

Crystal Growth Kinetics of Calcite in a Dense Fluidized-Bed Crystallizer

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The growth kinetics of calcite crystals are studied in a batch-fluidized-bed crystallizer, which is maintained at a constant pH. The growth experiments are conducted in the metastable region explored as part of this research. The crystal growth rates are evaluated from the consumption rates of calcium ions, using the cured natural calcite or silica sand as seeding materials. Several operation variables are investigated, including supersaturation, pH, ionic strength, superficial velocity, and particle size and type of seed. The significant factors that affect the crystal growth rate are identified. Then the crystal growth data of constant pH and ionic strength are analyzed by the two-step growth model. The mass-transfer coefficients are obtained and compared at various crystal sizes and superficial velocities. Finally, a growth-rate equation of calcite crystal, which is based on the two-step growth model, is proposed for design purposes.

Introduction

Recently a pellet reactor, which is a reactive, fluidized-bed, growth-type crystallizer, was developed for water softening, fluoride and phosphate removal, and heavy-metal recovery. The water feed and the chemical reagents required to cause deposition are fed to the reactor containing suspended seeds (Dirken et al., 1990; Seckler et al., 1990; van Dijk and Wilms, 1991). In water softening, the undesired species, calcium ion, reacts to form calcium carbonate and then grows on the seeds, which are later removed from the reactor after they exceed a certain size. In the operation of the pellet reactor the flow pattern in the fluidized bed resembles an ideal plug flow with a limited backmixing, which gives a higher conversion as compared to the ideal mix-flow reactor (Levenspiel, 1972). In the design of a pellet reactor for water softening, we need to know the crystal growth kinetics of calcium carbonate and the hydraulics of a fluidized bed. It seems that the former is less understood than the later. The reported growth-rate equation of calcium carbonate is rather empirical, for example, an overall growth-rate model (van Dijk and Wilms, 1991):

$$\frac{d[\text{Ca}^{2+}]}{dt} = K_T A \{ [\text{Ca}^{2+}] [\text{CO}_3^{2-}] - K'_{sp} \}, \quad (1)$$

where $-(d[\text{Ca}^{2+}]/dt)$ is the depletion rate of $[\text{Ca}^{2+}]$ in the solution, which is the same as the crystal growth rate of CaCO_3 in moles per unit time; K_T is a constant; A is the specific surface area of seed crystals; K'_{sp} is the concentration solubility product of calcite; and $\{[\text{Ca}^{2+}][\text{CO}_3^{2-}] - K'_{sp}\}$ represents the driving force for crystal growth.

Several mechanisms regarding crystal growth have been proposed in the literature (Mullin, 1993), among which the two-step growth model is considered the most useful from the chemical engineering point of view. At steady-state conditions, the two steps can be described mathematically by the following equations:

$$G = K_d(\sigma - \sigma_i) \quad \text{mass transfer} \quad (2)$$

$$= K_r \sigma_i^r \quad \text{surface reaction.} \quad (3)$$

Although this model is a simplified scheme for crystal growth, it reveals a great deal of useful kinetic information, which was recently summarized by Tai (1997). The systems studied are mostly soluble salts using seeded techniques. Similar studies of sparingly soluble salts are limited, because large single-seed crystals of the system are difficult to prepare and the application of the technique was limited in the past.

The single-seed crystals of calcite, one of the barely soluble salts, were successfully prepared by using the gel growth

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technique, and the crystal growth kinetics of calcite grown in a stirred vessel was investigated using the two-step growth model to analyze the growth-rate data (Tai et al., 1993). It is concluded that the mass-transfer resistance and surface-reaction resistance are significant. The mass-transfer resistance is related to the relative velocity between crystal and solution and increases with a decrease in relative velocity. Under the well-suspended conditions, the relative velocity between crystal and solution in a fluidized bed is lower than that in a stirred vessel, thus the mass-transfer resistance as well as the surface-reaction resistance for crystal growth of calcite should be considered in a fluidized bed. Therefore, an overall growth-rate equation, such as Eq. 1, is not an adequate expression for scale-up purposes, because different hydrodynamic behaviors exist between scales.

Besides the hydrodynamic effects, the crystal growth of barely soluble salts are affected by the chemical properties of the supersaturated solution. For example, Stubicar et al. (1990, 1992) found that the solution properties, including ionic strength, pH, and lead-to-fluoride ion activity ratio, influenced the crystal growth kinetics of lead fluoride; α -PbF₂ and β -PbF₂ appeared at different pH values, and crystal growth rates were higher at higher ionic strength.

The aim of this study is to investigate the crystal growth kinetics of calcite in a dense fluidized-bed crystallizer using the pH-stat method. Experiments were first conducted in a pH-stat vessel to identify the metastable region, where nucleation is suppressed, for the CaCl₂-Na₂CO₃-H₂O system. Then the growth experiments were performed in the metastable region using three crystal sizes, namely 460, 650, and 920 μ m, which are prepared by sieving the cured natural calcite. The crystal size is the mean size of the apertures of the two closest ASTM standard sieves. For example, crystals of size 460 μ m means the fraction of crystals retained between ASTM sieves Nos. 40 and 35, with nominal apertures of 420 and 500 μ m, respectively. The crystal growth rate was evaluated from the depletion rate of [Ca²⁺] in the solution. The growth-rate data were then interpreted in terms of the two-step growth model, thus the mass-transfer coefficient and surface-reaction coefficient were determined. The effects of operating variables, such as ionic strength, pH, superficial velocity, crystal size, and seed type, were explored. The obtained crystal growth-rate data will serve as the design basis of a pellet reactor for water softening.

