Uninfiltrated Collagen in Hybrid Layers produced after Reduced Acid-etching Time on Primary and Permanent Dentin

Débora LS Scheffel, Cláudia Huck, Diana G Soares, Fernanda G Basso, Carlos A de Souza Costa, Martha G Brackett, David H Pashley, Josimeri Hebling

ABSTRACT

Aim: This study evaluated the influence of acid-etching time on collagen exposure in adhesive interfaces established on primary and permanent dentin.

Materials and methods: Flat dentin surfaces were produced on sound primary molars and premolars (n=8). The surfaces were divided into mesial and distal halves, and each half was etched with phosphoric acid for 5 or 15 seconds. The teeth were randomly allocated into two groups according to the adhesive system applied: Prime & Bond NT or Prime & Bond 2.1. After the adhesive application, the specimens were processed for Goldner’s trichrome staining. The thickness of the uninfiltrated collagen zone (UCZ) in the hybrid layer was measured under optical microscopy. Data were analyzed by analysis of variance and Tukey tests (α = 0.05).

Results: The thickness of UCZ was adhesive dependent. Within the same substrate, the specimens treated with Prime & Bond 2.1 presented thicker UCZ when etched for 15 seconds.

Conclusion: The thickness of UCZ in hybrid layers is directly affected by acid-etching time and by the adhesive system applied. Primary dentin seems to be more susceptible to collagen exposure than is permanent dentin.

Clinical significance: Both acid-etching time and adhesive system can influence the amount of exposed collagen interfering on resin–dentin bond quality, especially on primary dentin.

Keywords: Collagen, Degradation, Dentin, Hybrid layer, Resin.
monomers and collagen fibrils, called the hybrid layer. The architecture of the hybrid layer has been considered mainly responsible for the micromechanical retention of adhesive restorations to dentin as well as for the sealing of the etched substrate. The establishment of a perfect hybrid layer, where water is completely displaced from demineralized dentin by the adhesive system and the exposed collagen fibrils are fully infiltrated by monomers, is not a clinical reality. The incomplete infiltration of demineralized dentin caused by the presence of excessive residual water and by the different molecular weights and hydrophilicity of resin monomers result in a zone of naked, unprotected collagen at the bottom of the hybrid layer that may be degraded by endogenous proteases over time. Resin–dentin bond aging process ultimately results in a functional failure of the hybrid layer.

The literature suggests that primary dentin may be more sensitive to resin–dentin bond degradation than permanent dentin. The lower mineral content and differences in structure have associated primary dentin with a lower buffering capacity, which may lead to the creation of 25 to 30% thicker hybrid layers in primary dentin compared with permanent dentin when the substrates are etched for the same period of time. Thicker hybrid layers are more likely to contain pores and imperfections, resulting in lower bond-strength values.

Demineralization of dentin by an acidic agent is time dependent, being directly influenced by the duration of the contact between acid-etching solution/gel and dentin surface. Increased etching time generally results in a deeper dentin demineralization and thicker hybrid layers that tend to be poorly resin infiltrated.

MATERIALS AND METHODS

Sound human premolars (n = 8) and primary molars (n = 8) were obtained from the Human Teeth Bank at the Araraquara School of Dentistry (UNESP, São Paulo, Brazil) under a protocol approved by the Institutional Ethics Committee (process #15/08) and stored in a 0.1% thymol solution at 4°C. To minimize differences related to posteruptive mineralization degree, only premolars that were in function in the oral cavity for at least 3 years were included in this study.

To facilitate the handling of primary teeth with advanced root resorption, the pulp chambers were filled with low-viscosity resin (Filtek Flow, 3M ESPE, St. Paul, MN, USA) and artificial roots were created with composite resin (Z250 3M ESPE, St. Paul, MN, USA). Flat mid-coronal dentin surfaces were prepared using a 0.3-mm-thick ISOMET saw (ISOMET 1000 Buehler, Lake Bluff, IL, USA) under water cooling. Then, each surface was divided into two halves (mesial and distal) by a groove made with a 0.15-mm-thick saw (ISOMET 1000). Each dentin half was selected and acid-etched with 35% phosphoric acid gel (Scotchbond Etchant 3M ESPE, St. Paul, MN, USA) for 5 or 15 seconds, followed by rinsing with deionized water for 10 seconds and blotted with absorbent paper in order to obtain a moist surface.

