Calcium Fluoride Precipitation and Deposition From 12 mmol/L Fluoride Solutions With Different Calcium Addition Rates

1. Introduction

The bioavailable fluoride (F) deposits in the oral environment are primarily in the form of calcium fluoride (CaF$_2$) or CaF$_2$-like precipitates [1-5]. These CaF$_2$ deposits in the dental plaque become an oral F reservoir [6-8], which gradually releases F ions into saliva [9] that tends to maintain a constant elevated level of F. Persistent elevated F in oral fluids has been shown to have a profound effect in caries protection [10-12].

The CaF$_2$ particles are the major reaction products after a topical fluoride application to tooth surfaces [1-3]. The source of calcium (Ca) necessary for CaF$_2$ precipitation after topical F application could be enamel, saliva, plaque fluid, oral mucosa or calculus [13].
It was suggested that CaF$_2$ was formed not only on enamel surfaces but also to some extent in the subsurface areas [14,15].

The CaF$_2$ particles were also formed during application of two-component rinses that contain fluoride (Na$_2$SiF$_6$) and calcium (CaCl$_2$) sources [16]. Na$_2$SiF$_6$ instead of NaF was used as the F source because the hydrolysis of SiF$_6^{2-}$ produces a continuous release of F$^-$ into the solution. The Na$_2$SiF$_6$ type of two-component rinses, containing either a conventional (12 mmol/L F) or reduced (6 mmol/L F) level of F, have been shown to be much more effective than a one-component NaF-based conventional rinse containing 12 mmol/L F in: (a) in vitro lesion remineralization [17,18], (b) increasing oral bioavailable F [5,19-21], and most importantly (c) in situ lesion remineralization [22,23]. The greater effectiveness of these two-component F-release rinses is due to two factors. Firstly, the two-component rinses provide a high Ca concentration that induces a large amount of CaF$_2$ deposits, whereas with a conventional one-component 12 mmol/L NaF rinse only small amounts of CaF$_2$ deposits are formed due to the limited amount of oral Ca available [24]. Secondly, the continuous release of F$^-$ from SiF$_6^{2-}$ controls the rate of formation and deposition of CaF$_2$ and is the key to high CaF$_2$ deposition during the rinse application [16].

Our current goal is to develop a new type of two-component system with fixed NaF concentration and continuous Ca release to produce an enhanced bioavailable CaF$_2$ deposition. This novel two-component system with continuous Ca release generated through the dissolution of a suitable Ca salt could have several advantages. Firstly, since it contains NaF (instead of Na$_2$SiF$_6$) and appropriate Ca salts, which are generally recognized as safe (GRAS) chemicals, it faces fewer regulatory hurdles. Secondly, because the Ca-release rate can be controlled and adjusted by choosing the type, particle size range, and amount of the dissolving Ca salt, a large range of release rates is available, which is needed to counteract the effects of mouthrinse ingredients on CaF$_2$ precipitation. The initial step in development of this type of mouthrinse or dentifrice is determination of the Ca-release rate that produces an enhanced CaF$_2$ deposition.

In the present study, the CaF$_2$ precipitation and deposition were examined in the simple model system with 12 mmol/L NaF solution to which CaCl$_2$ solution was continuously added at average rates of (5, 7.5, 10, 12.5, 15 or 20) mmol L$^{-1}$ min$^{-1}$ mimicking dissolution rates of moderately soluble Ca salts. These studies are particularly important for determination of conditions at which the most efficient deposition of CaF$_2$ in/on porous model substrate could occur during short-term (1 min) application of Ca-release rinse in which Ca could be continuously released into a NaF solution by dissolution of a solid Ca salt.

