The Retardation of Enamel Dissolution Rates by Adsorbed Long-Chain Ammonium Chlorides

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Dissolution rate studies were conducted with hydroxyapatite and enamel in the presence of adsorbed surfactants. In general, the ability of the surfactant to retard the dissolution rate was directly related to its ability to adsorb onto apatite. Cetylpyridinium chloride adsorbed poorly onto apatite, and its influence on the dissolution rate was marginal. The long-chain protonated amines were much more effective as rate retarding agents, sometimes of the order of 1,000- to 10,000-fold. These compounds were also found to adsorb much more strongly. A systematic dependence of the dissolution rate on chain length was found for these amines.

In vitro studies1-2 have demonstrated that certain organic compounds retard the dissolution of enamel. Although other investigators3-9 have suggested a relationship between the phenomenon of adsorption at the enamel-liquid interface and the reduction in enamel solubility, little definitive information has been presented to correlate adsorption isotherms and dissolution rates. Equilibrium considerations for adsorption concomitant with dissolution experiments have been omitted, and conclusions have been based on one short-time interval. Therefore, previous studies of the influence of organic surface adsorbing agents on enamel solubility or antisolubility have not differentiated between rates of dissolution and solubility of enamel.

This report describes the first step of a quantitative study of the environmental conditions which can alter the adsorption process and their relation to the reduction in enamel dissolution rates. These experiments were first conducted with apatite, which is known to be the main constituent of the inorganic part of enamel. The results of this phase of the research were then correlated with enamel dissolution rate experiments.

Materials and Methods

The primary amine hydrochloride surfactants, n-decyl-, n-dodecyl-, n-tetradecyl-, and n-hexadecyl ammonium chloride, were prepared by bubbling anhydrous hydrogen chloride gas through a prefiltered solution (ca 4%) of free amine† in benzene. The hydrochloride was recrystallized three times from benzene. Practical grade cetylpyridinium chloride‡ was recrystallized once from acetone. A commercial synthetic hydroxyapatite§ (Victor apatite) was dried overnight at 110°C and was used for the adsorption study. The synthetic apatite which was used for the dissolution experiments was prepared according to the method of the Tennessee Valley Authority|| and will be designated as TVA apatite. The chemical composition and specific surface areas are given in Table 1. The low surface area of the TVA apatite sample necessitated the use of two differently prepared hydroxyapatite samples during this investigation.

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† Aldrich Chemical Co., Milwaukee, Wis.
‡ Eastman Organic Chemicals, Rochester, NY.
§ Stauffer Chemical Co., Victor Division, New York, NY.
|| Division of Chemical Development, Wilson Dam, Ala.
face areas were determined by the B.E.T. gas adsorption technic. *

Adsorption experiments were carried out by adding 10 ml of known adsorbate concentrations to known weights of Victor apatite (100 mg or 500 mg) into pyrex culture tubes. The tubes were rotated at 40 rpm in a constant-temperature bath by means of an end-over-end tumbling apparatus. At adsorption equilibrium, samples were withdrawn through 20 gauge 6-inch needles attached to 10 ml syringes and filtered through Swinny hypodermic adapters which contained 0.22 micrometer (µm) filters.† The predetermined adsorption equilibrium time was approximately 30 minutes, but the tubes were allowed to rotate overnight. At high adsorbate concentrations, the tubes were centrifuged before the samples were withdrawn. The clear supernatant solution was analyzed for the equilibrium surfactant concentration. The dodecylammonium chloride concentration was determined according to the antagonistic titration method of Ino, Kondo, Meguro, and Yoda,10 using sodium lauryl sulfate as the titrant and rhodamine 6G as the adsorption indicator. The precision of the assay was ± 2%. An ultraviolet analysis was used for cetpyridinium chloride (CPC) at a wavelength of 259 mµ. The number of millimoles of adsorbed surfactant was calculated by multiplying the concentration change by the volume.

