Morphological aspects in remineralizing potential of Silver Diamine Fluoride

LAURA IDORĂȘI¹, EMANUELA LIDIA CRĂCIUȘNECU¹,², ADRIAN TUDOR STAN¹,², COSMIN SINESCU¹,², ANA CODRUȚA CHIȘ⁵, DARIAN ONCHIŞ-MOACĂ⁴, MIHAI ROMÎNU¹,², MEDA LAVINIA NEGRUȚIȘ¹,²

¹Department of Prostheses Technology and Dental Materials, Faculty of Dental Medicine, Victor Babeș University of Medicine and Pharmacy, Timișoara, Romania
²Research Center in Dental Medicine Using Conventional and Alternative Technologies, Timișoara, Romania
³Research Institute for Biosafety and Bioengineering, Faculty of Agriculture, King Michael I of Romania Banat University of Agricultural Sciences and Veterinary Medicine, Timișoara, Romania
⁴Signal, Image and Machine Learning Team, Department of Computer Science, West University of Timișoara, Romania

Abstract

Objectives: The purpose of this study was to demonstrate the efficacy of Silver Diamine Fluoride (SDF) antibacterial solution in penetrating the demineralized areas of enamel. Materials and Methods: It was considered a group of four extracted teeth (with no color fading, fissures, decay, or demineralization). Each tooth was sectioned in two equal parts, in mesio-distal direction, using a dental handpiece and a special rounded, flat bur. Each specimen was demineralized, for one minute, with 45% orthophosphoric acid, on occlusal and proximal zones. The specimens were then washed and dried with water-air dental syringe. All the probes were inspected with an optical microscope and enamel thickness was digitally measured. Advantage Arrest (Elevate Oral Care, USA), which contains SDF, was applied on the previous demineralized zones. The penetration of the substance was visually inspected with the optical microscope and electronically measured. Results: It was observed an improvement in remineralizing the white spots on enamel surfaces, the optical microscope being able to detect both demineralization and the penetration of SDF through enamel. Conclusions: Based on our in vitro study, SDF (Advantage Arrest) was capable to induce/increase enamel remineralization, through SDF penetration.

Keywords: silver nanoparticles, white spot lesions, enamel remineralization, Silver Diamine Fluoride penetration, antibacterial activity.

Introduction

The most prevalent diseases of the oral cavity are the carious lesions. They occur due to the demineralization of tooth’s fissures or plain surfaces. The intense action of the organic acids causes the demineralization, with those organic acids originating from fermentation of carbohydrates and degradation of the organic matrix [1].

Speaking in terms of morphology, teeth have a complex structure consisting of enamel, pulp–dentin complex, and cementum. Enamel formation is made by specific cells called ameloblasts, which originate from ectoderm. Enamel’s role is to cover the anatomic crowns of teeth, varying in thickness depending on the area in which is located; it is averaging 2 mm at the incisal edge of the frontal teeth, 2.3–2.5 mm at the premolars’ cusps and maximum 3 mm at molars’ cusps. The cusps of posterior teeth have different mineralization centers, forming lobs that converge together. In some areas, the coalescence may not be complete, resulting in pits and fissures that have enamel reduced in thickness. Simultaneously, the enamel on smooth surfaces, such as proximal areas, has a decreased thickness (0.872 mm to 1.015 mm) [2].

Dentin represents the largest portion of a tooth, being in percent of 70% mineralized, with an organic content calculating for 20% of the matrix and a 10% of water. The physical characteristics of teeth are mainly attributed to enamel. It is also believed that collagen performs as an active, safeguarding protein of the underlying hydroxyapatite (HA) crystallite frame. There are two types of collagen. Type I collagen is known as procollagen, being secreted from cells (fibroblasts, odontoblasts, osteoblasts) into the extracellular gaps. In those gaps is type I collagen converted into tropocollagen. Tropocollagen can self-assemble into fibrils, which are built from the undulate packing of the individual collagen molecules. In dentin, type I collagen gets about 56% of mineral in its fibrils’ pores. Dentin matrix protein-1 initiates the nucleation and modulation of mineral phase’s morphology. During dentinogenesis, there exist three types of mineralization: “matrix vesicle-derived mineralization (in mantle dentin), molecule-derived mineralization – ECM (in majority of dentin), and blood serum-derived mineralization (in peritubular dentin)” [2–4].