The teeth were randomly divided into two groups (n = 4) according to the adhesive system used (Table 1). The adhesives were applied according to the manufacturer’s instruction and light-cured for 10 seconds. Four additional layers of the adhesive systems were added and photoactivated in order to permit cutting the specimens in a microtome. The teeth were stored in water at 37°C for 24 hours. All procedures involving photoactivation were conducted with the same light unit, Optilux 500 (Kerr, Danbury, CT, USA), after checking the irradiance using a radiometer (450 ± 10 mW/cm², Model 100, Optilux Radiometer Kerr, Danbury, CT, USA).

Table 1: Commercial name (manufacturer), classification and composition (major components) of adhesive systems

<table>
<thead>
<tr>
<th>Commercial name</th>
<th>Classification</th>
<th>Components</th>
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<tr>
<td>Prime &amp; Bond NT (Dentsply De Trey, Konstanz, Germany)</td>
<td>Total-etch simplified system</td>
<td>Resins and di-trimethyl-propane, amorphous silica, PENTA (dipentaerythritol penta acrylate monophosphate), photoinitiators, stabilizers, and methylamine hydrochloride acetone.</td>
</tr>
<tr>
<td>Prime &amp; Bond 2.1 (Dentsply Caulk, Milford, EUA)</td>
<td>Total-etch simplified system</td>
<td>Resins and trimethylpropane di-, PENTA, butylated hydroxytoluene, 4-methyl amino ethyl benzoate, Photoinitiators, and methylamine hydrochloride acetone.</td>
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**Histological Procedures**

After 24 hours in water at 37°C, premolar roots were removed in a cutting machine (ISOMET 1000) and the teeth were sectioned to obtain two or three blocks (2 mm thick × 2 mm wide × 5 mm long) per tooth, so that each block contained both halves (etched for 5 and 15 seconds). The slabs were fixed in a 10% formalin solution for 48 hours and slightly demineralized in a 10% Morse solution for 48 hours without agitation. Then, the blocks were rinsed in running tap water for 24 hours, neutralized in a 5% sodium sulfate solution for the same period of time, and rinsed again with water for 24 hours. The specimens were dehydrated in a series of increasing ethanol solutions (70 to 100%), cleared in xylol, and embedded in paraffin under vacuum. Approximately 5-μm-thick serial sections were cut from each block in a microtome (820 Spencer Microtome, American Optical Corp., Buffalo, NY, USA) and stained with Goldner’s trichrome.

**Optical Microscopy Analysis**

One 5-μm section from each block was randomly chosen and analyzed under 400× magnification in light microscopy (Olympus BX51 and Camedia C5060 camera, Olympus Corp., Tokyo, Japan). Using the UTHSCSA Image Tool software (University of Texas Health Science Center, San Antonio, TX, USA), five measurements of the thickness of the UCZ (red color) at the bottom of the hybrid layer were made in each section by demarcating one central and four equidistant points (two to the right and two to the left). The measurements were repeated by the same calibrated operator within a 1-week interval, and the means were calculated.

**Statistical Analysis**

The intraexaminer agreement for the two UCZ thickness measurements in the resin–dentin bonds was determined by an intraclass correlation coefficient (ICC). Data sets were tested by a three-way analysis of variance at three fixed criteria: Etching time (two levels), adhesive system (two levels) followed by a Tukey’s test. All statistical tests were considered at a significance level of 5%.