Experimental Procedure

Determination of relative supersaturation

The expression of the driving force used for crystal growth of barely soluble salts is the relative supersaturation proposed by Nielsen and Toft (1984):

$$\sigma = (K_{ip}/K_{sp})^{1/2} - 1, \quad (4)$$

where K_{ip} is the ionic product, defined as $K_{ip} = a_{Ca^{2+}} \cdot a_{CO_3^{2-}}$; and K_{sp} is the activity solubility product of CaCO₃. The activity of species, $a_{Ca^{2+}}$ or $a_{CO_3^{2-}}$, is the product of the activity coefficient and the concentration of the respective species. The concentrations of ionic species were computed from the measured pH, total calcium concentration, and total

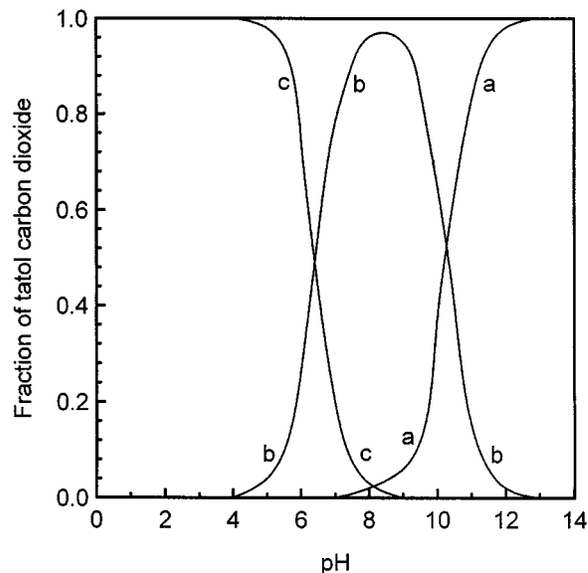


Figure 1. Fraction of total CO₂ present as the respective ions at various hydrogen-ion concentration: (a) CO₃²⁻; (b) HCO₃⁻; (c) H₂CO₃ (ASTM, 1994).

carbonate concentration by successive approximation for the ionic strength, using a computer program that contains mass-action equations, mass balance equations, charge-balance equations, and the modified Debye-Hückel equation for calculating the activity coefficient (Nancollas, 1996; Tai et al., 1993). In the operation of a pellet reactor, supersaturation should be controlled in a suitable range; a low supersaturation gives a low growth rate, which means a low removal rate of species, and a high supersaturation causes nucleation, which messes up the operation. The supersaturation is related to the concentration of CO₃²⁻, which is a function of pH at a fixed concentration of total carbonate. Therefore, as judged from Figure 1, which is a diagram showing the fraction of total carbon dioxide present as the respective ions at various pHs (ASTM, 1974), a suitable operational range of pH is roughly between 8.5 and 10.5, a pH lower than 8.5 giving a low CO₃²⁻ concentration and a pH higher than 10.5 absorbing too much CO₂ from air. The effect of pH on the relative supersaturation, which is obtained by computing, is clearly shown in Figure 2. It means that the control of pH is important in the experiment of studying the crystallization kinetics of CaCO₃.

Identification of metastable region of calcite

A key to the successful operation of pellet reactor is good control of supersaturation, which is generated by the chemical reaction. To suppress nucleation of calcite in growth experiments, supersaturation should be kept in the metastable region, which is the area between supersolubility and solubility curves (Nielsen and Toft, 1984). Since the supersolubility curve of the CaCl₂-Na₂CO₃-H₂O system is not available, experiments were first performed to identify the metastable region. Under constant temperature (25°C) and constant pH

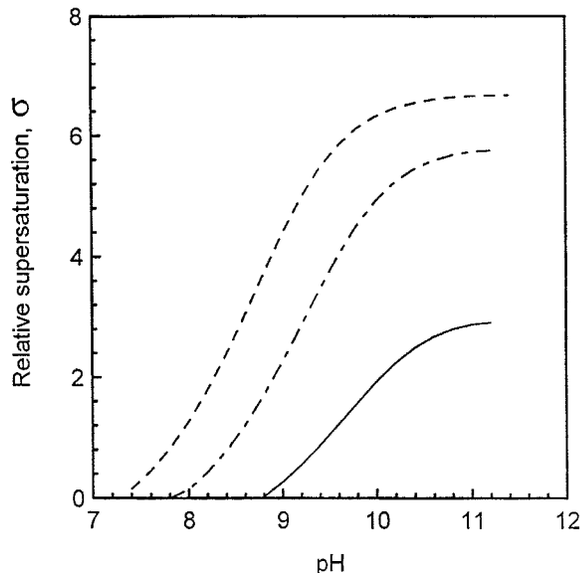


Figure 2. Relative supersaturation of calcium carbonate salt as a function of pH: $T = 25^{\circ}\text{C}$ and $[\text{Ca}]_T = 0.0005 \text{ kmol/m}^3$.

--- $[\text{CO}_3]_T = 0.05 \text{ kmol/m}^3$; - · - $[\text{CO}_3]_T = 0.005 \text{ kmol/m}^3$; — $[\text{CO}_3]_T = 0.0005 \text{ kmol/m}^3$.

(8.5, 9.5, and 10.5, respectively), 100 mL of CaCl_2 and Na_2CO_3 solutions were mixed and stirred for 2 h. For a specific concentration of calcium chloride, the solution remained clear at a low concentration of sodium carbonate. When the concentration of Na_2CO_3 increased, the solution became turbid due to nucleation. The boundaries were recorded for various CaCl_2 concentrations, such as those shown in Table 1 for a pH of 8.5. Then the concentrations or activities of Ca^{2+} and CO_3^{2-} at the boundaries were calculated using the same computer program as was used for estimating supersaturation. Finally, the metastable region was constructed and all the growth experiments were performed in this region.

Crystallization system

The crystallization system, containing a pH-stat fluidized-bed crystallizer, a storage tank, and a pH control system, as shown in Figure 3, was used to measure the growth rate of calcite crystal. The main part of the fluidized-bed crystallizer is a PVC column with a distributor at the bottom. Immediately above the column is an enlarged section to prevent seed crystals from carrying over to the storage tank. A supersaturated solution, which was prepared by mixing CaCl_2 and Na_2CO_3 solutions, of 6 L with a desired pH was charged into the storage tank and another 0.5 L of the same solution was fed into the fluidized bed. The solution in the storage tank was then pumped through a distributor and into the fluidized bed. After that, the solution overflowed to the storage tank. When the flow rate and pH became steady, the pump was stopped and approximately 25 g of seed crystals were introduced into the crystallizer. The pump was restarted and the growth experiment began. The pH of the solution was maintained constant during the operation, using a pH-stat apparatus

Table 1. Demarcation of Metastable Region for Calcium Carbonate (CaCO_3) at pH = 8.5 and $T = 25^{\circ}\text{C}$

$[\text{Na}_2\text{CO}_3]$ (kmol/m^3)	$[\text{CaCl}_2]$ (kmol/m^3)						
	0.05	0.025	0.015	0.005	0.0025	0.0015	0.0005
0.05							T
0.025							T
0.0175							T
0.015							C
0.005	T*			T	T	T	C
0.004							T
0.0035							T
0.00325							C
0.003							C
0.0025	T			T	T		C
0.00225							C
0.002							C
0.0015				T			C
0.00125	T						
0.001				C			
0.000875	T						
0.0005	T	T	T	C	C		
0.00045	T						
0.0004	T		T	C			
0.000375			C				
0.00035	T						
0.0003				C			
0.00025	T	T		C			
0.000225	T	C					
0.0002	C**	C					
0.00015	C						
0.00005	C						

*T means turbid.

