate evolution as a function of domain thickness. Three domain thicknesses (2.0, 2.2, 4.0 nm with alternating defect density high vs. low) are investigated. Low (blue) and high (red) defect density (dd) domains are shown for comparison. The dotted line represents a shorter reaction period similar to those analyzed in Figure 3.

272 The results in Figure 7 illustrate the impact of varying domain thickness on the overall reactivity and 273 material flux during dissolution. The grey-colored lines represent different thicknesses of the alternating 274 domains (**Table 1**). We compare the results with systems having constant defect densities. Overall, the 275 reduction in reactivity of the crystals having domains of contrasting defect density is found again, 276 similar to observations of alloy corrosion (Renner et al., 2006). Our results suggest a complex impact 277 of the domain thickness on the dissolution of a series of domains. The change of domain thicknesses 278 causes a non-linear change in reactivity and rates of the compound. Accordingly, both slope and position 279 of inflection points suggest a complex interplay in-between defect density/ etch pit number, domain 280 thickness, and material flux/ reaction rate. As an example, the crystal with domain thicknesses = 4.0 nm 281 shows a higher rate compared to the other examples (2.0 nm and 2.2 nm). Nevertheless, the elevated 282 reactivity that is represented by the crystal without any domains is not reached during the entire reaction 283 period. Moreover, the reactivity of this crystal (d = 4.0 nm) having high-reactivity domains drops 284 temporarily below the reactivity of the low-defect density crystal. Geometrical constraints related to the 285 evolution of rough surface topography during the etch pit growth are a potential explanation.

Figure 7 illustrates another important characteristics of any reacting crystal, exemplified by the bluecolored rate curve. During the comparatively long reaction period, the reaction rate increases as shown by the deviation from the dashed line. Such behavior can be explained by the occurrence of large etch pit walls having high kink site densities as the major portion of the overall crystal surface. Similar results have been reported for the dissolution reaction of grains of single crystals (Briese et al., 2017; Petrik and Harbrecht, 2013). Thus, an additional important parameter that exerts control on the system's reactivity is the domain size and the related opportunity to form large etch pits with a characteristic high surface roughness and topography.

294 Previous attempts to quantify the surface rates vs. etch pit nucleation focused on pit size distribution analysis (MacInnis and Brantley, 1993). Such approaches compared the varying size, distribution, and 295 296 coalescence of pits with reaction rates of single crystal surfaces (Luttge, 2005). Experimental 297 approaches showed how spatial heterogeneities in both surface kinetics and material transport properties 298 evolve during single crystal dissolution (Peruffo et al., 2013). As a result, the morphology of the 299 characteristic etch features may reveal direction-dependent dissolution kinetics. Our approach utilizes 300 this knowledge and applies it to more complex model systems in order to extract important new 301 information about the evolution of reactivity. More specifically, the temporal evolution of rate spectra 302 informs about material flux variability and relates it to etch pit formation, growth, and coalescence 303 (Fischer and Luttge, 2017). Rate acceleration and rate jerk data suggested the pulsating behavior of 304 surface reactivity (Fischer and Luttge, 2018). Overall, the modeling results focusing on varying defect 305 density underscore the critical role of reactivity anisotropy in crystalline material for the evolution of 306 surface roughness and void volume pattern due to dissolution. More specifically, these results show 307 how the spatial distribution of high vs. low reactivity domains in crystal grains impacts the evolution 308 of void volume patterns in materials that are characterized by an identical mean defect density. This 309 approach might thus serve as an example of how the present KMC results are applicable to explain the 310 formation of initial pattern of void volume in solids. Examples include the corrosion of steel (Sun et al., 311 2015), the formation of secondary porosity due to contrasting defect density pattern in feldspar grains 312 and cements (González-Acebrón et al., 2012; Götze et al., 2000), and the dissolution in battery 313 electrolyte (Steiger et al., 2014). For such systems, we expect the existence of concurrent mechanisms 314 that affect surface reactivity. Such mechanisms include passivation processes during mineral surface 315 reactions due to the formation of altered surface layers (e.g., Daval et al., 2011; Wild et al., 2016).

#### 317 **5** Conclusions

318 The analysis of Kinetic Monte Carlo simulation results by employing the rate spectra concept provides 319 new insight into the reactivity of crystalline material. The combination of new results about the 320 reduction in reactivity of highly-reactive crystal domains with the quantitative concept of rate spectra 321 may serve as a base for the development of a powerful tool to predict the reactivity of complex 322 crystalline materials. In detail we conclude: The reactivity of a crystalline system can be governed by 323 its reaction history. The dissolution of low-reactivity domains within crystalline matter defines the 324 subsequent reactivity. Crystal domains with high defect densities inherit a lower reactivity close to 325 reacting crystal domains with lower defect densities. Such a mechanism contributes to the explanation 326 of the reduction in surface reactivity of polycrystalline materials. Our results provide a quantitative 327 background to answer questions related to corrosion and dissolution in the material and earth sciences, 328 such as the predetermination of void volume formation with consequences for permeability evolution, 329 as well as for radioactive waste management problems that involve materials with contrasting defect 330 densities.

331

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