### RESULTS

Variations in thickness of UCZ in the hybrid layers are presented in Table 2. A high intraexaminer correlation (ICC=0.88) was obtained between the first measurement of UCZ thickness and after 7 days. Red-stained exposed collagen fibrils were identified at all resin–dentin bonds. Only the primary factors, “substrate” and “etching time,” exerted a statistically significant effect on these values (p < 0.05). For the same type of substrate, increasing etching time from 5 to 15 seconds increased the thickness of UCZ. However, this difference was statistically significant only when Prime & Bond 2.1 was used. Comparing the same etching time, thicker UCZ values were observed for primary dentin compared with permanent dentin, although statistical significance was verified only when the adhesive system Prime & Bond 2.1 was applied. Thus, for this system, a greater exposure of collagen fibrils was observed in primary dentin etched for 5 seconds compared with permanent dentin etched for the same period of time. However, there was no difference in the thickness of UCZ zone between primary and permanent dentin etched for 15 seconds.

Interfaces produced by the system Prime & Bond 2.1 are shown in Figures 1A and B. Incomplete infiltration of demineralized dentin was observed in all specimens, regardless of substrate or etching time. This was evidenced by the presence of uninfiltred collagen fibrils available for reaction with Goldner’s trichrome (stained red).

**DISCUSSION**

Ideally, after acid-etching and adhesive application, demineralized dentin should be completely encapsulated by monomers to form a perfect hybrid layer that effectively seals dentin. However, many studies have shown that the complete infiltration of organic matrix seldom occurs, permitting the presence of naked collagen within the hybridized zone.

### Table 2: Thickness (μm) of exposed collagen zone in dentin–adhesive bond produced on primary and permanent teeth vs etching time

<table>
<thead>
<tr>
<th>Adhesives</th>
<th>Primary dentin</th>
<th>Permanent dentin</th>
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<tr>
<td></td>
<td>5 seconds</td>
<td>15 seconds</td>
</tr>
<tr>
<td>Prime &amp; Bond NT</td>
<td>1.99 (0.66)\textsuperscript{a,b}</td>
<td>2.18 (0.31)\textsuperscript{A,B}</td>
</tr>
<tr>
<td>Prime &amp; Bond 2.1</td>
<td>1.82 (0.41)\textsuperscript{A,b}</td>
<td>2.33 (0.31)\textsuperscript{A,c}</td>
</tr>
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\textsuperscript{a,b}Values represent mean (standard deviation), n=8. Capital letters compare vertical column data, while lowercase letters compare horizontal lines. Different letters are significant at p < 0.05.
Water present in etched dentin keeps collagen fibrils expanded and can be classified as free (unbound) and collagen-bound water. While free water represents approximately 75 to 79% of total water content in demineralized dentin, 21 to 25% is bound water. The application of polar solvents, such as ethanol and acetone may chemically dehydrate dentin, creating a more favorable environment for adhesive infiltration. However, Agee et al demonstrated that when applied for 60 seconds on dentin, ethanol and acetone remove respectively, 25 and 34% of unbound water, leaving behind most of the total water content permeating collagen fibrils. Remaining water impairs monomer infiltration and is used by proteases to degrade unprotected collagen in the hybrid layer.

Optical microscopy coupled with Goldner’s trichrome stain has been used to detect exposed collagen fibrils in resin–dentin bonds. This technique was validated by many studies in which the unprotected collagen fibrils were stained by the trichrome dye and turned apparent red color under light microscopy. The presence of the red coloration allows measurement of UCZ thickness at the bottom of the hybrid layer.

In the present study, a direct relationship between etching time and UCZ thickness was observed for both primary and permanent dentin, in agreement with results reported by Wang et al. Those authors noted that the demineralization of permanent dentin after application of 35% phosphoric acid for 15 seconds was two times deeper than the demineralization observed after 10 seconds. This indicates that despite the buffering capacity of dentin, greater dissolution of mineral phase is observed when etching time is extended, making dentin demineralization a time-dependent process.