**C means clear.

tus to control the amount of NaOH solution added to the storage tank. A 3-mL sample of the solution was withdrawn from the crystallizer every 30 min by a syringe fitted with a

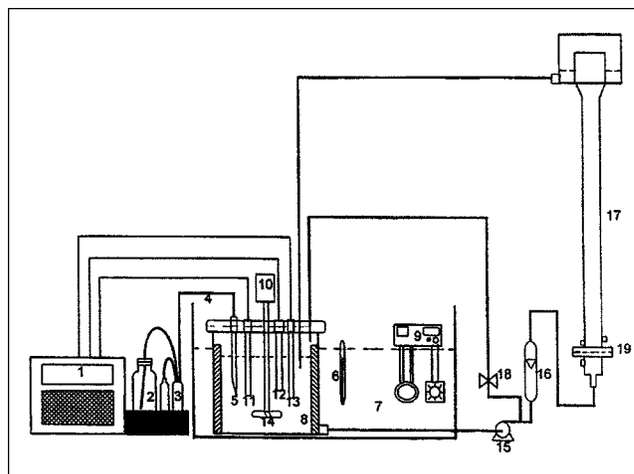


Figure 3. pH-stat crystallization system.

1. pH and temperature indicator; 2. reagent bottle; 3. pumping system of reagent; 4. reagent delivering line; 5. burette; 6. thermometer; 7. water bath; 8. storage tank; 9. temperature controller; 10. motor; 11. glass-electrode; 12. reference electrode; 13. thermocompensator; 14. axial-flow impeller; 15. magnetic motor; 16. flowmeter; 17. fluidized-bed crystallizer; 18. recycle valve; 19. distributor.

0.22- μm filter and then diluted to 50 mL with deionized water. The calcium-ion concentration of the diluted solution was then determined by an atomic absorption spectrometer and the growth rate of calcite crystal was estimated from the consumption rate of the calcium ion. A typical run lasted 260 min.

Determination of crystal growth rate

The total mass and surface area of crystals are given by

$$W = n\rho_p\alpha L^3 \quad (5)$$

$$A = n\beta L^2, \quad (6)$$

where ρ_p is the crystal density; α and β are the volume and surface area shape factor, respectively; and n the number of crystals.

The crystal growth rate expressed in $\text{kg}/\text{m}^2\text{s}$ is

$$R_g = \frac{1}{A} \frac{dW}{dt} \quad (7)$$

or

$$R_g = \frac{3\alpha\rho_p}{\beta} \frac{dL}{dt}. \quad (8)$$

Combine Eqs. 7 and 8, the linear growth rate G can be written as

$$G = \frac{dL}{dt} = \frac{\beta}{3\alpha\rho_p A} \frac{dW}{dt}. \quad (9)$$

During a growth experiment, the change of crystal mass in a

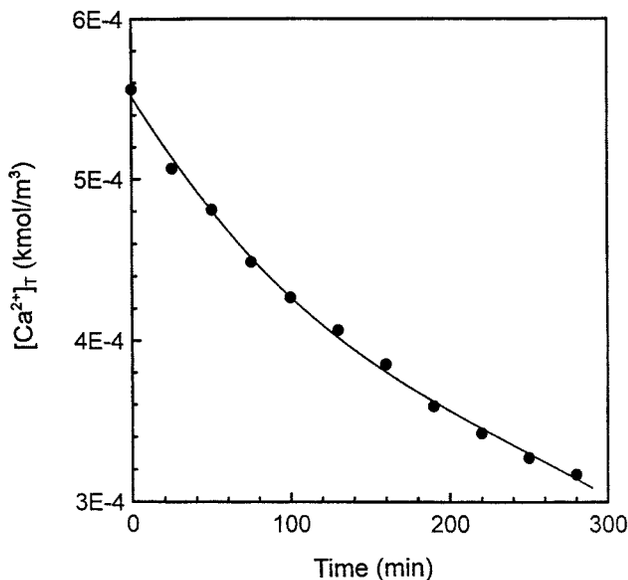


Figure 4. Typical change in the total calcium concentration over time for run O14.

crystallizer is related to the change of calcium-ion concentration as follows:

$$\frac{dW}{dt} = MV \left(- \frac{d[\text{Ca}^{2+}]}{dt} \right), \quad (10)$$

where M is the molecular weight of CaCO_3 , and V is the solution volume.

Equation 10 is substituted into Eq. 9 to give

$$G = \frac{\beta MV}{3\alpha\rho_p A} \left(- \frac{d[\text{Ca}^{2+}]}{dt} \right). \quad (11)$$

Combining Eqs. 5 and 6, we have

$$\frac{\beta}{\alpha A} = \frac{L\rho_p}{W}. \quad (12)$$

Substituting Eq. 12 into Eq. 11, we obtain the linear growth-rate expression:

$$G = \frac{LMV}{3W} \left(- \frac{d[\text{Ca}^{2+}]}{dt} \right). \quad (13)$$

Once the concentration profile of $[\text{Ca}^{2+}]$, which is determined by an atomic absorption spectrometer, is available, the linear crystal growth rate can be evaluated by assuming that the crystal size and total crystal weight are the same as for the seed crystal. This assumption will not cause much error because the total increase in crystal size or weight is negligible as compared with the seed crystal. For example, the growth rate of calcite is roughly 1×10^{-10} m/s, and thus an increase in crystal size for a growth time of 260 min is estimated to be 1.5 μm , which is negligible when compared with the seed sizes of 460 μm or larger. A typical calcium-ion concentration profile trend is shown in Figure 4.

