

LETTER:

CALCITE DISSOLUTION UNDER TURBULENT FLOW
CONDITIONS: A REMAINING CONUNDRUM

PROBLEM NASTANKA FASET IN RAZTAPLJANJA KALCITA
V TURBULENTNEM TOKU

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UDK 551.435.8:549.742.111

INTRODUCTION

A significant body of experimental and theoretical work has examined the dissolution rates of calcite, and other carbonate minerals, under varying chemical and hydrodynamic conditions (see Morse & Arvidson (2002) for a comprehensive review). The relationships derived from this work have been applied extensively to the development of mechanistic models of speleogenesis (e.g. Dreybrodt 1996; Dreybrodt *et al.* 2005; Birk *et al.* 2005; Rehl *et al.* 2008; Kaufmann 2009; Szymczak & Ladd 2011). However, the primary focus of these models has been on the early stages of cave formation, with less attention toward the later stages of cave evolution and turbulent flow conditions. Recent efforts have begun to develop mechanistic models for processes governing later stages of cave evolution, considering factors such as turbulent flow structures (Hammer *et al.* 2011) and open channels (Perne 2012). However, such studies remain limited, in part due to significant quantitative uncertainties in a va-

riety of processes that become important beyond the incipient speleogenesis stage (Covington *et al.* 2013).

While speleogenetic models have not typically been run much beyond the transition from laminar to turbulent flow conditions, experimental and theoretical studies of carbonate dissolution have constrained dissolution rates and rate-controlling mechanisms under turbulent flow (e.g. Rickard & Sjöberg 1983; Dreybrodt & Buhmann 1991; Liu & Dreybrodt 1997). However, a direct application of these results and comparison to field observations leads to an apparent conundrum. According to the theory, solutional forms such as scallops and flutes should not exist in limestone; however, they clearly do exist at a wide variety of sites and scales. This conundrum suggests that there may be problems with the theory, problems with our understanding of scallop formation, or both.

THE CONUNDRUM

Dissolution rates under turbulent flow conditions are typically calculated by dividing the fluid into two regions, a turbulent core that is well-mixed, and a diffusion boundary layer (DBL) that lies between the turbulent core and the dissolving wall (Dreybrodt & Buhmann

1991). Within the DBL, flow is presumed to be laminar and species cross the layer via the process of molecular diffusion. There are two end-member regimes in which overall dissolution rates are limited by either diffusion rates (the transport-limited regime),

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Received/Prejeto: 26.11.2013

$$F_D = \frac{D(C_{eq} - C_b)}{\epsilon}, \quad (1)$$

or surface reaction rates (the reaction-limited regime),

$$F_s = k_1(1 - C_b/C_{eq}), \quad (2)$$

where F_D and F_s are the transport-, and reaction-limited fluxes, respectively, D is the diffusion constant, C_{eq} is the equilibrium concentration of calcium, C_b is the concentration of calcium in the bulk solution, ϵ is the DBL thickness, and k_1 is the kinetic rate constant for calcite dissolution in the linear kinetic regime. The mixed kinetic regime occurs when the diffusion-limited and surface-reaction-limited rates are similar (Rickard & Sjöberg 1983). In this case, both processes are important. For simplicity, I only consider linear dissolution kinetics