Osorio et al demonstrated that rougher dentin surfaces were produced when etching time was decreased from 15 to 7 seconds, resulting in an increased surface area for interaction with the adhesive system, and consequent improvement in bond strength to dentin. This led us to test whether reducing etching time restricts dentin demineralization and enhances resin–dentin bond quality. However, Figure 1 shows that collagen fibrils still remain exposed at the base of adhesive bond even when the substrates (primary and permanent dentin) were etched for only 5 seconds. This finding suggests that the extent of dentin demineralization is not the only factor that influences monomer infiltration. Other factors, such as molecular weight, hydrophilicity of the adhesive systems, and substrate moisture may also play an important role in the process of hybrid layer stabilization.

The composition of the adhesive system seems to be extremely influent on monomer infiltration. The results observed for Prime & Bond NT were different from those observed for Prime & Bond 2.1. Data collected for Prime & Bond NT require acceptance of the null test hypothesis.

Figs 1A and B: Resin–dentin bonds produced by Prime & Bond 2.1 system applied on: (A) Primary dentin etched for 5 seconds; (a’) primary dentin etched for 15 seconds; and (B) permanent dentin etched for 5 seconds, and (b’) permanent dentin etched for 15 seconds.

A: Adhesive system; MD: Mineralized dentin (green); exposed collagen (*red). Goldner Trichrome 400×
that reducing etching time from 15 to 5 seconds does not change the thickness of the UCZ. However, Prime & Bond 2.1 groups showed a higher sensitivity between depth of demineralization and monomer infiltration when acid-etching time was increased.

Both Prime & Bond NT and Prime & Bond 2.1 use acetone as the volatile solvent. However, Prime & Bond NT contains nanofiller particles to increase its viscosity so that it will not spread too thin and evaporates too rapidly. This system was less sensitive to etching time than Prime & Bond 2.1. It is likely that the diffusion of adhesive monomers into the collagen network is also time dependent.6 According to the manufacturer’s directions, after adhesive application and solvent evaporation, Prime & Bond 2.1 is immediately photoactivated, while Prime & Bond NT must remain at rest for 20 seconds on dentin surface prior to light activation. Adhesive infiltration may be more efficient when one waits a few seconds after adhesive application and photoactivation.5 Thicker UCZ was observed for primary teeth compared with permanent teeth, which was statistically significant when the substrates were etched for 5 seconds. These results can be explained based on structural and morphological differences between the dentin of primary and permanent teeth.27-29 It has been shown that, regardless of acid-etching time, primary dentin is more sensitive to acid than is permanent dentin.4 Primary dentin contains a larger number of dentinal tubules per unit area than permanent dentin29 and is more porous in the intertubular region.28,29 Since dentinal tubules are considered the preferred diffusion pathways for acid penetration in the first seconds of application,6 primary dentin demineralization may also etch faster than does permanent dentin.4 Additionally, primary dentin has lower mineral content,27 which confers greater solubility to primary dentin and a lower buffering capacity, resulting in higher demineralization when etched for the same time than permanent teeth.12,18

Reducing etching time is a clinical procedure that may minimize the discrepancy between the dentin demineralization depth and monomer infiltration, increasing the quality of resin–dentin bonds.13,15 However, there is still concern that very short etching periods are not sufficient to fully dissolve the smear layer and the underlying dentin mineral.14 It is especially relevant for permanent dentin that seems to present smear layer even after 5 seconds in contact with phosphoric acid, as observed in Figure 1. The maintenance of smear layer results in lower bond strength for permanent teeth.5 It is also important to highlight that the favorable results obtained with reduced etching times in the present study and in the literature were mainly produced on sound dentin.13,17 Therefore, further investigation should evaluate the influence of shortening etching time on caries-affected dentin, since this substrate is often used for bonding.

CONCLUSION
The depth of demineralization and the amount of uninfiltrated dentin collagen are time and adhesive dependent. In general, thicker UCZs were observed for primary teeth compared with permanent teeth.

CLINICAL SIGNIFICANCE
By reducing acid-etching time, one can decrease the depth of demineralization in order to minimize the amount of exposed collagen and increase resin–dentin bond quality and stability.

REFERENCES