Dissolution experiments with TVA hydroxyapatite were performed on previously equilibrated (18 hour) adsorbed samples. A 100 mg/10 ml slurry containing apatite and surfactant was quantitatively transferred to a 250 ml volumetric flask fastened on a wrist-action shaker.‡ The temperature was maintained at 30 C. Two hundred milliliters of a buffer or buffer-surfactant solution (previously equilibrated at 30 C) were added and the shaker and a timer were simultaneously started. Surfactant was incorporated into the buffer to maintain the equilibrium adsorption concentration. Samples were withdrawn at various time intervals with 7-inch needles into 10 ml syringes and were filtered through 0.22 µm filters contained in Swinny hypodermic adapters. The total number of milligrams of hydroxyapatite dissolved at any time was calculated from the amount of phosphate present. Phosphate ion in solution was determined according to the method of Gee, Domingues, and Deitz.11 In order to verify that the apatite was dissolving stoichiometrically, calcium ion concentrations were selectively determined by means of an atomic absorption spectrophotometer.§ Since the acid slurry contained dissolved apatite, the amount present was calculated from a duplicate sample which was filtered. This amount was subtracted from the total amount going into solution. Therefore, graphical representations gave the net amount of hydroxyapatite dissolved as a function of time.

The enamel dissolution studies were performed on six selected teeth embedded in a wax matrix mount. The mount, in the shape of a doughnut, was secured to the bottom of a 180 ml jacketed beaker. Temperature was maintained constant at 30 C. One hundred milliliters of a buffer or buffer-surfactant solution were added, the solution was stirred at 1,800 rpm, and samples were withdrawn at the various time intervals. As was the case with the hydroxyapatite powder, the surfactant was preadsorbed (for 30 minutes in this instance) under the same experimental conditions, ie, pH, buffer strength, and surfactant concentration, as those for dissolution. After the adsorption time, the buffer-surfactant solution was decanted, and fresh buffer-surfactant solution was added. The dissolution of the enamel was then followed. After each experiment, the mount was washed with fresh buffer to remove any adsorbed surfactant. Figure 1 shows the effect, and therefore demonstrates the need for the preadsorption of the adsorbing agents.

* Perkin-Elmer Shell Sorptometer, Perkin-Elmer Corp., Norwalk, Conn.
† Millipore Filter Corp., Bedford, Mass.
‡ Burrell Corp., Pittsburgh, Pa.
§ Perkin Elmer, Model 303, Norwalk, Conn.

### TABLE 1
**Composition and Surface Areas of Hydroxyapatite Samples**

<table>
<thead>
<tr>
<th></th>
<th>TVA</th>
<th>Victor</th>
</tr>
</thead>
<tbody>
<tr>
<td>% Phosphate theoretical</td>
<td>56.7</td>
<td>56.7</td>
</tr>
<tr>
<td>% Phosphate experimental</td>
<td>56.9</td>
<td>54.4</td>
</tr>
<tr>
<td>% Calcium theoretical</td>
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<td>39.9</td>
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<tr>
<td>% Calcium experimental</td>
<td>38.4</td>
<td>35.4</td>
</tr>
<tr>
<td>Ca/Po4 Molar ratio theoretical</td>
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<tr>
<td>Ca/Po4 Molar ratio experimental</td>
<td>1.64–1.68</td>
<td>1.54</td>
</tr>
<tr>
<td>Specific surface area (M²/gm)</td>
<td>1.17</td>
<td>66.1</td>
</tr>
</tbody>
</table>
Fig 1.—Effect of preadsorption of tetradecylammonium chloride on enamel dissolution at pH 4.5 (0.05 M acetate buffer) at 30 C.*

* All determinations done at 30 C.

Fig 2.—Adsorption isotherms of dodecylammonium chloride onto Victor apatite in sodium chloride solutions showing the influence of ionic strength. Arrows signify approximate critical micelle concentrations.
Results

The results of adsorption studies with the surfactant dodecylammonium chloride (DAC) and CPC are presented in Figures 2-4. The S-shaped curves are in contrast to Langmuirian type of adsorption, where the curves are initially linear with no inflection. Experiments performed in pH 4.5 and 6.0, 0.05 M acetate buffer showed differences in the adsorption behavior of the surfactants investigated. Further studies, considering the ionic strengths of the solutions, showed that these differences could be explained by an ionic strength effect.