Results and Discussion

Metastable region of CaCl_2 - Na_2CO_3 - H_2O system

Using the concentration data at the boundary shown in Table 1 for pH at 8.5 and other similar data for pH at 9.5 and 10.5 not shown here (Chen, 1985), $\mu[\text{Ca}]_T$ is plotted against $\mu[\text{CO}_3]_T$ in Figure 5 to mark the supersolubility curves, in which $\mu[\text{Ca}]_T$ and $\mu[\text{CO}_3]_T$ represent $-\log[\text{Ca}]_T$ and $-\log[\text{CO}_3]_T$, respectively. The solubility curves are also plotted in the figure using the solubility product of CaCO_3 . The metastable regions of the three pHs do not agree when they are projected to the same pH plane. However, if $[\text{Ca}^{2+}]$ and $[\text{CO}_3^{2-}]$ at the boundaries, which were calculated from the total concentration data using a computer program for estimating supersaturation, are used in the plotting (as shown in Figure 6), the metastable region is independent of the pH value, and the solubility and supersolubility curves are bent at both ends. Similar curves for the CaCl_2 - Na_3PO_4 - H_2O system have been reported (Füredi-Milhofer et al., 1975). This type of curve means that the formation of complex ions is significant (Söhnel and Garside, 1992). When the concentra-

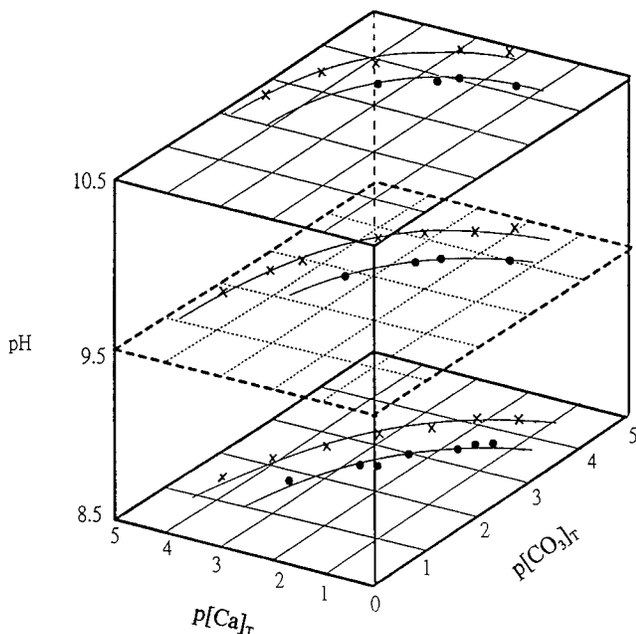


Figure 5. Metastable region of calcium carbonate for the $\text{CaCl}_2\text{-Na}_2\text{CO}_3$ system at pH 8.5, 9.5, and 10.5.
 —×— supersolubility curve; —·— solubility curve.

tion of Ca^{2+} and CO_3^{2-} , that is, $[\text{Ca}^{2+}]$ and $[\text{CO}_3^{2-}]$, in Figure 6 is replaced by the activity of the respective ion, $a_{\text{Ca}^{2+}}$ or $a_{\text{CO}_3^{2-}}$, the solubility and supersolubility curves become almost straight and parallel lines in the concentration range studied in this experiment, as shown in Figure 7. This simplifies the plotting of the metastable region.

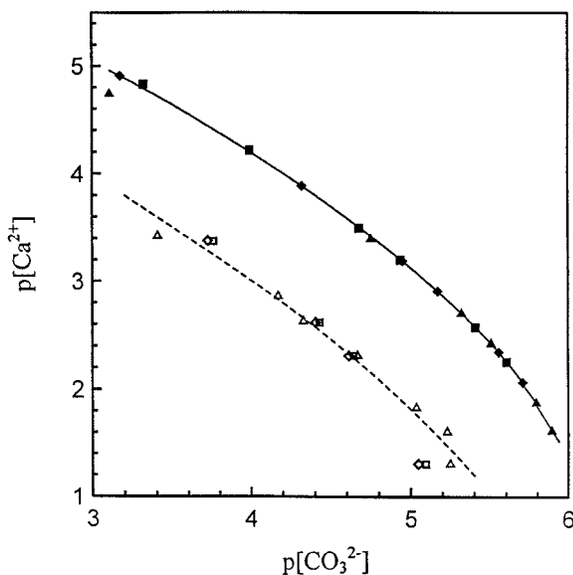


Figure 6. $p[\text{Ca}^{2+}]$ vs. $p[\text{CO}_3^{2-}]$: metastable region of calcium carbonate.
 — solubility curve: \blacktriangle pH = 8.5, \blacklozenge pH = 9.5, \blacksquare pH = 10.5; --- supersolubility curve: \triangle pH = 8.5, \diamond pH = 9.5, \square pH = 10.5.

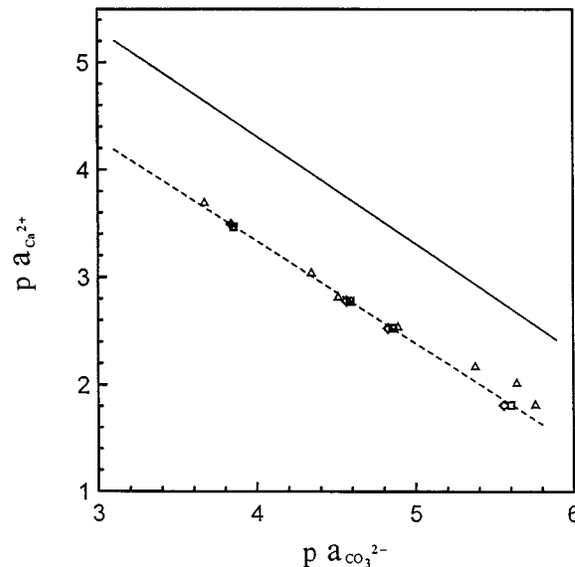


Figure 7. $pa_{\text{Ca}^{2+}}$ vs. $pa_{\text{CO}_3^{2-}}$: metastable region of calcium carbonate.

— solubility curve; --- supersolubility curve: \triangle pH = 8.5, \diamond pH = 9.5, \square pH = 10.5.

Factors affecting calcite growth rate in a dense fluidized bed

In the operation of a fluidized bed, the superficial velocity of the crystals should be lower than the terminal velocity, otherwise, the crystals would be carried over the top of the bed. At low superficial velocity, a boundary between solution and suspension is clearly observed in a dense bed. When the superficial velocity approaches the terminal velocity, particles move freely in the bed and the boundary no longer exists. In this case, it is called a lean fluidized bed. The dense bed is more advantageous as far as seed loading is concerned. Therefore, factors that affect the calcite growth rate in a dense bed are investigated, including supersaturation, pH, ionic strength, superficial velocity, particle size, and type of seeds.