2. Materials and Methods

2.1 Systems With Continuous Calcium Addition

Standard solutions with concentration of 0.1000 mol/L $\pm$ 0.0005 mol/L of NaF and CaCl$_2$ (Orion Research, Beverly, MA, USA) were used to prepare reactant solutions. The volume of 150 mL of 12 mmol/L NaF solution, with 30 mmol/L KCl as a background electrolyte was placed in a jacketed, 250 mL capacity glass vessel exposed to air and connected to a circulating bath set at 37 ºC. A vigorous stirring at the rate of about 1880 rad/min (300 rotations per minute) was employed using a magnetic stirrer. The 0.1 mol/L CaCl$_2$ titrant solution containing 30 mmol/L KCl as background electrolyte was continuously added at a constant rate (in mL/min) with an automatic burette (Dosimat 665, Brinkman Instruments, Westbury, NY, USA). Concomitantly, a 24 mmol/L NaF titrant solution containing 30 mmol/L KCl was added at the same titration rate as the CaCl$_2$ titrant solution to compensate for decrease in total F concentration due to continuous increase of reactant solution volume. The titration rate of these two solutions was adjusted to produce the desired average rate of total Ca increase of (5, 7.5, 10, 12.5, 15, or 20) mmol L$^{-1}$ min$^{-1}$ to the reactant solution.

The desired average rate of total Ca increase expressed in mmol L$^{-1}$ min$^{-1}$ is defined as the amount of Ca (in mmol) that was added in the total volume of reactant solution (in L) after 1-min titration. The added titrants cause the continuous increase in the total volume of suspension, and therefore the real rate of total Ca increase is not constant and slightly differs from the average rate of total Ca increase rate during the 1-min time interval. As a result, the actual total Ca concentration in the reactant solution does not increase linearly with time (Fig. 1, data points) and is slightly higher than the hypothetical total Ca concentration that increases with a constant rate (Fig. 1, solid line).

1 Certain commercial materials and equipment are identified in this paper to specify the experimental procedure. In no instance does such identification imply recommendation or endorsement by the National Institute of Standards and Technology, the National Institutes of Health, or the ADA Foundation or that the material or equipment identified is necessarily the best available for the purpose.
dependence to the logarithmic values of $F^-$ and $Ca^{2+}$. The activity coefficients (and corresponding relative charges of ions in solution)

The activity coefficients of $F^-$ and $Ca^{2+}$ in reactant solutions were converted into activities, respectively. The measured activities of $F^-$ or $Ca^{2+}$ in each standard solution were calculated from the equation $a(F^-)$ or $a(Ca^{2+})$ in standard solutions were calculated with Chemist©. The degrees of saturation ($S$) at different times were also calculated with Chemist© using equation $S = (IAP/K_{sp})^{1/3}$, where $IAP$ is ionic activity product, $IAP = a(Ca^{2+}) \cdot a^2(F^-)$, and $K_{sp}$ is solubility product of $CaF_2$ ($pK_{sp} = 10.41$).

The $H^+$ or OH$^-$ ions were not consumed or produced during CaF$_2$ precipitation. Therefore, it was not necessary to have a pH buffer in the solutions.

Turbidity of the suspension was measured as absorbance by a fiber optic spectrophotometer containing a photodetector amplifier (PDA1, World Precision Instruments (WPI), Sarasota, FL, USA), a visible light detector (VISD, WPI), and a miniature dipping probe (DIP-UV, WPI). Standard uncertainty of turbidity was assumed to be ± 0.005 absorbance unit based on observed standard deviations for four runs.