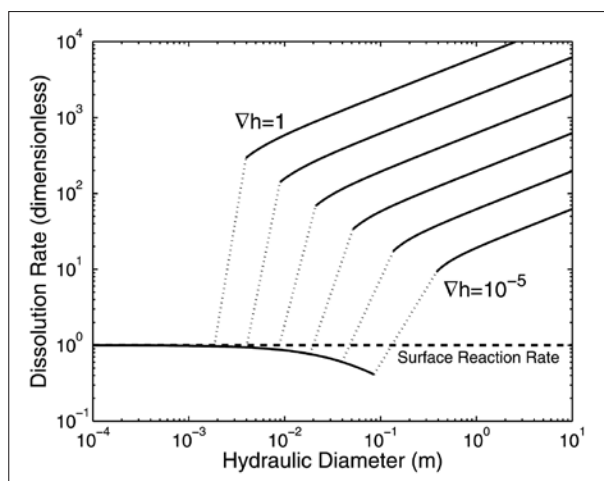


Fig. 1: Diffusional (solid) and surface reaction (dashed) dissolution rates as a function of hydraulic diameter, D_{H1} , depicted for different hydraulic head gradients, ∇h . Sharp jumps in diffusion rates occur at the onset of turbulent flow. The laminar/turbulent transition is indicated with dotted lines. For the parameter space shown, the theory suggests that diffusional rates only control dissolution for low-gradient conduits just below the laminar/turbulent transition. Elsewhere, diffusional rates are much higher than surface reaction rates. Figure reproduced with permission from Covington *et al.* (2012).

here, as is common below about 80% saturation, though our conclusions are not sensitive to this choice. In the case of a sufficiently thick DBL, or a sufficiently thin turbulent core, conversion of CO_2 can also be rate-limiting (Dreybrodt & Buhmann 1991). Though non-linear kinetics and CO_2 conversion limitation are not explicitly considered here, if anything, they would exacerbate the presented conundrum by providing further means to reduce surface reaction rates.

In order to determine the typical conditions under which rates were limited by either reaction or transport, Covington *et al.* (2012) calculated surface reaction and diffusion rates for a wide variety of head gradients and

hydraulic diameters using the Darcy-Weisbach equation and Colebrook-White relation. For these calculations, the fiducial parameter values given in Dreybrodt *et al.* (2005) were used, with $k_1 = 4 \times 10^{-11} \text{ mol cm}^{-2} \text{ s}^{-1}$, $D = 10^{-5} \text{ cm}^2 \text{ s}^{-1}$, and $C_{eq} = 2 \times 10^{-6} \text{ mol cm}^{-3}$. For ease of comparison, Covington *et al.* (2012) converted the “surface” rate law to the form

$$F_s = \alpha_s(C_{eq} - C), \quad (3)$$

where $\alpha_s = 2 \times 10^{-5} \text{ cm s}^{-1}$ was determined by dividing k_1 by C_{eq} . Here I have slightly modified the notation of Covington *et al.* (2012), replacing α with α_s to clarify that the constant applies to the chemical reaction rate at the surface. In many previous manuscripts (e.g. Buhmann & Dreybrodt 1985) α without a subscript represents the combined effects of diffusion and reaction.



Fig. 2: Scallops formed on a limestone surface within a cave, indicating contrasts in dissolution rates as a result of turbulent flow structures. Major divisions on the ruler are in cm (Photo: Matija Perne).

Covington *et al.* (2012) compared the rates predicted by Equations 1 and 3 by nondimensionalizing the two equations by dividing by the term $\alpha_s(C_{eq} - C)$, such that the dimensionless dissolution rates are independent of chemistry and become

$$F_s^* = 1, \quad (4)$$

and

$$F_D^* = \frac{D}{\alpha_s \epsilon}, \quad (5)$$

where the stars denote dimensionless rates. Then ϵ was calculated for a wide variety of head gradients and hydraulic diameters to examine the relative magnitude of the reaction- and transport-limited equations. The DBL thickness was calculated using equations 2.13 and 2.14 from Dreybrodt *et al.* (2005), which are equivalent to Equations 9 and 10 of this manuscript. For these calculations a fractional roughness of 0.05 was assumed. The results of this calculation (Fig. 1) show that, according to the equations used above, the surface reaction rate is almost always limiting, and no cases in the turbulent flow

regime show transport-limited rates. In fact, diffusion rates under turbulent flow are typically several orders of magnitude larger than the surface reaction rate. This conclusion leads to the conundrum. The model suggests that dissolution rates should almost always be controlled by the surface reaction rate in the turbulent flow regime, and, therefore, that dissolution rates should be independent of any spatial variations in DBL thickness. On the con-

trary, caves and channels formed in limestone frequently contain scallops (Fig. 2) and flutes that apparently form as the result of systematic contrasts in DBL thickness as a result of turbulent flow structures (Curl 1966; Goodchild & Ford 1971; Blumberg & Curl 1974; Curl 1974). The remainder of this note will explore this conundrum in more detail, and discuss possible resolutions.