Figures 2 and 3 illustrate the effect of increasing ionic strength, \( \mu \), (by the addition of sodium chloride) on the adsorption of DAC onto Victor apatite in water and pH 4.5 (0.05 M) acetate buffer respectively. A decided effect was observed showing a distinct displacement of the curves to lower equilibrium surfactant concentrations. The maximum amount adsorbed remained the same. The adsorption of CPC (Fig 4), on the other hand, showed an apparent reversal of this ionic strength dependency. The curves indicate no displacement but showed a decrease in the maximum amount adsorbed. Sodium chloride (0.05 M) essentially prevented any significant adsorption. Figure 4 also shows that no measurable adsorption of CPC occurred in pH 4.5, 0.05 M acetate buffer.

Retardations of the dissolution rates of the TVA hydroxyapatite sample and enamel were observed with all members of the long-chain homologous series of the protonated, primary amines studied, decyl-, dodecyl-, tetradecyl-, and hexadecylammonium chlorides. CPC, although another cationic surface active agent, caused no significant re-
FIG 5, 5a—Dissolution profiles of preadsorbed TVA apatite in various dodecylammonium chloride solutions in pH 4.5, 0.05 M acetate buffer.
FIG 6. 6a—Dissolution profiles of preadsorbed TVA apatite in various tetradecylammonium chloride solutions in pH 4.5, 0.05 M acetate buffer.
Fig 7.—Dissolution profiles of preadsorbed enamel in various dodecylammonium chloride solutions in pH 4.5, 0.05 M acetate buffer.

Fig 8.—Dissolution profiles of preadsorbed enamel in various tetradecylammonium chloride solutions in pH 4.5, 0.05 M acetate buffer.
tardations with either apatite or enamel.

Figures 5 through 8 are representative of typical TVA apatite and enamel dissolution profiles in the presence of various surfactant concentrations in pH 4.5 and 0.05 M acetate buffer. In a control experiment, the exact experimental conditions, including preadsorption, were maintained except that no surfactant was present. In all cases, the apatite dissolved congruently. Although the molybdate reagent in the phosphate assay interacted with the higher surfactant concentrations, the assay was not significantly altered.

For means of comparison, the retardation factors for both the TVA apatite and the enamel rate runs were calculated by taking the ratios of the times necessary to attain the same amount of dissolution in the surfactant experiment as in the control run. The retardation factor was then averaged over a particular interval. Figure 9 shows the retardation factor as a function of the different surfactant concentrations. Results from the enamel study demonstrated a similar type of rate retardation dependence on surfactant concentrations.

The correlation of adsorption data with rate retardation data for DAC can be seen by referring to Figures 3 (Curve A) and 9 (Curve B). No rate retardations were observed until appreciable adsorption took place, and then the retardations were found to increase with increasing adsorption.

When the effect of CPC on the dissolution of apatite was explored at two concentrations (0.08 and 2.0 mM), one concentration above and one below the measured CMC of 0.12 mM, no significant rate reductions were observed. The estimated ionic strength for these runs was approximately 0.02. This was sufficiently high so that the adsorption of CPC was expected to be very low (Fig 4). Similar results were obtained in the enamel dissolution studies, where the presence of CPC at 1 or 10 mM concentration showed no retardation.

Dissolution experiments with TVA apatite were also performed in pH 5.0 and 0.05 M acetate buffer in selected surfactant solutions. Figure 10 is an example of apatite dissolution in the presence of tetradecylammonium chloride (TAC) at pH 5.0. At this pH, plots of the retardation factor versus surfactant concentration were displaced by about 35% to lower concentrations of TAC when compared to the retardation results in pH 4.5, 0.05 M acetate buffer. A similar shift of about 50% was observed when 0.1 M sodium chloride was added to the dissolution media (pH 4.5).

Discussion

Adsorption Behavior.—The S-shaped nature of the adsorption isotherms illustrates nonideal behavior when compared to the Langmuir model for adsorption. It suggests a cooperative mechanism for adsorption involving lateral interactions of the hydrocarbon portion of the molecules. Also, the possibility for more than one monomolecular layer cannot be completely disregarded in the case of DAC. Using the value of 66.1 M²/gm (Table 1) for the surface area of Victor hydroxyapatite, a complete monomolecular layer would require the adsorption of 0.54 millimoles of the dodecylammonium ion. This assumes an area of 20.5 A² per molecule which is the reported value for condensed films of long-chain amines. The experimental value for the maximum amount adsorbed from water was 0.64 millimoles/gm. Although this is in fair agreement for a monolayer, a somewhat larger area per molecule or a smaller effective apatite surface area (due to solvent adsorption or to the ability of a gas molecule to penetrate areas during gas adsorption that are inaccessible to the larger organic molecule during adsorption from solution) would suggest bilayer adsorption.

