INTRODUCTION

Almost half century of research strongly supports that fluoride (F) exerts its effect on caries and on dental fluorosis by distinct mechanisms. Thus, while the effect on caries is local, based on maintenance of oral F concentration [reviewed by Cury and Tenuta (1)], the effect on fluorosis is systemic, based on blood F concentration (2), which is influenced by innumerable metabolic variables.

For instance, it is recognized that blood fluoride concentration is not controlled by homeostasis (3), and thus the pattern of F use and how it affects plasma F levels may be of importance (4). Furthermore, considering that enamel formation is a continuous process and that the clinically visualized defects are dependent on the thickness of enamel formed under exposure to a given F dose (5), it is expected that short peaks of plasma F concentrations due to an acute dose would have less importance than the continuous exposure to chronic lower F doses on fluorosis development (4,6). A previous survey in a city that lacks a well controlled system of community water fluoridation showed that fluctuations of water F concentration above the optimal concentration (0.7 ppm F) did not increase fluorosis prevalence (7). In fact, in the city where water F concentration oscillated above (higher than 0.8 ppm F) and below (lower than 0.6 ppm F) the optimum, the fluorosis risk was 8.3 times lower than in the another city with constant optimum water F concentration (7).

This epidemiological observation raised the question on how fluctuations in F exposure chronically would affect enamel fluorosis. Thus, it was hypothesized that the systemic effect of F on fluorosis would depend on both the mean effect of each oscillating F dose and the average time of exposure of ameloblasts to them during

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enamel maturation, and a first experimental study was conducted in rats to evaluate the effect of symmetrical oscillations in water F concentration on metabolic variables and fluorosis development.

MATERIAL AND METHODS

Experimental Design

This blind study was approved by the Animal Ethics Committee of the State University of Campinas (Protocol #1579-1). Twenty Wistar females rats aged 35 days were randomly divided into 5 groups of 4 animals each. Four groups received ad libitum water containing 0 (control), 12.5, 25 or 37.5 µg F/mL during 78 days, which is sufficient time to allow the full renewal of rat incisors (8). During the same 78 days, the 5th group (oscillating) received water containing 12.5 or 37.5 µg F/mL alternately every 72 h. The F solutions were prepared with NaF and the concentrations were checked. The rats had free access to pellet chow (Labina; Purina, Paulínia, SP, Brazil; total F concentration of 17 µg F/g). The amount of water ingested by each group was assessed daily and animals were weighed at the beginning and the end of the experiment.

On the 79th day, the rats were anesthetized and blood was collected by disruption of the brachial plexus using heparin as an anticoagulant. One femur, the mandible bone and the lower incisors of the animals were removed and coded for the analysis by a blinded examiner. At the time of sacrifice, animals of the oscillating group had been drinking water at 37.5 µg F/mL for the last 24 h.

Plasma Fluoride Determination

Plasma samples were obtained by centrifugation at 2,000 g for 10 min. F concentration in plasma was determined using an inverted F-specific electrode (Orion Research model 94-09; Orion Research Incorporated, Cambridge, MA, USA) and a micro-reference electrode, under microscope, after buffering with TISAB III (9).

Fluoride Determination in the Femur (Whole and Surface), Mandible (Alveolar Bone) and Incisors

The femur was used for F determination in the whole bone and bone surface (10). The femur was dried for 24 h at 60°C, sectioned transversally to obtain 2 slices measuring approximately 5 mm from the diaphysis region. For whole F determination, one slice was pulverized, the powder was sieved to obtain particles of 140 µm to 1,000 µm and left at 60°C for additional 24 h. Ten milligrams (±0.01 mg) were weighed and transferred to plastic test tubes containing 1.0 mL of 0.5 M HCl. After 1 h, at room temperature and under agitation, 1.0 mL of TISAB II (1.0 M acetate buffer, 1.0 M NaCl, 0.4% CDTA, pH 5.0) containing 0.5 M NaOH was added.

To determine F concentration on the bone surface, the other femur slice was sealed with acid-resistant nail varnish leaving a circular hole (2.0 mm diameter) exposed (10). Three layers of bone were sequentially removed by immersing each slice in tubes containing 0.25 mL of 0.5 M HCl for 15, 30 and 60 s, under constant agitation. The extracts were neutralized and buffered with equal volumes of TISAB II, as described before.

For F determination on the surface alveolar bone where there was an incisor tooth, one hemi-mandible from each animal was dried for 24 h at 60°C. The lingual plate of the alveolar socket was removed, thereby exposing the rest of the alveolar socket. All other regions were isolated with acid-resistant nail varnish. Using an automatic pipette, 0.1 mL of 0.5 M HCl was applied to the alveolar socket and the acid was flushed for 15 s by aspiration and injection. The extract was transferred to a test tube and buffered as described before.