SURFACE RATES FROM THE PWP EQUATION

The above analysis makes use of fiducial kinetic constants from Dreybrodt *et al.* (2005). However, these constants are motivated by typical values from experiments, where the effects of transport and reaction are both present. Therefore, a more accurate approach is to use the full Plummer-Wigley-Parkhurst (PWP) equation (Eqn. 6, Plummer *et al.* 1978) to calculate true theoretical surface rates. If the PWP rates were high enough, then they could explain away the conundrum.

For comparison, I calculate PWP rates that result from a temperature and partial pressure of CO_2 that approximately reproduce the fiducial value of $C_{\text{eq}} = 2 \times 10^{-6} \text{ mol cm}^{-3}$ used above. This equilibrium concentration is roughly obtained (for an open system) with $T = 10 \text{ }^\circ\text{C}$ and $P_{\text{CO}_2} = 0.01 \text{ atm}$, values which are also quite typical in the cave environment. The PWP rates are depicted in Fig. 3, alongside the simplified linear relationship with two different values of α_s .

The form of the PWP equation over the relevant range is approximately linear for this example, except for the highly undersaturated end, suggesting that use of a linear function is reasonable in this case, except at nearly zero dissolved Ca. However, comparing the fiducial value of α_s used to create Fig. 1 above, with the best-fit value for the linear region of the PWP equation (Fig. 3), suggests that the value of α_s from Dreybrodt *et al.* (2005) is too small by about a factor of 3 to 4 in order to approximate the PWP rate. Consequently, the surface reaction line in Fig. 1 should be shifted up by a similar factor of 3 to 4. While this does imply that the surface rates are higher

than would be predicted by the equation and parameter values given in Tab. 2.1 of Dreybrodt *et al.* (2005), the more accurate surface rates obtained from PWP are still much lower than the diffusive rates are predicted to be for the majority of the turbulent region of the parameter

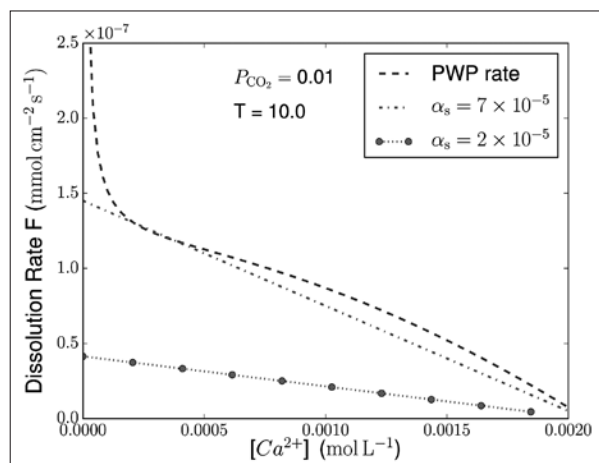


Fig. 3: The dissolution rate as a function of Calcium concentration for $P_{\text{CO}_2} = 0.01 \text{ atm}$ and $T = 10 \text{ }^\circ\text{C}$. This example is constructed to have a similar value of C_{eq} and temperature to the fiducial case from Dreybrodt *et al.* (2005).

space (Fig. 1). For the convenience of future studies, a fitting function for PWP rates as a function of temperature and P_{CO_2} is provided in the Appendix. This relation is not used explicitly here, but provides a quick means of estimating PWP rates in a given setting.