In enamel, after the dentin mineralization at dentino–enamel junction (DEJ) was initiated, ameloblast cells will start to secrete enamel matrix proteins: ameloblastin, amelogenin, enamelin and proteinases at the dentin surface, being responsible for a prompt mineralization of ~30% of enamel. The ribbons (first formed enamel crystals) expand between the existing dentin crystal and elongate at the mineralization front where enamel proteins have been
developed. Meanwhile, the ameloblast start secreting huge quantities of enamel matrix proteins. In the moment in which the whole thickness of enamel is formed, the ameloblasts start becoming “protein-resorbing” cells. The mineral content will be slowly diminished from the enamel outer toward the DEJ [4, 5].

The dynamic process of demineralization may be reversed if the white spot lesions are detected in early stages. Caries progression appears when there is a demineralization–remineralization imbalance. While the demineralization progresses, one or more white spots appear due to enamel continue mineral privation. This fact causes visual changes which start with the subclinical stage (white spots) and followed by cavitation [1].

There have been massive searches trying to find the best options for detection of early signs of caries lesions and their treatment. As treatment, all the existent substances try to enhance the enamel’s remineralization (for white spots lesions) or dentin disinfection for profound lesions.

Some authors showed, in a systematic review (2017), that the remineralizing agents, such as casein phosphopeptides along amorphous calcium phosphate (CPP–ACP), casein phosphopeptides including amorphous calcium phosphate and fluoride (CPP–ACFP), Icon (DMG Chemisch-Pharmazeutische Fabrik GmbH, Hamburg, Germany) can induce regression of white spots lesions, in size and their visual appearance, improving esthetics at the same time [6].

Other authors evaluated, in comparison, the capacity of a sealant and an infiltrant of penetrating the fissure caries lesions. They mentioned that the infiltrant was developed for proximal lesions and the “resin infiltration technique” has not been evaluated for fissures lesions. The study revealed that the resin infiltration was much more efficient than the fissure sealing [7, 8].

Other studies demonstrated that silver nanoparticles (AgNPs) can inhibit the deoxyribonucleic acid (DNA) replication of bacteria, when remaining in contact with Ag⁺ ions and major structural changes in the bacteria’s membrane. “The bactericidal effect of AgNPs is size dependent, so that AgNPs mainly in the range of 1–10 nm attach better to the cell’s membrane surface, disturbing the main functions. They are possibly able to interact with sulfur- and phosphorus-containing compounds, such as DNA, having a great contribution to the bactericidal effect”. Accordingly, the AgNPs having dimensions from 2 nm to 5 nm are capable to penetrate the dentinal tubules and to inactivate the bacteria’s metabolism. The incorporation of AgNPs in dental materials may be as a monomer, usually 2-(tert-Butylamino)ethyl methacrylate, for improving the Ag solubility in the resin solution, such a product being Advantage Arrest (Elevate Oral Care, USA), which contains SDF [9].

Aim

This study aimed to evaluate, in comparison, the efficacy of Advantage Arrest in penetrating the tooth’s structure, in two different morphological areas: the proximal and the occlusal fissures.

Materials and Methods

Four extracted teeth, with no discoloration, carious lesions or demineralization were selected and maintained in physiological serum. Each tooth was sectioned in two equal parts, in mesio-distal direction, as shown in Figure 1, resulting eight specimens. A demineralization agent, 45% orthophosphoric acid, was applied on each sample and maintained for one minute, on two different enamel areas: occlusal and proximal. The agent was then washed out and the sample dried with water-air dental syringe, as it can be observed in Figure 1. The final step consisted of visually and digital, optical microscope inspection and the enamel thickness was electronically measured (Figure 2).

Advantage Arrest (Elevate Oral Care, USA) (Figure 3) has been applied on both demineralized areas, proximal and occlusal, without touching the inner surface of the teeth.