Figure 8 shows the calcite growth rate as a function of supersaturation at various levels of ionic strength from 0.0025 to 0.0340 kmol/m^3 . For all levels of ionic strength, which is adjusted by adding NaCl solution, the growth rates increase with an increase in supersaturation. At the same supersaturation, the growth rates increase with ionic strength from 0.0025 kmol/m^3 to 0.0185 kmol/m^3 ; however, a further increase in ionic strength is no longer effective. Besides, the effect of ionic strength is less significant at higher supersaturations. One other thing worth noting is that the slope of the line, $\log G$ vs. $\log \sigma$, changes for the lowest ionic strength of 0.0025 kmol/m^3 , implying a change in the strength of various types of resistance to crystal growth. The effects of ionic strength have been reported for the crystal growth of vaterite and lead fluoride. Kralj et al. (1990) found a 10% increase in the growth rate of vaterite crystal when the ionic strength varied between 0.015 kmol/m^3 and 0.315 kmol/m^3 . As compared with the range studied in this experiment (0.0185 kmol/m^3 to 0.034 kmol/m^3), the effect of ionic strength was not observed. On the other hand, Stubicar et al. (1993) reported that the growth

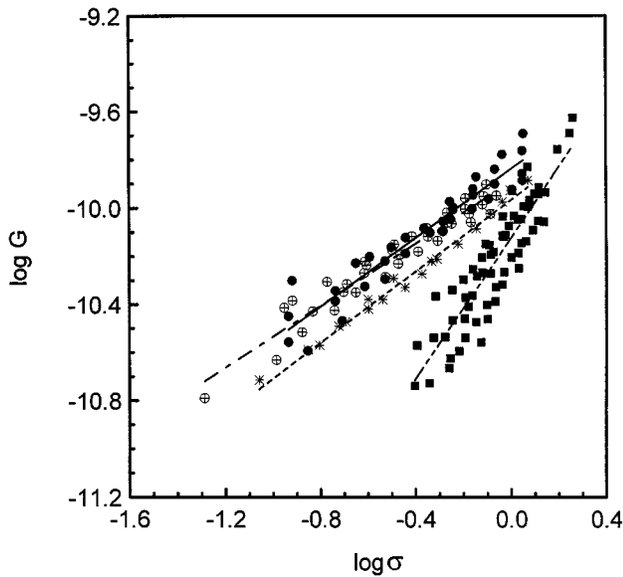


Figure 8. $\log G$ vs. $\log \sigma$ effect of ionic strength on the growth rate of CaCO_3 crystal at $L = 460 \mu\text{m}$, $\text{pH} = 9.5$, $T = 25^\circ\text{C}$.
 \circ $I = 0.0340 \text{ kmol/m}^3$; \bullet $I = 0.0185 \text{ kmol/m}^3$; $*$ $I = 0.0105 \text{ kmol/m}^3$; \blacksquare $I = 0.0025 \text{ kmol/m}^3$.

rate of PbF_2 is about five times greater at a high ionic strength of 0.1 kmol/m^3 as compared with that in water, and a different growth mechanism is speculated. All the findings on the effect of ionic strength are similar to the results of this experiment.

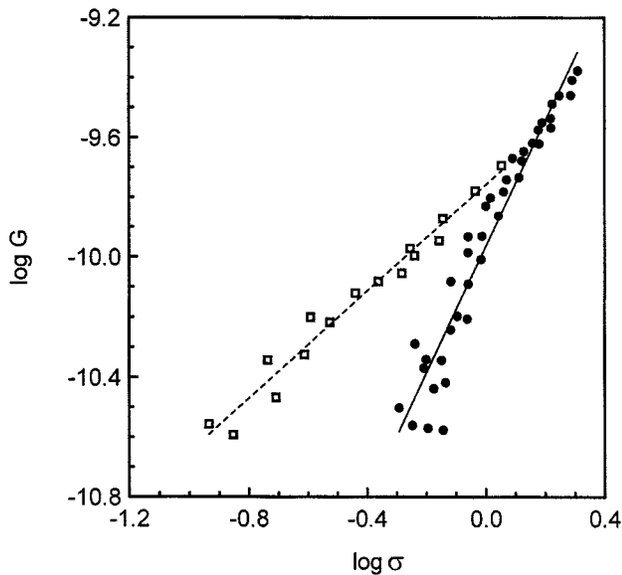


Figure 9. Crystal growth rates of calcium carbonate, $\log G$ vs. $\log \sigma$, at two pH values: $I = 0.018 \text{ kmol/m}^3$; $L = 460 \mu\text{m}$; $T = 25^\circ\text{C}$; $u = 2.36 \times 10^{-2} \text{ m/s}$.
 \bullet $\text{pH} = 8.5$; \square $\text{pH} = 9.5$.

The effects of pH on the growth rate of calcite are shown in Figure 9 for two levels of pH, namely 8.5 and 9.5, which is about the range used in a pellet reactor for water softening (van Dijk and Wilms, 1991). Similar to the effects of ionic strength, different slopes of the $\log G$ - $\log \sigma$ plot are obtained at different pH, and the effects of pH diminish at higher supersaturations. It is possible that the main resistance to crystal growth is influenced by ionic strength and pH in the same way. The effects of pH on the growth rate of lead fluoride were studied by Stubicar et al. (1993), using the constant-composition method. The result is consistent with that of calcite crystal studied in this experiment, that is, the crystal growth rate increases with pH when the pH is lower than the isoelectric point, which is 9 ~ 10 and 5.6 for calcite (Reed, 1989) and α -lead fluoride, respectively.

The growth rates of calcite were measured at various superficial velocities, that is, 1.42, 2.36, 3.54 and 4.72 m/s, for the seed crystals of size $460\text{-}\mu\text{m}$. Although the bed voidage and bed expansion changes a great deal, the calcite growth rates are rather constant, as shown in Figure 10. Assuming that the solution velocity does not influence the surface-reaction step, the constant growth rate is consistent with Tournie et al.'s (1979) conclusion that the mass-transfer coefficient is practically independent of the liquid velocity, which they arrived at by analyzing the extensive dissolution data for both lean and dense fluidized beds reported in the literature. On the other hand, Figure 11 shows the effects of particle size on the calcite growth rate at a superficial velocity of 4.72 m/s. The calcite growth rate increases with an increase in crystal size. The influence of superficial velocity and crystal size on the mass-transfer coefficient in a fluidized bed is discussed later.