COMPARING SURFACE-REACTION AND DIFFUSION RATES

The equations given above for diffusion-limited (Equation 1) or surface-limited rates (Equation 3) are only val-

id under conditions where dissolution is truly limited by the respective process. In steady state, diffusion and sur-

face reaction rates must be equal, and the rate of diffusion is given by

$$F_D = \frac{D(C_s - C_b)}{\epsilon} \tag{6}$$

where C_s is the concentration at the surface, and C_b is the concentration in the bulk flow. For the diffusion-limited case $C_s \rightarrow C_{eq}$, and we recover Equation 1 above. Similarly, surface reaction rates are dependent on the concentration at the surface and are given by

$$F_s = \alpha_s(C_{eq} - C_s) \tag{7}$$

Again, if dissolution is reaction-limited $C_s \rightarrow C_b$ and we recover Equation 3. Setting diffusion and reaction rates

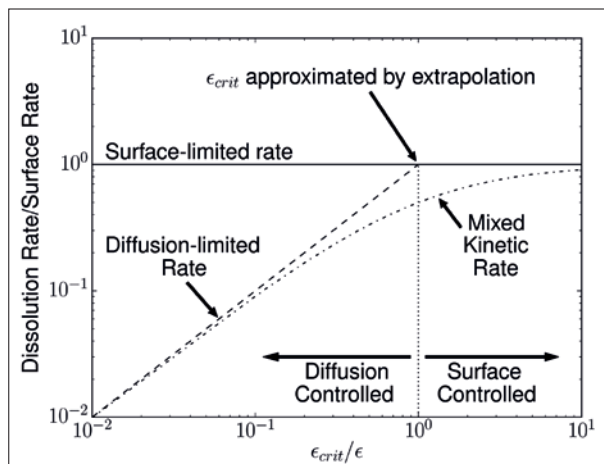


Fig. 4: The critical DBL thickness, below which dissolution rates are surface reaction-limited, can be estimated by extrapolating the diffusion-limited rate until it intersects the surface-limited rate determined by the PWP Equation. This leads to the relation in Equation 10. Though this approximation scheme can be employed more generally, the lines plotted in this figure represent exactly the case where the rates can be approximated using α_d and α_s .

equal, solving for C_s , and plugging this back into Equation 7, one can see that the dissolution rate accounting for both reaction and diffusion is given by

$$F = \frac{\alpha_s \alpha_d}{\alpha_s + \alpha_d} (C_{eq} - C_b) \tag{8}$$

where $\alpha_d = D/\epsilon$. The lines depicting the diffusion-limited rates in Fig. 1 are equivalent to α_d/α_s , so the analysis and conclusions of Covington *et al.* (2012) can also be cast in terms of the relative magnitudes of α_d and α_s . Therefore, if dissolution rates are well-represented by the linear relation in Equation 7, then the relative importance of diffusion and reaction in determining the overall rates can be quantified by comparing α_d and α_s . A critical diffusion boundary layer thickness, at which diffusion and reaction are equally important can be calculated by setting $\alpha_d = \alpha_s$, leading to

$$\epsilon_{crit} = D/\alpha_s \tag{9}$$

When $\epsilon \gg \epsilon_{crit}$ then rates are diffusion-limited, and when $\epsilon \ll \epsilon_{crit}$ rates are reaction-limited. If $\epsilon \sim \epsilon_{crit}$ then dissolution will occur via mixed kinetics. At $\epsilon = \epsilon_{crit}$ diffusion suppresses dissolution rates to half of what they would be from surface reaction alone.

The above analysis relies on a surface reaction rate of the linear form given by Equation 7. However, it can be generalized using the full PWP dissolution rate. In the limit of a thick DBL the diffusion-limited equation can be applied, while in the limit of thin DBL the PWP surface rate can be applied. An estimate of ϵ_{crit} can be