2.2 Fluoride Deposition in Model Substrate

Hydrophilic cellulose-ester filter discs with pore sizes of up to 0.2 µm and diameter of 8.0 mm (Bioanalytical Systems, West Lafayette, IN, USA) were used as the porous model substrates [26]. Filter discs (five in each run) were pre-soaked for 1 min in 30 mmol/L KCl solution and then placed in the reactant solution for the time interval from 0 s to 60 s to test 1-min F deposition. After deposition, the discs were rinsed twice in 50 mL of a stirred solution saturated with respect to CaF$_2$ for 5 s. The deposit from each filter disc was extracted by immersing in 1 mL of a stirred solution saturated with respect to CaF$_2$ for 30 min. This solution was then neutralized with 1 mL of Total Ionic Strength Adjuster Background (TISAB II, Orion Research) solution that also contained 0.5 mol/L NaOH, and F concentration was measured by F-electrode (Orion, Mo 94-09). The amount of extracted F from a filter disc was expressed as mass of F per unit surface area of a disc (in µg/cm$^2$). The amount of extracted Ca in some selected samples was determined spectrophotometrically as an arsenazo complex using a microanalytical procedure [27]. Particle sizes and morphology of deposited precipitate were examined by JEOL scanning electron microscope (JEOL JSM-5300, JEOL USA, USA).
Peabody, MA, USA). Before SEM examination, the filter discs were rinsed for 5 s in solution saturated with respect to and then for 5 s in ethanol.

The 1-min F depositions were determined in 12 mmol/L NaF solutions to which 0.1 mol/L calcium chloride solution was continuously added at average rates of (5, 7.5, 10, 12.5, 15 or 20) mmol L$^{-1}$ min$^{-1}$. The 1-min F deposition was also determined in three different control systems: (I) a two-component F-release rinse that after mixing of equal volumes (75 mL) of components contained 2 mmol/L Na$_2$SiF$_6$ (12 mmol/L F) and 10 mmol/L CaCl$_2$ and 50 mmol/L of acetate buffer at pH = 5, (II) a two-component rinse that after mixing of equal volumes (75 mL) of components contained 12 mmol/L NaF, 10 mmol/L CaCl$_2$, and 30 mmol/L KCl, and (III) a conventional one-component 12 mmol/L NaF rinse containing 30 mmol/L KCl.

The data for F depositions in filter discs ($n=10$) were expressed as mean value ± standard deviation (SD) for each Ca-addition system and control systems. The differences in the mean values (F depositions) among the groups (Ca-release systems and control systems) were tested with one way ANOVA followed by an all pair wise multiple comparison procedure (Student Newman-Keuls method) using the SigmaStat software.

3. Results

3.1 CaF$_2$ Precipitation

The changes of $c$(F$^-$) in reactant solutions during 1-min Ca addition at the rates of (5 to 20) mmol L$^{-1}$ min$^{-1}$ ($n=2$ for each system) are shown in Fig. 2a. In all systems $c$(F$^-$) continuously decreased with time indicating continuous CaF$_2$ precipitation.

The $c$(F$^-$) decreases correlated to Ca-addition rates. The increases in $c$(Ca$^{2+}$) in all systems (not shown) were very low at the beginning of reaction whereas in the later times, the increases became proportional to the rates of Ca addition. The pHs of suspensions were 5.8 ± 0.2 and they did not change with time.

Turbidities were very low at the very beginning of the reactions (induction times) indicating the presence of no or very small amounts of CaF$_2$ crystals (Fig. 2b). Induction times were from 0 s to 13 s for the systems with Ca-addition rates from (20 to 5) mmol L$^{-1}$ min$^{-1}$, indicating an inverse correlation. After the initial induction period the turbidities continuously increased with time in each of the systems in correlation with increasing Ca-addition rates.

![Fig. 2.](image)

Fig. 2. (a) Concentration of fluoride, $c$(F$^-$), and (b) absorbance as a function of time for precipitation systems with initial $c$(NaF) = 12 mmol/L, $c$(KCl) = 30 mmol/L, and different Ca addition rates (expressed in mmol L$^{-1}$ min$^{-1}$) of 5 (□), 7.5 (■), 10 (△), 12.5 (▲), 15 (○), and 20 (●).
The amounts of the precipitated CaF$_2$ calculated from measured $c(F^-)$ in experimental Ca-addition systems showed positive correlation (0.99) with corresponding absorbance values indicating the reliability of these short-term kinetics data.

From the amounts of precipitated CaF$_2$, the precipitation rates ($R$) expressed as $R=d c(CaF_2)/dt$ were calculated for all Ca-addition systems. The $R$ vs. time curves for selected systems are shown in Fig. 3a.