F concentration in the mandible incisors was determined at 3 transversal slices measuring approximately 1 cm from the apical, middle and incisal regions. These fragments were separately pulverized and the powder was dried at 60°C for 24 h. Five milligrams (±0.01 mg) were weighed and transferred to plastic test tubes, to which 1.0 mL of 0.5 M HCl was added for F extraction as described for whole femur F determination.

F in all extracts was determined using an ion-specific electrode (Orion Research model 94-09; Orion Research Incorporated) and a reference electrode (Flex-Ref; WPI Inc., Sarasota, FL, USA) both coupled to a potentiometer (Orion Research model EA 940; Orion Research Incorporated), previously calibrated with standards with known F concentration, and prepared under the same conditions as the samples. All results were expressed as µg F/g dry weight and the amount of bone removed from the surface of femur or alveolar bone was calculated according to Bezerra de Menezes et al. (10).
Dental Fluorosis Assessment

Due to the limitations of hardness and transverse microradiography to quantify dental fluorosis in rats exposed to low F concentrations (4,11), our research group has developed a quantitative method of image analysis named Dental Fluorosis by Image Analysis (DFIA), based on the known effect of F changing the iron pigmentation of the rat incisor enamel surface, which was quantified.

In order to acquire the input images, one mandible incisor from each animal was fixed with wax on a glass slide (Fig. 1) with the enamel surface of incisal third of the crown upward parallel to a macro lens (Medical Nikkor; Nikon, Tokyo, Japan) attached to a vertical stand at a fixed distance of 7 cm above the teeth. An image was acquired with the digital camera (Nikon D70; Nikon) using standardized light intensity and focus. A central area of 690 x 170 pixels was cropped from the digital image and converted to gray-scale. The gray-scale cropped image is used as the input image for the DFIA method. The input image is segmented in order to isolate the darker bands from the lighter bands (Fig. 1). In Figure 1, the black area corresponds to the original orange bands and the white area corresponds to the original white bands. For each segmented area, the method computes the average of the pixel intensities in the input image. The DFIA index is obtained by the difference between \( I_{HI} \) (the average of the higher intensity pixel values, originally the white bands) and \( I_{LI} \) (the average of the lower intensity pixel values, originally the orange bands). Higher values of DFIA index should correspond to more severe fluorosis.

Statistical Analysis

Linear regression analysis was used to correlate the administered constant F concentrations in water (independent variable) with F concentration in plasma, bones and incisor, and dental fluorosis found (SPSS software version 12.0; SPSS Inc., Chicago, IL, USA). Using the linear equations, the estimated mean F concentrations to which the oscillating group (12.5/37.5 µg F/mL) would have been subjected during the all experiment was calculated. Differences among the groups were assessed by ANOVA and Tukey's post-hoc test using SAS software version 8.01 (SAS Institute Inc., Cary, NC, USA). Data that did not fit the assumptions of normal distribution of errors and equality of variances were transformed (12). All analyses were performed at 5% significance level.

RESULTS

No statistically significant difference (p>0.05) in water intake (daily) and rat weights (either initial or final) was observed among groups.

A statistically significant dose-response effect (p<0.0001) was found between the F concentrations in the drinking water given constantly to the rats during 78 days and the dependent variables F concentrations in plasma, bone (whole, surface and alveolar) and incisor, and the fluorosis index DFIA, with correlations for the linear regression models between 0.87 and 0.98 (Table 1).

The images of rats lower incisors of the control group (0 µg F/mL) show that normal enamel displays an even orange coloration resulting from iron pigmentation, while groups exposed to higher F concentrations in the drinking water (12.5, 25 and 37.5 µg F/mL) show a succession of white and pigmented bands, with an increase in loss of iron pigmentation according to the increase in the F concentration used (Fig. 2).

The mean F concentration to which the oscillating
group (12.5/37.5 µg F/mL) would have been subjected during the experiment was calculated for each dependent variable using the linear equations and varied from 24.6 to 31.8 µg F/mL (Table 1).

The group receiving oscillating doses did not differ significantly (p>0.05) from the group receiving 25 µg F/mL constantly (Table 2).

DISCUSSION

Rats have been extensively used as a model to study fluorosis because plasma F concentrations causing enamel fluorosis in these animals and in humans are similar (4,6), as well as the structure and formation of the enamel (13).

In the present study, to estimate the effect on dental fluorosis of oscillating F doses to which rats were subjected chronically by alternately drinking water containing 12.5 and 37.5 µg F/mL, the dose-response effect of drinking constantly distilled water (control) or water containing 12.5, 25.0 or 37.5 µg F/mL was determined on some biological parameters related to F metabolism and fluorosis.