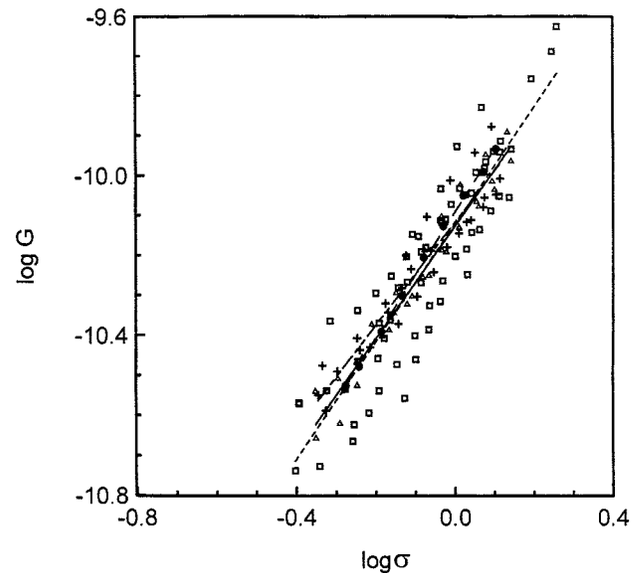


Figure 10. Growth rates of calcium carbonate, $\log G$ vs. $\log \sigma$, at various superficial velocities: $L = 460 \mu\text{m}$; $\text{pH} = 9.5$; $T = 25^\circ\text{C}$; $I = 0.0025 \text{ kmol/m}^3$.
 \triangle $u = 1.42 \text{ m/s}$; \square $u = 2.36 \text{ m/s}$; $+$ $u = 3.54 \text{ m/s}$; \bullet $u = 4.72 \text{ m/s}$.

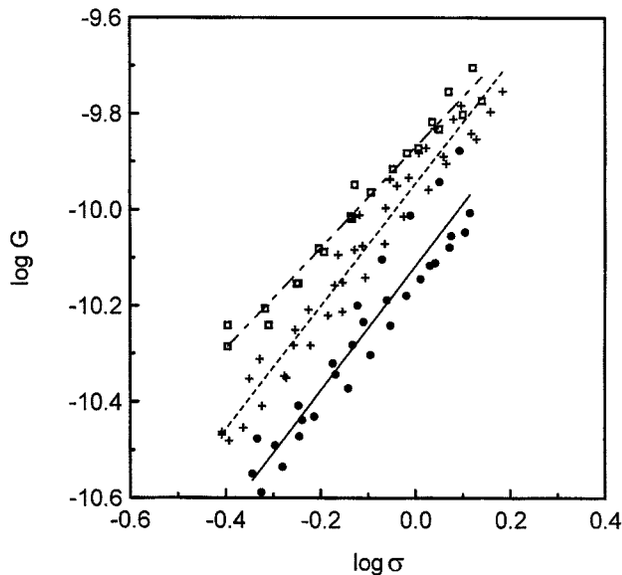


Figure 11. $\log G$ vs. $\log \sigma$ effect of crystal size at $\text{pH} = 9.5$; $I = 0.0025 \text{ kmol/m}^3$; $u = 4.72 \text{ m/s}$; $T = 25^\circ\text{C}$.

□ $L = 920 \mu\text{m}$; + $L = 650 \mu\text{m}$; ● $L = 460 \mu\text{m}$.

In the operation of a water-softening pellet reactor, silica sand or quartz sand is used as the seeding material (van Dijk and Wilms, 1991). It is therefore desirable to investigate the growth phenomena of calcite on the surface of silica sand. The concentration profile of the calcium ion is shown in Figure 12 for an experimental run using silica sand as seed. When compared with Figure 4, it is clearly seen that an induction time of calcite growth is required for silica sand. The induc-

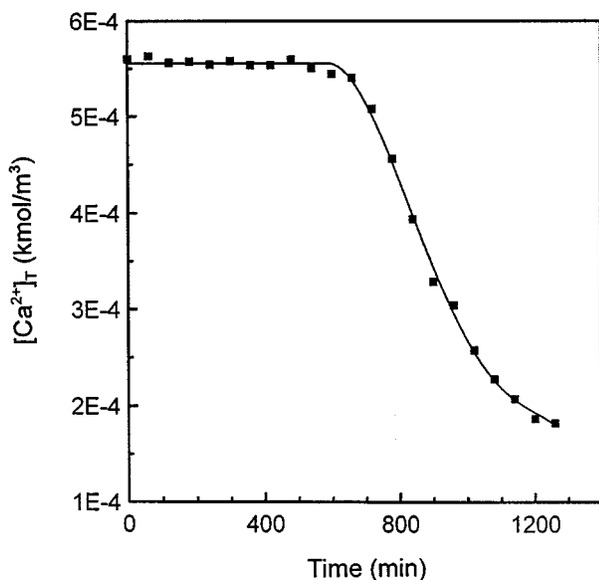


Figure 12. Induction time for calcium carbonate growth using silica sand as seed under the following operation conditions: $\text{pH} = 9.5$; $u = 2.36 \text{ m/s}$; $\sigma = 1.3$; $L = 460 \mu\text{m}$.

tion time is 600 min for the specific operating conditions, and is shorter at higher supersaturation. The calcite growth rate on the silica sand seed is lower than that on the calcite seed, as shown in Figure 13, in which curves A and B are the growth rates of sand seed after a growth period of 15 h and 20 h, respectively. The growth rate of sand seed will eventually approach the growth rate of calcite seed when the surface of sand seed is fully covered with calcite.

Mass-transfer coefficient of calcite growth in a dense fluidized bed

According to two different sources summarized by Tai et al. (1993), the surface reaction order of calcite growth is approximately 2. This is also true for some other systems, such as copper sulfate pentahydrate (Tai and Pan, 1985) and potassium alum (Tai et al., 1987). Taking $r = 2$, Eqs. 2 and 3 are combined to give

$$\frac{\sigma}{\sqrt{G}} = \frac{1}{K_d} \sqrt{G} + \frac{1}{\sqrt{K_r}} \quad (14)$$

Thus the mass-transfer coefficient, K_d , can be evaluated from the slope of the plot, (σ/\sqrt{G}) vs. \sqrt{G} . Figure 14 illustrates the plotting of Eq. 14 for three different crystal sizes at the same pH and ionic strength. The obtained K_d are listed in Table 2. Using the table, we can compare K_d for the different crystal sizes suspended at a superficial velocity of 3.54 m/s during the first three runs. On the other hand, the K_d values of the 460- μm -sized crystals operated at various superficial velocities can be compared using the last three rows in the table. Despite the variation in particle size and superficial velocity, the K_d are rather constant. The deviations from the average value of K_d , 1.32×10^{-10} , are less than 10%.