For Ca-addition rates of (5, 7.5, 10, 12.5, 15, and 20) mmol L$^{-1}$ min$^{-1}$ the maximum precipitation rates ($R_{max}$) of (0.10 ± 0.01, 0.13 ± 0.01, 0.16 ± 0.02, 0.20 ± 0.03, 0.22 ± 0.01, and 0.31 ± 0.02) mmol L$^{-1}$ s$^{-1}$ were reached at the times of (33 ± 2, 29 ± 2, 23 ± 2, 18 ± 2, 11 ± 2, and 6 ± 1) s. The $R_{max}$ showed positive correlation (0.99) with corresponding Ca-addition rates and negative correlation (0.95) with corresponding times.

The calculated degrees of saturation ($S$) are shown in Fig. 3b. For Ca-addition rates of (5, 7.5, 10, 12.5, 15, and 20) mmol L$^{-1}$ min$^{-1}$, the maximum $S$ were 9.2 ± 0.3, 9.6 ± 0.2, 9.8 ± 0.4, 10.3 ± 0.3, 8.8 ± 0.2, and 8.3 ± 0.3, respectively. All $R$ vs. time and $S$ vs. time curves have similar general trends increasing up to a maximum value and then continuously decreased. In each system the maximum $S$ was reached at earlier time than the corresponding maximum $R$.

3.2 CaF$_2$ Deposition

For the Ca-addition rates of (5, 7.5, 10, 12.5, 15, and 20) mmol L$^{-1}$ min$^{-1}$, the corresponding 1-min F depositions (mean ± SD, $n = 10$) were (3.78 ± 0.31) μg/cm$^2$, (11.45 ± 0.89) μg/cm$^2$, (9.31 ± 0.68) μg/cm$^2$, (8.20 ± 0.56) μg/cm$^2$, (6.63 ± 0.43) μg/cm$^2$, and (2.09 ± 0.28) μg/cm$^2$, respectively (Fig. 4).

The differences in the F deposition mean values among the systems with different Ca-addition rates were statistically significant (ANOVA, $p < 0.05$). The Ca/F molar ratio of 0.508 ± 0.013 ($n = 12$, two 1-min deposits from each of six systems were analyzed) indicated precipitation of pure CaF$_2$ in all systems. The SEM of the cubical CaF$_2$ crystals deposited in/on the model substrate for the system with a Ca-addition rate of 7.5 mmol L$^{-1}$ min$^{-1}$ is shown in Fig. 5.

The 1-min F depositions determined for three control systems (I, II, and III) with the same initial F concentrations (12 mmol/L F) were used for comparison with F deposition from Ca-addition systems. F depositions (mean ± SD, $n = 10$) from control systems (I)
Fig. 4. 1-min F depositions (mean ± SD, n = 10) for Ca-addition system that contained 12 mmol/L NaF solutions with different Ca-addition rates from (5 to 20) mmol L⁻¹ min⁻¹ and for three control systems: (I) a two-component F release rinse that contained 2 mmol/L Na₂SiF₆ (12 mmol/L F), 10 mmol/L CaCl₂, and 50 mmol/L of acetate buffer at pH = 5, (II) a two component rinse that contained 12 mmol/L NaF, 10 mmol/L CaCl₂, and 30 mmol/L KCl, and (III) a conventional one component 12 mmol/L NaF rinse containing 30 mmol/L KCl.

Fig. 5. Scanning electron micrographs of CaF₂ crystals deposited in/on model porous substrate (0.2 μm filter paper) from the solution with c(NaF) = 12 mmol/L, c(KCl) = 30 mmol/L, and Ca-addition rate of 7.5 mmol L⁻¹ min⁻¹ at the reaction time of 1 min.
two-component F-release rinse (2 mmol/L Na₂SiF₆ and 10 mmol/L CaCl₂), (II) two-component rinse with immediate mixing of F and Ca components (12 mmol/L NaF and 10 mmol/L CaCl₂), and (III) conventional one-component F rinse (12 mmol/L NaF), were (6.79 ± 0.40) μg/cm², (0.40 ± 0.21) μg/cm², and (0.11 ± 0.02) μg/cm², respectively (Fig. 4). The F deposition mean values for control systems I, II, and III were significantly higher (ANOVA, p < 0.05) than F deposition mean values for control systems I, II, and III.