A significant increase in the value of all variables analyzed was observed according to the increase in F concentration in the water for the constant groups (Table 1), in agreement with other studies with rats (14). Also, the effect of increasing levels of F exposure was satisfactory reflected (r=0.88) in increasing fluorosis severity.

Table 1. Correlation (r) between fluoride concentration in rats plasma, bone and incisor, and dental fluorosis by DFIA index (y) as a function of constant fluoride concentration in water (x), linear equation and estimated F concentration in the water given to the rats of the oscillating group (12.5/37.5 µg F/mL).

<table>
<thead>
<tr>
<th>Y</th>
<th>x</th>
<th>r*</th>
<th>Equation</th>
<th>Estimated [F] in water for the oscillating group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma</td>
<td>[F] in water</td>
<td>0.872</td>
<td>log y = 0.381+0.013x</td>
<td>28.7 ± 4.2</td>
</tr>
<tr>
<td>Whole femur</td>
<td>[F] in water</td>
<td>0.981</td>
<td>y = 250.960+33.771x</td>
<td>24.6 ± 3.2</td>
</tr>
<tr>
<td>Femur surface (10.0 µm)</td>
<td>[F] in water</td>
<td>0.917</td>
<td>y = 764.653+30.427x</td>
<td>31.8 ± 3.7</td>
</tr>
<tr>
<td>Femur surface (24.7 µm)</td>
<td>[F] in water</td>
<td>0.941</td>
<td>y = 535.008+37.751x</td>
<td>31.8 ± 4.2</td>
</tr>
<tr>
<td>Femur surface (46.2 µm)</td>
<td>[F] in water</td>
<td>0.948</td>
<td>y = 458.152+31.762x</td>
<td>29.7 ± 5.9</td>
</tr>
<tr>
<td>Alveolar bone surface</td>
<td>[F] in water</td>
<td>0.973</td>
<td>y = 461.328+34.286x</td>
<td>25.7 ± 3.9</td>
</tr>
<tr>
<td>Apical (tooth)</td>
<td>[F] in water</td>
<td>0.976</td>
<td>y = 99.350+16.580x</td>
<td>26.0 ± 3.8</td>
</tr>
<tr>
<td>Middle (tooth)</td>
<td>[F] in water</td>
<td>0.979</td>
<td>y = 96.388+15.685x</td>
<td>26.0 ± 3.6</td>
</tr>
<tr>
<td>Incisal (tooth)</td>
<td>[F] in water</td>
<td>0.984</td>
<td>y = 89.387+14.401x</td>
<td>24.9 ± 2.7</td>
</tr>
<tr>
<td>Dental fluorosis</td>
<td>[F] in water</td>
<td>0.881</td>
<td>log y = 0.021 + 0.017x</td>
<td>24.8 ± 6.4</td>
</tr>
</tbody>
</table>

* p<0.0001 for all variables
quantified by the DFIA index method developed for this research.

The estimated water F concentration to which the rats were subjected by the oscillating concentrations was very close to 25 µg F/mL for all metabolic variables analyzed (Table 1), which is the expected mathematical mean of drinking water containing 12.5 and 37.5 µg F/mL given at symmetrically alternating periods. Indeed, the group subjected to the oscillating dose did not differ significantly from the group drinking constantly water containing 25 µg F/mL in any biological variable analyzed and on fluorosis development (Table 2).

These findings are supported by the relationship between F concentration in blood and bone. It is known that there is no homeostatic mechanism to maintain blood F concentration (3), and that F will either be taken up from blood by bone or, when the F exposure is interrupted, the concentration in the blood could be maintained for some days due to the F coming from the exchangeable bone sub-compartment (15), but later will decrease (14). Thus, F accumulated in the bone could contribute to increase fluorosis by sustaining, for a certain period of time, blood F concentration systemically or locally in fluids adjacent to maturing enamel.

It has been hypothesized that after an acute F dose, the gradual release of F from the adjacent bone could be sufficient to cause enamel fluorosis (16) and the similar F concentration on the surface of alveolar bone for the oscillating and 25 µg F/mL groups (Table 2) show some evidence favoring this hypothesis, even for chronic dose. Accordingly, F concentration in the femur and other long bones would contribute to a steady state plasma F concentration (14) and the findings showed that the oscillating and the 25 µg F/mL groups did not differ concerning femur and plasma F concentrations (Table 2). Therefore, the same F concentrations in incisors of oscillating and 25 µg F/mL groups (Table 2) could be explained by a local and systemic effect.