The CPC plateau value in water reached only 16% of the DAC plateau value. If CPC and DAC were oriented similarly at the solid surface, the difference in the plateau values could indicate that the pyridinium-head group does not allow close packing at the interface and therefore gives poorer adsorption. It is possible that the nature of the pyridinium group may not interfere with the interactions of the aliphatic tails in a micelle, but prevents good interactions at a planar surface.

The addition of sodium chloride to the adsorption media altered the adsorption process differently for DAC and CPC. Such behavior may be rationalized in terms of the effects of salt on the critical micelle concentration and the apatite adsorption affinity for the two surfactants.

Dissolution Rate Retardation.—In this discussion, the adsorbed surfactants at the
enamel-liquid or hydroxyapatite-liquid interface are considered to form a molecular film. Therefore, in order for a dissolved ion to reach the bulk solution, it must pass through the interfacial film and the diffusion layer regions. The results of this investigation suggest that the rate determining step is controlled by this film.

Figure 11 shows a plot of the logarithm of the surfactant concentration at equal retardations versus carbon chain length of the ammonium ions. This linear relationship corresponds to about a threefold increase in rate retardation per increase in carbon num-

**Figure 9.** Retardation profiles of TVA apatite as a function of surfactant concentrations in pH 4.5, 0.05 M acetate buffer.
ber, and it is analogous to Traube's rule.17 This relationship reflects the greater adsorbing tendency (binding strength) of the higher homologues due to the greater number of methylene groups. Ionic transport, therefore, may take place through an all-or-none process across the film, the probability of which depends on the monolayer state and composition. This single step process can be accounted for by the fluctuation desorption of a surfactant molecule at a site of dissolution. A similar mechanism has been used to explain the retardation of water evaporation rates by condensed monolayers.18 A similar example of the correlation between adsorption and rate retardation was found when dissolution and DAC adsorption experiments were conducted with and without 0.10 M sodium chloride at pH 4.5. The observed greater retardations in the presence of sodium chloride may be attributed to the greater surfactant adsorbing tendency at the higher ionic strength. The greater rate retardation in pH 5.0 buffer may also be attributed to the same tendency because of a somewhat greater ionic strength at the higher pH value.

It should be pointed out that the long time dissolution data (Fig 5a, 6a) in the presence of the long-chain ammonium chlorides do not rule out the possibility of a solubility change of hydroxyapatite. Such changes would, however, be of a much smaller order of magnitude than the kinetic effects of the surfactant. In view of the complex solubility behavior of apatite itself, it is not surprising that the adsorption of these surfactants may influence the apparent solubility. Studies are under way to demonstrate the self-consistent interrelationships of both the kinetic and the apparent thermodynamic changes brought about by these adsorbing agents.

Conclusions

The influence of two types of cationic surfactants on the dissolution rates of enamel and hydroxyapatite was investigated. Surfactant adsorption onto hydroxyapatite showed that environmental conditions can significantly alter the adsorption process and concomitantly affect dissolution rates. Ionic strength, surfactant type, and micelle formation all played important roles in the adsorption process. Under the experimental dissolution conditions (pH 4.5), the adsorption of CPC was low and did not modify the dissolution rate of hydroxyapatite or enamel. On the other hand, the adsorption of dodecylammonium chloride onto apatite was measured, and significant rate reductions were observed with it and with the three other members of the homologous series. The greater retardation efficiency
per increased chain length was found to obey Traube's rule. The greater rate reduction with equivalent concentrations of DAC in the presence of sodium chloride was attributed to its greater adsorbing tendency. Hydroxyapatite dissolution results were in good agreement with the results of enamel dissolution experiments, and it was expected that the same factors would be operating in both systems.

It was proposed that the fluctuation desorption of a surfactant molecule provides a reasonable explanation for the rate determining process in the retardation of dissolution rates by long-chain ammonium chlorides.

References

2. MANLY, R.S., and MANLY, K.F.: Influence