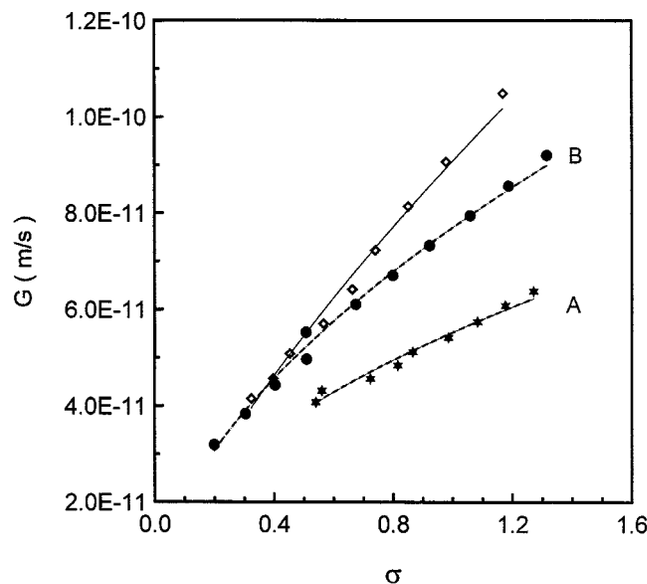


Figure 13. Comparison of growth rates of different seed types.

◇ calcite; ● silica sand after 20 h of growth time; ★ silica sand after 15 h of growth time.

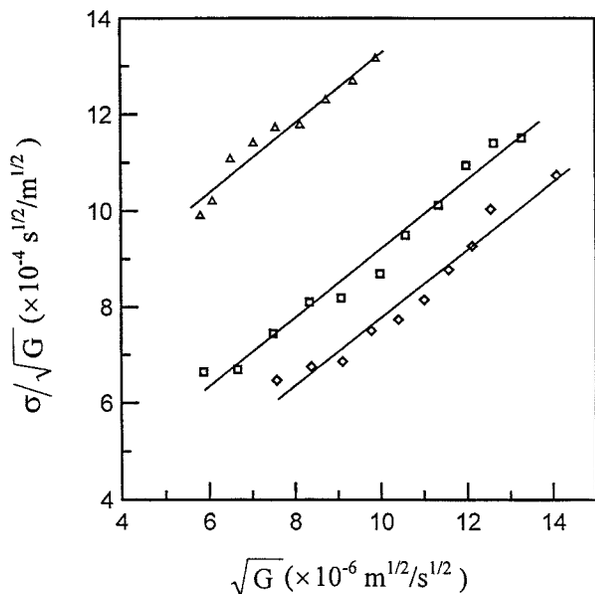


Figure 14. Equation 14 for calcium carbonate system at pH = 9.5 and $I = 0.0025 \text{ kmol/m}^3$.

Run No.	Crystal Size (μm)
Δ Run A-2-2-M14	460
\square Run A-4-2-O13	650
\circ Run A-4-1-O20	920

The mass transfer between fluidized spheres and liquid solution has been widely studied in many operations, such as dissolution, crystallization, ion exchange, leaching, and adsorption. Numerous correlations to estimate the mass-transfer coefficient are proposed in the literature and summarized by Tournie et al. (1997). However, none of them is derived from or applied to the crystal growth of a barely soluble system. One of the correlations originally proposed by Fan and modified by Kunii and Levenspiel (1969) for a liquid fluidized bed with low voidage has the following form when neglecting the diffusion term:

Table 2. Mass-Transfer and Surface-Reaction Coefficients of Calcite Crystal Estimated by Using Eq. 14 at pH = 9.5 and $I = 0.0025 \text{ kmol/m}^3$

Run No.	Particle Size (μm)	Superficial Velocity (m/s)	K_d (10^{-10} m/s)	K_r (10^{-9} m/s)
A-4-1-O20	920	3.54	1.45	13.78
A-4-2-O13	650	3.54	1.42	2.25
A-2-2-M14	460	3.54	1.33	0.30
A-1-4-M08,16	460	2.36	1.11	0.31
A-2-4-M12,13	460	1.42	1.29	0.34

$$Sh\left(\frac{K_d L}{D}\right) = 1.5[(1 - \epsilon) Re_p]^{1/2} Sc^{1/3}, \quad \epsilon \leq 0.84 \quad (15)$$

and

$$Re_p = \frac{Lu\rho}{\mu}, \quad 5 < Re_p < 120, \quad (16)$$

where u is the superficial velocity and ϵ the bed voidage. Equation 15 is derived by using the experimental data of β -naphthol and benzoic acid, which are soluble systems. To test the applicability of Eq. 15 to a barely soluble salt, calcite, the diffusivity of calcite in water should be found first. Unfortunately, the diffusivity data of calcite are not available in the literature, and the existing correlations for predicting diffusivity are only good for dilute solutions, which is quite different from the supersaturated solution we are dealing with.