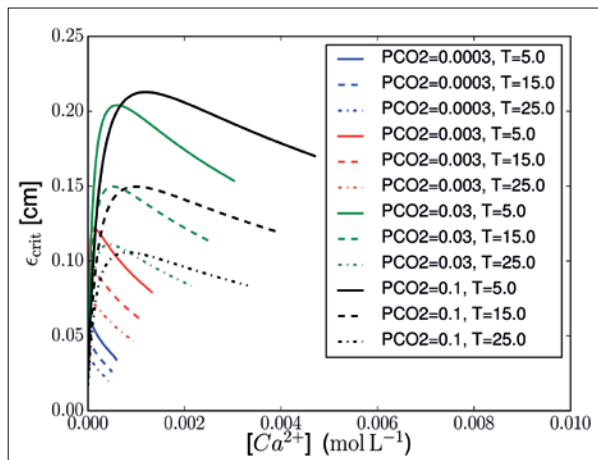


Fig. 5: The critical DBL thickness, below which dissolution rates are surface reaction-limited, shown for a wide range of temperature, P_{CO_2} values, and dissolved loads. Lines terminate on the right-hand side at saturation.

obtained by extrapolating the diffusion-limited equation until it intercepts the PWP surface rate (Fig. 4). Quantitatively, this results in the relation

$$\epsilon_{crit} = \frac{D(C_{eq} - C)}{F_{pwp}} \tag{10}$$

When the DBL thickness is near this critical value, dissolution will occur via mixed kinetics, but when the DBL is much larger or smaller than ϵ_{crit} then dissolution rates will be limited by either diffusion or surface reaction, respectively. Using this relation, and the full PWP equation, one can calculate critical DBL thicknesses for a range of temperatures, P_{CO_2} values, and calcium concentrations. This calculation (Fig. 5) shows that the critical DBL thickness is on the order of magnitude of 1 mm for essentially the entire range of temperature, P_{CO_2} , and dissolved load found in natural karst systems, except in highly undersaturated conditions where $C > 0$ and $\epsilon_{crit} \ll 1$ mm.

The approximation $\epsilon_{\text{crit}} \sim 1$ mm can also be reproduced using a rough estimate of PWP rates. In most natural settings, the forward reaction rate is dominated by reaction III (κ_3 in the PWP equation) (Dreybrodt 1988). This rate is simply a function of temperature and is on the order of 10^{-10} mol cm $^{-2}$ s $^{-1}$ (Dreybrodt 1988, p. 127). Critical DBL thickness can then be estimated as

$$\epsilon_{\text{crit}} \sim \frac{DC_{\text{eq}}}{\kappa_3} = \frac{(10^{-5} \text{ cm}^2 \text{ s}^{-1})(2 \times 10^{-6} \text{ mol cm}^{-3})}{10^{-10} \text{ mol cm}^{-2} \text{ s}^{-1}} = 0.2 \text{ cm} \quad (11)$$

which is quite similar to the result of 1 mm obtained from the larger parameter study (Fig. 5).

Within speleogenesis models DBL thickness is usually estimated from a relationship for pipe flow that employs the Sherwood Number (Sh),

$$\epsilon = D_H / \text{Sh}, \quad (12)$$

where Sh is given by an empirical relationship such as

$$\text{Sh} = \frac{(f/8)(\text{Re} - 1000)\text{Sc}}{1 + 12.7\sqrt{f/8}(\text{Sc}^{2/3} - 1)} \quad (13)$$

(Dreybrodt *et al.* 2005). Here, Re is the Reynolds number, Sc is the Schmidt number, and f is the Colebrook-White friction factor. Using these relations, one can calculate typical values of DBL thickness, (ϵ), for a selection of head gradients and hydraulic diameters (Fig. 6). Again, for this calculation a relative pipe roughness of 0.05 is assumed, though the result is not particu-