The specimens have been inspected with optical, digital microscope (Figure 4), with a focus range of 0–40 mm, a still image capture resolution of 160×120, 320×340, 640×480, 1280×1024, 1600×1200 pixels, digital zoom of 5× and sequence mode and magnification ratio of 20×–800×.

For the statistical analysis, we apply first the non-parametric Mann-Whitney–Wilcoxon analysis [10, 11] to
decide whether the population disseminations are identical without charging them to follow the normal distribution. In the data frame list named SDF penetration, we gather all the values measured after SDF penetration, meanwhile another data column indicates the area type (0 – proximal, 1 – occlusal). In other words, the differentiating factor is the area type. Without supposing that the data has normal distribution, we need to set at $\alpha=0.05$ significance level if the SDF penetration data of proximal and occlusal area have identical data distribution.

To apply an analysis of variance (ANOVA)-like test, which for the case of only two columns is identical with the $t$-test, we first try to correct the lack of normal distribution. Parametric methods, such as $t$-test and ANOVA tests [12] to be meaningful, they assume the dependent variable as approximately normally distributed for every group to be compared.

In addition, statistical analysis has been done using the IBM Statistical Package for the Social Sciences (SPSS) software. It considered elements of descriptive statistics, Box-Plot diagrams, and the $t$-test for the studied variables.

\section*{Results}

It was observed a possible improvement in remineralizing of the white spots on flat and occlusal surfaces, the optical microscope detecting the demineralization and the penetration of the Advantage Arrest through enamel. The areas of demineralization and SDF penetration were measured (in four different points) with the TAGARNO measurement software provided by the digital microscope. The values have been collected and compared in Table 1. It has been detected a remineralization percentage of 52.675\% for proximal areas and of 41.001\% for occlusal areas.

### Table 1 – The values recorded for each area of demineralization, SDF penetration and average percentage of SDF penetration

| Specimen No. | Demineralization | Remineralization | Remineralization | Demineralization | Remineralization | Remineralization | Remineralization |
|--------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|
|              | [mm]             | [mm]             | [mm]             | [mm]             | [mm]             | [mm]             | [mm]             |
| 1.           | 0.356            | 0.163            | 45.787           | 0.560            | 0.145            | 25.893           |
| 2.           | 0.283            | 0.268            | 94.700           | 0.430            | 0.110            | 25.581           |
| 3.           | 0.430            | 0.179            | 41.628           | 0.520            | 0.130            | 25.000           |
| 4.           | 0.520            | 0.083            | 15.962           | 0.250            | 0.070            | 28.000           |
| 5.           | 0.131            | 0.113            | 86.260           | 0.163            | 0.079            | 48.466           |
| 6.           | 0.137            | 0.106            | 77.372           | 0.270            | 0.171            | 63.333           |
| 7.           | 0.211            | 0.151            | 71.564           | 0.304            | 0.208            | 68.421           |
| 8.           | 0.344            | 0.090            | 26.163           | 0.275            | 0.127            | 46.182           |

SDF: Silver Diamine Fluoride.

\subsection*{Statistical analysis}

The null hypothesis is that the SDF penetration values for proximal and occlusal areas are equivalent populations. To test this hypothesis, we applied the test to compare the independent samples and we obtain the following values:

Wilcoxon rank sum test: $W=4096$, $p$-value $<2.2e-16$, alternative hypothesis: true location shift is not equal to 0.
As the p-value turns out to be smaller than 2.2e-16, and is less than the 0.05 significance level, we reject the invalid hypothesis. Next, we continue our analysis with the plot in Figure 5. In this figure, ‘A’ stands for the proximal area and ‘B’ for the occlusal area. We can immediately observe from this plot that in general the SDF penetration is higher in the case of the proximal area A.