The data on c(F⁻) changes with time in the reactant solutions indicated that the rates of CaF₂ precipitation (R) are controlled by the rates of Ca addition. This result is supported by the turbidity data, where an increase in turbidity correlated to the formation of CaF₂ particles.

All R vs. time curves have similar general trends increasing up to a maximum rate of precipitation value and then continuously decreased with time (Fig. 3a). All calculated logR vs. time and logR vs. –logS curves (not shown) have similar trends as the R vs. time curves showing the continuous increase initially, and the continuous decrease in the later times. The increase of logR with –logS curve up to the maximum value is typical for the time interval in which nucleation and crystal growth are concomitant processes and the further linear decrease of logR with –logS indicates that crystal growth was dominant in that time interval [28].

These precipitation kinetics data could be used for the explanation of the F deposition data. For the experimental systems with moderate Ca-addition rates of (7.5, 10, and 12.5) mmol L⁻¹ min⁻¹ that produced significantly larger F depositions, the maximum precipitation rates (Rmax) of (0.13, 0.17 and 0.20) mmol L⁻¹ s⁻¹ were reached in (29, 23, and 18) s, respectively. It indicated that the continuous nucleation and relatively slow crystal growth during the time intervals (0 to 29) s, (0 to 23) s, and (0 to 18) s for Ca-addition rates of (7.5, 10, and 12.5) mmol L⁻¹ min⁻¹, respectively, produced crystals that were small enough to penetrate in the small substrate pores. The majority of these single CaF₂ crystals that were initially formed or penetrated in/on the model substrate could be later fixed in surface pores by inter-growth and/or agglomeration (Fig. 5). For the experimental system with the highest Ca-addition rate of 20 mmol L⁻¹ min⁻¹ that produced very low F deposition, Rmax of 0.31 mmol L⁻¹ s⁻¹ was reached in 6s. It indicated that CaF₂ crystal growth almost immediately became dominant process in this system (Fig. 3a, decrease of R vs. time curve after 6s) and correspondingly, the CaF₂ crystals became too large for deposition in/on the porous model substrate. It is important to note that even lower F deposition was determined from the control system II with immediately mixed Ca and F components. It could be explained by an extremely fast growth and agglomeration of formed crystals that instantaneously became too large to penetrate in/on much smaller substrate pores. The still lower F deposition from the one-component conventional F rinse (control system III) could be explained by the absence of calcium in reactant solution and, therefore, the adsorption of F ions was the only possible F deposition mechanism.

The cubical shape of CaF₂ crystals (Fig. 5) formed in these Ca-addition systems is typical for precipitation from pure fluoride solutions. The CaF₂ crystals formed in fluorosilicate solutions (control system I) had spherical shape, similar as crystals formed in phosphate solutions [29,30].

The 1-min CaF₂ depositions in experimental systems do not positively correlate to the CaF₂ precipitation rates. The largest CaF₂ depositions were obtained from the Ca-addition systems in which precipitation rates moderately increased with time and reached maximum values after prolonged time intervals. It seems that continuous formation and penetration of submicron-size crystals during these time intervals were prerequisites for large 1-min CaF₂ depositions. These results indicate that in the F systems with continuous Ca addition, the rate of CaF₂ precipitation that controls CaF₂ deposition is critical factor for deposition effectiveness.