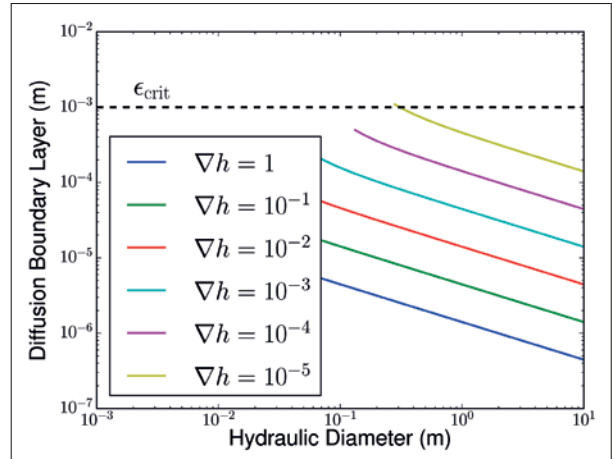


Fig. 6: DBL thickness (ϵ) under turbulent flow conditions as estimated from Sherwood number for a wide range of hydraulic parameters. Each line represents a choice of head gradient. The lines terminate on the left at $\text{Re} = 4000$, where the Colebrook-White formulation breaks down. Typical values of DBL thickness as estimated from Sherwood Number are much less than the 1 mm magnitude of the critical value determined from the PWP equations.

larly sensitive to this choice. Typical values of the DBL thickness under turbulent conditions, as calculated from Sherwood Number, are much less than the order of magnitude estimate of a critical DBL thickness of 1 mm. This is true for all turbulent flow cases except those at very low head gradients (i.e. 10^{-5}).

CONCLUDING THOUGHTS AND POTENTIAL RESOLUTIONS

A more careful analysis reinforces the conundrum. The theory suggests that limestone dissolution rates under turbulent flow conditions should be typically limited by surface reaction rates. However, field observations of scallops and other solutional forms clearly suggest otherwise. A recent model developed using PWP dissolution equations and computational fluid dynamics also found that limestone dissolution flutes were unstable under the calculated dissolution rates, though flutes in gypsum, which has much higher surface rates, proved to be somewhat more stable (Hammer *et al.* 2011). Superficially, this result is in agreement with the analysis presented here, and may provide further evidence of difficulties with the current theory. The central purpose of this letter is to clearly state the problem, rather than to solve it, in hopes of stimulating some discussion on the subject. However, I here discuss a few potential resolutions.

Perhaps the most suspicious component of the turbulent dissolution model is the semi-empirical equa-

tion used to calculate Sherwood Number (Eqn. 13), and consequently DBL thickness (Eqn. 12). This equation is based on experiments in smooth pipes, and therefore ought to be used with caution when calculating DBL thickness for the rough and irregular surfaces found on natural bedrock channel walls. However, the experiments of Blumberg & Curl (1974) provide a means of checking this as a potential resolution. Mass transfer data from their experiments with flutes in gypsum suggest a typical DBL thickness of $\epsilon = 0.0089 L \text{ Sc}^{-1/3}$, where L is the flute length and Sc is the Schmidt Number, with $\text{Sc} \gtrsim 1000$ for water at the relevant temperatures. This results in a DBL thickness that is roughly $\approx 0.1\%$ of the flute length. In combination with the estimate of critical DBL thickness of 1 mm for limestone, this result suggests that such solutional forms should develop, but only on length scales of a meter or greater. This again is in conflict with field observations, where scallops and flutes often form on length scales of centimeters (Fig. 2).

Another possibility is that the rates estimated by the PWP equation are significantly lower than rates on natural surfaces. In fact, several studies of dissolution rates measured on rotating disks produce rates that are approximately twice those given by the PWP equation (Dreybrodt & Buhmann 1991; Liu & Dreybrodt 1997). These studies have suggested that increased surface roughness, and therefore surface area, on the disks may result in the higher apparent dissolution rates. Similarly, roughness on natural limestone surfaces on a smaller scale than the DBL thickness could result in an apparent increase in surface reaction rates that would reduce the critical DBL thickness. However, the observed discrepancy of a factor of two is not sufficient to explain away the conundrum by itself. If natural surfaces are still significantly rougher at scales below the DBL thickness than the disks used in experiments, this could potentially resolve the problem. Other factors, such as microbial films, could also influence surface reaction rates, and perhaps in some cases increase them.