![Figure 5 – Statistical analysis plot.](image)

We apply the log10 transform to transform the data into a close to normal distribution because the dependent variable increases more rapidly with increasing independent variable values. We consider \( n=32 \) since we have 32 measurements for each area and \( k=2 \) since we have two categories. We plan to test if the means of the groups are equal, the \( H_0 \) hypothesis for this statistical test. We extract the data on each category, and we compute \( \text{varIntra} \) variable, which is the mean within categories, equal in our case is 435.8386. Next, we compute the \( \text{varInter} \) variable, which is the mean between categories, equal in our case with 3406.676. We can compute now the \( F \)-value for our statistics: \( F=\frac{\text{varInter}}{\text{varIntra}}=7.816371 \) and the critical \( F \)-value, which was \( \text{critF}=q_f(0.95, \, df_1=k-1, \, df_2=n-k)=4.170877 \), with \( df \) being the corresponding data frame and 0.95 the significance level. For taking the decision, we use the following formula if \( (F>\text{critF}) \) reject \( H_0 \), else accept \( H_0 \). Therefore, we reject the \( H_0 \) and this decision can be interpreted as a statistical recommendation for the more efficient clinical use of SDF penetration in the proximal area.

The average value of the SDF penetration percentage for the proximal area was 65.843%, with a standard deviation of 21.454%, the percentage reaching a maximum of 94.70%. In the case of the percentage of SDF penetration for occlusal area, the average was 51.252%, with a standard deviation of 20.284% and a maximum of 94.237% (Table 2; Figure 6).

![Figure 6 – The Box-Plot diagram associated with the measurements for proximal area and occlusal area, respectively, for demineralization and SDF penetration (remineralization), respectively. SDF: Silver Diamine Fluoride.](image)

Analyzing the median value for the proximal area SDF penetration (%), one observes that in more than 50% of the measurements the SDF penetration percentage was 68.3%. In the case of the occlusal area SDF penetration, the median value was 47.5%. The values of the upper quartiles (Q3) showed that in 25% of the measurements, the SDF penetration percentage was higher than 85.9% for the proximal area SDF penetration (\%), respectively higher than 69.5% for the occlusal area SDF penetration (\%) (Table 2; Figure 7).

![Figure 7 – The Box-Plot diagram associated with the proximal area SDF penetration and occlusal area SDF penetration, respectively. SDF: Silver Diamine Fluoride.](image)

There are significant differences for the measurements done for the proximal area demineralization and the proximal area remineralization.
area SDF penetration \((t=6.60, p=0.00<0.05)\), respectively for the occlusal area demineralization and the occlusal area SDF penetration \((t=9.74, p=0.000<0.05)\) (Table 3).

Significant differences have been observed between the SDF penetration percentages for the proximal area and the occlusal area \((t=3.161, p=0.003<0.05)\) (Table 4).

In Figures 8–11, some examples of occlusal and proximal measurements of the specific specimens are provided.

### Table 3 – T-test for proximal area, respectively occlusal area

|                   | Mean | Standard deviation | Standard error of the mean | 95% Confidence interval of the difference | t   | df | Sig. (2-tailed) |
|-------------------|------|--------------------|---------------------------|------------------------------------------|-----|----|-----------------|
| Pair 1            | 0.11 | 0.10               | 0.02                      | 0.08 to 0.15                             | 6.60| 31 | 0.000           |
| Proximal area demineralization vs. Proximal area SDF penetration | 0.19 | 0.11               | 0.02                      | 0.15 to 0.23                             | 9.74| 31 | 0.000           |

### Table 4 – T-test for the proximal area SDF penetration (%) and the occlusal area SDF penetration (%)

|                   | Mean       | Standard deviation | Standard error of the mean | 95% Confidence interval of the difference | t   | df | Sig. (2-tailed) |
|-------------------|------------|--------------------|---------------------------|------------------------------------------|-----|----|-----------------|
| Pair              | 14.5914    | 26.1099            | 4.6156                    | 5.1777 to 24.0050                        | 24.0050|31|0.003            |
| Proximal area SDF penetration (%) vs. Occlusal area SDF penetration (%) | 0.311 mm | 0.255 mm            | 0.333 mm                   | 0.206 to 0.433 mm                        | 3.161| 31 | 0.003            |

SDF: Silver Diamine Fluoride.
Discussions

There is still a difficulty in detecting the early signs of caries lesion, considering the methods for caries detection. The early detection of carious lesions is important for improving the treatment and preventing the major loss of dental structures.