Although a correlation to predict the mass-transfer coefficient could not be established for the calcite growth, Eq. 15 is used to test the consistency of the mass-transfer coefficient as shown in Table 2. Values of $[(1 - \epsilon) Re_p]^{1/2}$ for crystals of 460, 650 and 920 μm , fluidized at various superficial velocities, are tabulated in Table 3. The values of $[(1 - \epsilon) Re_p]^{1/2}$ at various superficial velocities are quite constant for each of the sizes. Therefore an average value is calculated for each size in Table 3. According to Eq. 15, the mass-transfer coefficient is proportional to $[(1 - \epsilon) Re_p]^{1/2}/L$. The ratios of $[(1 - \epsilon) Re_p]^{1/2}/L$ to that of 920 μm are shown in the last column

Table 3. Values of $[(1 - \epsilon) Re_p]^{1/2}$ for Different Sizes of Calcite Fluidized at Various Superficial Velocities

Particle Size, L (μm)	Superficial Velocity (m/s)	Bed Voidage ϵ	Re_p	$[(1 - \epsilon) Re_p]^{1/2}$	$[(1 - \epsilon) Re_p]^{1/2}$ Average	$[(1 - \epsilon) Re_p]^{1/2}/L$
						$\left\{ \frac{[(1 - \epsilon) Re_p]^{1/2}/L}{[(1 - \epsilon) Re_p]^{1/2}/L} \right\}_{L=920 \mu\text{m}}$
460	1.42	0.68	7.14	1.52	1.57	0.97
	2.36	0.77	12.05	1.67		
	3.54	0.84	18.08	1.67		
	4.72	0.91	24.11	1.44		
650	1.42	0.50	10.23	2.26	2.61	1.13
	2.36	0.58	17.06	2.67		
	2.81	0.64	20.46	2.71		
	3.54	0.71	25.55	2.72		
920	1.42	0.60	24.17	3.11	3.23	1
	2.81	0.64	29.00	3.22		
	3.54	0.70	36.11	3.29		
	4.72	0.77	47.20	3.31		

of Table 3. Thus, the ratios of the mass-transfer coefficient for the three sizes are:

$$\frac{(K_d)_{L=460 \mu\text{m}}}{0.97} = \frac{(K_d)_{L=650 \mu\text{m}}}{1.13} = \frac{(K_d)_{L=920 \mu\text{m}}}{1.0} \quad (17)$$

The ratios are rather constant, meaning that the mass-transfer coefficient of calcite growth in a fluidized bed is independent of crystal size and superficial velocity. This result is consistent with Table 2.

The surface-reaction coefficients at various crystal sizes are also listed in Table 2, which increases with crystal size and are rather constant for the same size. The size-dependent K_r is responsible for the effect of particle size on the calcite growth rate shown in Figure 11. Similar results have been reported for calcite growth in a stirred tank (Tai et al., 1993) and potassium alum in a fluidized bed (Budz et al., 1984; Tai et al., 1987). In Table 2 we also note that the values of K_r are higher than those of K_d , especially for larger sizes. It is interesting to see if there is a controlling step for the calcite growth, which can be judged from the surface-reaction effectiveness factor η (Garside, 1971) or the ratio of σ_{2-}/σ . If η or σ_{2-}/σ approaches 1, the crystal growth process is surface-reaction control. On the other hand, if η or σ_{2-}/σ approaches 0, the growth process is mass-transfer control. The calculated values of η and σ_{2-}/σ range between 0.191 and 0.355, and 0.433 and 0.577, respectively, for the experimental run A-2-2-M14, in which the mass-transfer resistance is the lowest among the experimental runs shown in Table 2 as judged from the relative magnitude of K_d to K_r . Therefore, the mass-transfer resistance and surface-reaction resistance are both significant, with the former being higher than the later in a fluidized bed operated at pH 9.5 and an ionic strength 0.0025 kmol/m³. For the largest size of 920 μm , the growth is almost mass-transfer controlled.

Growth-rate expression of calcite crystals

Once the surface-reaction order, mass-transfer coefficient, and surface-reaction coefficient are available, the crystal growth rate can be expressed according to the two-step model. For a surface-reaction order $r=2$, the expression of crystal growth rate is as follows (Mullin, 1993):

$$G = K_d \left[\left(1 + \frac{K_d}{2K_r\sigma} \right) - \sqrt{\left(1 + \frac{K_d}{2K_r\sigma} \right)^2 - 1} \right] \sigma \quad (18)$$

The values of K_d and K_r can be determined from the experiment using a lab-scale fluidized bed. Because K_d is almost a constant, which is independent of crystal size and superficial velocity, and K_r is related to crystal surface properties, which should be independent of hydrodynamics, the experimental data of K_d and K_r obtained in a lab-scale apparatus can be used for scale-up purposes.

The units of crystal growth rate, G , in Eq. 18 is in m/s. The linear growth rate can be easily converted to mass growth rate or molar growth rate, such as in Eq. 1, by using Eqs. 9 and 10. Equation 18, which is based on the two-step growth model, is more able than Eq. 1, which is an overall growth-rate

equation, to represent the crystal-growth-rate expression of calcite.

Conclusion

The metastable region of the $\text{CaCl}_2\text{-Na}_2\text{CO}_3\text{-H}_2\text{O}$ system is explored in this study. The solubility and supersolubility curves, which are the boundaries of the metastable region, are almost parallel straight lines in the concentration range studied, when the precipitation diagram is constructed by plotting $pa_{\text{Ca}^{2+}}$ against $pa_{\text{CO}_3^{2-}}$. The nucleation of calcite is suppressed when the growth experiments are conducted in the metastable region, using a fluidized-bed crystallizer.

Factors that affect the calcite growth rate in a dense fluidized bed are identified, including supersaturation, pH, ionic strength, and particle size and type of seeds. On the other hand, the effects of superficial velocity on growth rate are less significant. An induction period for crystal growth is required for the silica-sand seed, which is one of the seeding materials used in a large-scale water-softening pellet reactor. The growth rate of this type of seed is slower before the seed surface is fully covered with calcite.

When the calcite-growth-rate data of constant pH and ionic strength are analyzed by the two-step growth model, the mass-transfer coefficients so obtained are independent of crystal size and superficial velocity. On the other hand, the surface-reaction coefficient is size dependent. Since the mass-transfer and surface-reaction coefficients are rather insensitive to hydrodynamics in a fluidized bed, a growth-rate equation can be given for design purposes according to the two-step growth model, in which the mass-transfer coefficient and surface-reaction coefficient are determined using a lab-scale fluidized bed.

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Notation

- [Ca²⁺] = calcium-ion concentration, kmol/m³
- [Ca]_T = total calcium concentration, kmol/m³
- [CO₃²⁻] = carbonate-ion concentration, kmol/m³
- [CO₃]_T = total carbonate concentration, kmol/m³
- D = diffusivity, m²/s
- G = linear crystal growth rate, m/s
- I = ionic strength, kmol/m³
- K_r = surface-reaction coefficient, m/s
- L = crystal size, m
- Re_p = particle Reynolds number
- Sc = Schmidt number
- Sh = Sherwood number
- T = temperature, °C
- t = time, s
- μ = solution viscosity, kg/ms
- ρ = solution density, kg/m³
- σ_{2-} = interfacial relative supersaturation

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