One might ask whether a very small contrast in dissolution rates would be sufficient to form flutes and scallops. However, flute experiments show that dissolution rates vary by roughly a factor of two over the length of a flute (Blumberg & Curl 1974), a number that is thought to be relatively constant across the parameter space (Curl

1966). A contrast in dissolution rates of a factor of two requires that the thickest portion of the DBL within the scallop is greater than or equal to the critical thickness, as this is the DBL thickness when surface rates are suppressed by roughly a factor of two. A thinner DBL would not allow sufficient contrast in rates.

Forms somewhat similar to scallops (mechanical erosion flutes) also are found in bedrock channels in relatively insoluble rocks, though the variety and prevalence of such forms is greatest in highly soluble rocks (Richardson & Carling 2005). One possible explanation is that so-called solutional forms, such as scallops and flutes, actually form by a combination of solutional and mechanical processes. For example, chemical processes could loosen individual grains that are later plucked from the surface by mechanical processes.

Finally, it could be that scallops form only in highly aggressive waters, where the critical DBL thickness is sufficiently small (Fig. 5). However, the author's experience would suggest that scallops are also present in locations without such highly aggressive water. Little systematic attempt has been made to study the locations in which scallops form, and whether forms differ according to hydrological or lithological settings. Such studies might also provide clues as to the correct resolution of the current conundrum.

ACKNOWLEDGMENTS:

I would like to acknowledge Franci Gabrovšek, Matija Perne, Joe Myre, Wolfgang Dreybrodt, and Douchko Romanov for stimulating discussions and useful comments. This material is based upon work supported by the National Science Foundation under Grant No. EAR

1226903. Any opinions, findings, and conclusions or recommendations expressed in this material are those of the author and do not necessarily reflect the views of the National Science Foundation.

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APPENDIX: AN APPROXIMATION FOR THE PWP EQUATION

The PWP equation is roughly a linear function of $(C_{\text{eq}} - C)$ between 10% and 90% of saturation for a wide range of temperatures and P_{CO_2} values. To facilitate future work with PWP rates, I present a fitting function for the rate constant α_s ,

$$\log(\alpha_s) \approx A + B_1 T + B_2 \log(P_{\text{CO}_2}) + C_1 T^2 + C_2 [\log(P_{\text{CO}_2})]^2, \quad (14)$$

where the best fit values of the parameters are given in Tab. 1. To determine this relationship, I calculated PWP rates for 100 calcium concentration values from 10% to 90% saturation for each of 100 different values of P_{CO_2} sampled evenly in log space in the range $3 \times 10^{-4} \leq P_{\text{CO}_2} \leq 0.1$ and for 11 different values of temperature in the range $0 \text{ }^\circ\text{C} \leq T \leq 24 \text{ }^\circ\text{C}$. For each choice of T and P_{CO_2} a best fit value of α_s for the relation $F_{\text{pwp}} = \alpha_s(C_{\text{eq}} - C)$ was calculated using least squares. This resulted in 1100 values of α_s for a range of T and P_{CO_2} conditions. Typical residuals between the linear relation and the full PWP equation are about 10%. The maximum residuals for any of the fits are around 70% and occur near saturation for cases with high P_{CO_2} .

Parameter	Best fit value
A	-4.30
B_1	1.40×10^{-2}
B_2	0.150
C_1	-2.83×10^{-5}
C_2	8.76×10^{-2}

Table 1: Best fit parameters for the equation for the linear approximation to the PWP dissolution rate equation.

The parameters in the best fit relation above (Eqn. 14) were determined via least squares fitting of these 1100 values of α_s . Eqn. 14 reproduces the best fit values of α_s within 10%. Therefore, this relation is not intended for precision work, but can allow quick estimation of PWP rates according to a simpler relation. In addition to the fitting relation, the Python code used to calculate PWP rates is available online at <http://www.speleophysics.com>. All calculations shown above employed the full PWP equation rather than the approximation given by Eqn. 14.