5. Conclusions

The rate of Ca addition controls rate of CaF₂ precipitation and deposition. The CaF₂ precipitation rates are proportional to the Ca-addition rates but the CaF₂ depositions are not proportional to the Ca-addition rates. The CaF₂ depositions from 12 mmol/L NaF solutions with continuous Ca-addition rates of 7.5 mmol L⁻¹ min⁻¹ to 12.5 mmol L⁻¹ min⁻¹ having maximum precipitation rates of 0.13 mmol L⁻¹ s⁻¹ to 0.20 mmol L⁻¹ s⁻¹, respectively, are significantly higher than depositions in systems without continuous Ca addition.
Acknowledgment

This investigation was supported, in part, by USPHS research Grant R01-DE5354 to the American Dental Association Foundation from the National Institutes of Health/National Institute of Dental and Craniofacial Research and is part of the dental research program conducted by the National Institute of Standards and Technology in cooperation with American Dental Association Foundation.

6. References

[1] B. Ögaard, G. Rölla, and K. Helgeland, Alkali Soluble and Alkali Insoluble Fluoride Retention in Demineralized Enamel in vivo, Scand. J. Dent. Res. 60, 200-204 (1983).

[2] G. Rölla, On the Role of Calcium Fluoride in the Cariostatic Mechanism of Fluoride, Acta Odontol. Scand. 46, 341-345 (1988).

[3] F. Lagerlöf, J. Ekstrand, and G. Rölla, Effect of Fluoride Addition on Ionized Calcium in Salivary Sediment and in Saliva, Scand. J. Dent. Res. 96, 399-404 (1988).

[4] G. Rölla and E. Saxegaard, Critical Evaluation of the Composition and Use of Topical Fluorides with Emphasis on the Role of Calcium Fluoride in Caries Inhibition, J. Dent. Res. 69 (Spec Iss), 780-785 (1990).

[5] G. L. Vogel, Z. Zhang, L. C. Chow, G. E. Schumacher, and D. Banting, Effect of in vitro Acidification on Plaque Fluid Composition with and without a NaF or a Controlled F Release Fluoride Rinse, J. Dent. Res. 79, 983-990 (2000).

[6] J. M. ten Cate, Review on Fluoride with Special Emphasis on Calcium Fluoride Mechanisms in Caries Prevention, Eur. J. Oral. Sci. 105, 461-465 (1997).

[7] J. M. ten Cate, Current Concepts on the Theories of the Mechanism of Action of Fluoride, Acta Odontol. Scand. 57, 325-329 (1999).

[8] J. Ekstrand and A. Oliveby, Fluoride in the Oral Environment, Acta Odontol. Scand. 57, 330-333 (1999).

[9] K. Sjögren, How to Improve Oral Fluoride Retention?, Caries Res. 35 (Suppl 1), 14-17 (2001).

[10] J. M. ten Cate and P. P. Duijsters, Influence of Fluoride in Solution on Tooth Demineralization. I. Chemical Data, Caries Res. 17, 193-199 (1983).

[11] H. C. Margolis and E. C. Moreno, Physicochemical Perspectives on the Cariostatic Mechanisms of Systemic and Topical Fluorides, J. Dent. Res. 69 (Spec Iss), 606-613 (1990).

[12] J. D. B. Featherstone, The Science and Practice of Caries Prevention, J. American Dental Assoc. 131, 887-899 (2000).

[13] M. J. Larsen and A. Richards, The Influence of Saliva on the Formation of Calcium Fluoride-like Material on Human Dental Enamel, Caries Res. 35, 57-60 (2001).

[14] H. Duschner, H. Götz, and B. Ögaard, Fluoride Induced Precipitates on Enamel Surface and Subsurface Areas Visualized by Electron Microscopy and Confocal Laser Scanning Microscopy, Eur. J. Oral. Sci. 105, 466-472 (1997).

[15] B. Ögaard, CaF₂ Formation—Cariostatic Properties and Factors of Enhancing the Effect, Caries Res. 35 (Suppl 1), 40-44 (2001).

[16] L. C. Chow and S. Takagi, Deposition of Fluoride on Tooth Surfaces by a Two-solution Mouth Rinse in vitro, Caries Res. 25, 397-401 (1991).