Moreover, the similar fluorosis observed for the oscillating and the constant 25 µg F/mL groups may also be explained by the timing of enamel formation under this metabolic condition. The main effect of chronic low F dose occurs during enamel maturation (17) and this phase takes approximately 10.5 days in rats (18). Thus, enamel under maturation during a given period would partly be exposed to both the oscillating concentrations. As rat lower incisors are totally renewed in approximately 60 days, the clinical outcome on enamel would reflect the mean effect of the oscillating F concentrations given each 3 days during the 78 days

<table>
<thead>
<tr>
<th>[F] in water (µg F/mL)</th>
<th>Plasma (µM)</th>
<th>Whole femur (µg F/g)</th>
<th>Femur surface (µg F/g)</th>
<th>Alveolar bone surface (µg F/g)</th>
<th>Incisor (µg F/g)</th>
<th>Dental fluorosis (DFIA index)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>2.1±0.3</td>
<td>249.2±91.5</td>
<td>736.8±131.4</td>
<td>448.7±77.5</td>
<td>114.1±29.1</td>
<td>1.1±0.3</td>
</tr>
<tr>
<td>12.5</td>
<td>4.3±0.5</td>
<td>714.9±41.8</td>
<td>1217.1±194.8</td>
<td>921.8±125.3</td>
<td>312.9±31.9</td>
<td>1.7±0.6</td>
</tr>
<tr>
<td>25.0</td>
<td>5.5±1.7</td>
<td>1016.9±105.7</td>
<td>1464.7±218.2</td>
<td>1292.5±186.0</td>
<td>457.1±62.1</td>
<td>3.4±1.2</td>
</tr>
<tr>
<td>37.5</td>
<td>7.5±2.4</td>
<td>1555.7±112.2</td>
<td>1922.1±192.6</td>
<td>1753.7±99.4</td>
<td>756.9±45.0</td>
<td>4.1±0.9</td>
</tr>
<tr>
<td>37.5</td>
<td>5.9±0.8</td>
<td>1082.7±108.6</td>
<td>1731.9±112.3</td>
<td>1343.9±133.9</td>
<td>529.8±69.1</td>
<td>2.8±0.7</td>
</tr>
</tbody>
</table>

*In order to fit the assumptions for this test, data of plasma fluoride and dental fluorosis were log-transformed. *This group differed statistically from all groups for all variables (p<0.05). *This group did not differ statistically from group received 25 µg F/mL for all variables (p>0.05).
of the experiment. Indeed, the same effect on fluorosis by oscillating group compared with the 25 µg F/mL is supported by F concentration in the incisor, which was similar for both groups (Table 2).

The present study gives support to the epidemiological observation that short fluctuations in F concentration in the water above optimal do not necessarily result in a higher fluorosis risk (7). This could be explained by the small thickness of enamel formed during each period of exposure to a given dose, suggesting that the mean exposure to F is the one that adequately reflects the resulting enamel fluorosis. The lower fluorosis risk found in a city with oscillating water F concentrations could thus be explained by longer periods in which the F content in water was kept below the optimal value, resulting in a lower average F dose when compared to a city in which water concentration was kept within the optimal range.

In summary, the findings of this study suggest that the biological effect of F on animals chronically exposed to symmetrical oscillating F doses reflect the effect of the mean of the oscillating doses. However, limitations on the extrapolation of the present results should be taken into account considering the model used and the necessity to conduct another study to evaluate the effect of oscillating F doses during asymmetrical periods of exposure, which is under investigation in our research lab.

RESUMO

Considerando que a concentração de fluoreto no sangue varia de acordo com a exposição ao fluoreto, e que a fluorose dental está relacionada com a quantidade de esmalte formado sob determinada dose de fluoreto, este estudo avaliou se a fluorose resultante da exposição a doses oscilantes de fluoreto seria semelhante àquela causada pela exposição a uma dose constante, representativa da média das oscilações durante um determinado tempo. Durante 78 dias, ratos receberam água com concentrações constantes de fluoreto de 0; 12,5; 25 ou 37,5 µg F/mL, ou concentrações oscilantes de 12,5 e 37,5 µg F/mL alternados a cada 72 h (média de exposição = 25 µg F/mL). Concentrações de fluoreto no plasma, fêmur e dentes incisivos dos ratos foram determinadas após o período experimental. O índice de fluorose, observado nos incisivos dos ratos, foi quantificado usando um método de análise de imagem desenvolvido para essa pesquisa, denominado de índice de fluorose por análise de imagem (DFIA, em Inglês). A concentração de fluoreto no plasma, fêmur e incisivo dos ratos, assim como o DFIA, aumentaram de forma linear para as concentrações constantes de fluoreto na água (p<0,0001, r=0,87-0,98). Não houve diferença significativa entre o grupo que recebeu doses oscilantes e o grupo que recebeu 25 µg F/mL (p=0,05). Os resultados sugerem a fluorose dental decorrente de exposição crônica de animais a doses de fluoreto oscilantes e simétricas reflete o efeito metabólico da média da oscilação.

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