Advantage Arrest’s main component is SDF, with the following chemical formula and concentrations: Ag(NH₃)₂F, content: 25–27% Ag, 5–6% fluoride (F), 9–10% ammonia (NH₃) (w/v) [13].

This product, having incorporated an antibacterial substance and a remineralizing one increases teeth resistance to acid attack by forming Ag–protein conjugates in decayed tooth’s surface. It, also, increases the mineral density by increasing HA and fluorapatite. Ag⁺ and F⁻ ions penetrate about 25 μm into enamel and 50–200 μm into dentin; F promotes remineralization, and Ag promotes antimicrobial action. Clinical characteristic and possible disadvantage to take into consideration is the discoloration of the tooth’s structure when applied on demineralization and the soft tissues [14–18].

Visual inspection and radiographic examinations are the most often used techniques for caries and structural abnormalities, such as denticles detection; unfortunately, none of these methods is sufficiently sensitive for detecting the first signs of caries. Furthermore, visual inspection and radiographs cannot detect the penetration and efficacy of any antibacterial solutions, which may be applied to stop bacterial metabolism [19].

The dental structures generate a response to inflammatory acid attack through the connective tissue called dental pulp. When the hard dental tissues are not able any more to create a healthy barrier to protect the pulp, it reacts through inflammatory response, through positive T-lymphocytes for the cluster of differentiation (CD)45RO protein, which means that no non-invasive treatments can be fulfilled [20].

To improve detection of early stages in carious lesions, several methods have been tried, such as dental optical coherence tomography (OCT) [21], heterodyne lock-in thermography [22], near-infrared (IR) transillumination and reflectance imaging (in vivo) [23], the most recent one being coherent speckle light scattering pattern [20].

In vivo, the most common used techniques are quantitative light-induced fluorescence, laser-induced fluorescence, fiber optic transillumination [24].

Tooth’s structure remineralization was studied with the aid of polarized light microscope, Shah & Birur detecting “27.306% decrease in lesion depth after 10 days of pH cycling” [25] and by using a fiber optic backscatter spectroscopic sensor, where with Kishen et al. managed to mark initial enamel demineralization and remineralization and to provide useful, significant information that can be used for monitoring changes in a tooth and make comparisons between teeth [26].

It has been shown in clinical and experimental studies that non-cavitary lesions can remineralize if the negative oral environment is altered, using plaque removal technique and F [27–29].

Other research showed that specific, functionalized AgNPs coated with ethylene glycol (EG) or polyvinylpyrrolidone (PVP) have antibacterial effects on different bacteria; the success depending on the AgNPs dose and of the type of bacteria [30].

In our study, there have been made new attempts in capturing both demineralization and SDF penetration, with a digital optical microscope; the SDF penetration for the proximal surfaces having an easy growth in comparison with SDF penetration on occlusal areas. This difference may appear due to the different morphology of the areas; the proximal area being a smooth zone in comparison with pits and fissures, which may not permit the penetration of antibacterial substances. Being an in vitro study, it is necessary some notifications to be considered; the effectiveness of SDF may be affected by numerous clinical factors, such as tooth surfaces topography, size and shape of gingival papilla, patient’s oral hygiene, the bacteria involved in demineralization, the surface’s roughness, the enamel structure (normal, hypocalcified) [2, 31–33].

Conclusions

SDF is efficient in remineralizing tooth’s enamel, and the digital, optical microscope is receptive for evaluation of both enamel demineralization and SDF penetration, in permanent teeth, for proximal and occlusal surfaces. In our study, we succeeded in demonstrating the positive effect of AgNPs on demineralized enamel. In addition to these results, we managed to show differences in remineralizing aspects of occlusal versus proximal enamel surfaces.

Conflict of interests

The authors declare that they have no conflict of interests. All authors read and approved the final manuscript.

Consent for publication

Consent for publication of the results of this study has been obtained from all the participants. All the specimens used in this research are extracted teeth for orthodontic purpose and the patients gave their consent for using the teeth as specimens for research purposes.

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