[17] L. C. Chow, S. Takagi, and S. Shih, Effect of a Two-solution Fluoride Mouthrinse on Remineralization of Enamel Lesions in vitro, J. Dent. Res. 71, 443-447 (1992).

[18] S. Takagi, L. C. Chow, S. Shih, and B. A. Sieck, Effect of a Two-Solution Fluoride Mouth Rinse on Deposition of Loosely Bound Fluoride on Sound Root Tissue and Remineralization of Root Lesions in vitro, Caries Res. 31, 206-211 (1997).

[19] G. L. Vogel, Y. Mao, C. M. Carey, and L. C. Chow, In vivo Fluoride Concentrations Measured for Two Hours after a NaF or a New Two-Solution Rinse, J. Dent. Res. 7, 448-452 (1992).

[20] G. L. Vogel, Y. Mao, C. M. Carey, L. C. Chow, and S. Takagi, Increased Overnight Fluoride Concentrations in Saliva, Plaque, and Plaque Fluoride after a Novel Two-solution Rinse, J. Dent. Res. 76, 761-767 (1997).

[21] G. L. Vogel, Z. Zhang, L. C. Chow, and G. E. Schumacher, Effect of a Water Rinse on “Labile” Fluoride and other Ions in Plaque and Saliva before and Following Conventional and Experimental Fluoride Rinses, Caries Res. 35, 116-124 (2001).

[22] L. C. Chow, S. Takagi, C. M. Carey, and B. A. Sieck, Remineralization Effect of a Two Solution Fluoride Mouth Rinse: An in situ Study, J. Dent. Res. 79, 991-995 (2000).

[23] L. C. Chow, S. Takagi, S. Frukhtbeyn, B. Sieck, E. E. Parry, N. S. Liao, G. S. Schumacher, and M. Markovic, Remineralization Effect of a Low Concentration Fluoride Rinse in an Intraoral Model, Caries Res. 36, 136-141 (2002).

[24] E. Saxegaard and G. Rölla, Kinetics of Acquisition and Loss of Calcium Fluoride by Enamel in vivo, Caries Res. 23, 406-411 (1989).

[25] C. W. Davies, Ion Association, Bulterworths, Washington D. C., (1962) pp. 41.

[26] S. Takagi, H. Liao, and L. C. Chow, Effect of a Low-fluoride-content Two-Component Rinse on Fluoride Uptake and on Dental Re-mineralization of Enamel Lesions: An in vivo Study, Caries Res. 35, 223-228 (2001).

[27] G. L. Vogel, L. C. Chow, W. E. Brown, A Microanalytical Procedure for the Determination of Calcium, Phosphate, and Fluoride in Enamel Biopsy Samples, Caries Res. 17, 23-31 (1983).

[28] M. Markovic and H. Füredi-Milhofer, Precipitation of Calcium Oxalates from High Ionic Strength Solutions. Part 6. Kinetics of Precipitation from Solutions Supersaturated in Calcium Oxalates and Phosphates, J. Chem. Soc., Faraday Trans. 1 84, 1301-1310 (1988).

[29] J. Christoffersen, R. M. Christoffersen, W. Kibalczyc, and W. G. Perdock, Kinetics of Dissolution and Growth of Calcium Fluoride and Effects of Phosphate, Acta Odontol. Scand. 46, 325-336 (1988).

[30] J. Christoffersen, R. M. Christoffersen, J. Arends, and S. Leonardsen, Formation of Phosphate-containing Calcium Fluoride at the Expense of Enamel, Hydroxyapatite and Fluorapatite, Caries Res. 29, 223-230 (1995).
About the authors: Milenko Markovic, Shozo Takagi, and Laurence C. Chow are PhD chemists and Stanislav Frukhtbeyn is research assistant with the American Dental Association Foundation in the Paffenbarger Research Center at National Institute of Standards and Technology. The National Institute of Standards and Technology is an agency of the U.S. Department of Commerce.