The success of such repairs depends not only on the type of the material used for the initial restoration and the repair procedure utilized, but also on the surface treatment applied to the surface to be repaired. Treating the surface is mandatory, owing to the lack of reactive methacrylate (MC) groups in the old resin, and because the sorption of water by the old restoration restricts the addition of new material.

However, the problem for the clinician is that the type and brand of the composite used in the previous restoration is often unknown. The problem is aggravated when different composites with organic matrices other than dimethacrylates have been introduced. Furthermore, various protocols of intraoral repair systems are exploring on the repair potential of indirect aesthetic restorations, ceramic, feldspathic porcelain, and zirconia.

The repair of MC-based composite restorations is widely considered to be a reliable procedure in

**INTRODUCTION**

With the introduction of dental adhesive technology, tooth-colored composite restorations have gained wide popularity in recent decades. Despite innovative improvements over the years, and the long-term stability of composite restorations, failures continue to occur.[1] In cases of failure or fracture of an adhesive restoration, the repair procedure offers a minimally invasive alternative to complete replacement of a restoration.[2] Repair is an alternative to complete removal that preserves the tooth, as it is often difficult to remove an adhesive restoration without removing an integral part of the tooth.[3,4] For all of these reasons, a qualitative, optimal method of repairing composite restorations has become a desirable alternative.[2] The evidence as it currently stands seems to favor repair over replacement, but it is not conclusive.[5] The success of such repairs depends not only on the type of the material used for the initial restoration and the repair procedure utilized, but also on the surface treatment applied to the surface to be repaired.

Treating the surface is mandatory, owing to the lack of reactive methacrylate (MC) groups in the old resin, and because the sorption of water by the old restoration restricts the addition of new material.[6] However, the problem for the clinician is that the type and brand of the composite used in the previous restoration is often unknown. The problem is aggravated when different composites with organic matrices other than dimethacrylates have been introduced. Furthermore, various protocols of intraoral repair systems are exploring on the repair potential of indirect aesthetic restorations, ceramic, feldspathic porcelain, and zirconia.[7,8] The repair of MC-based composite restorations is widely considered to be a reliable procedure in

**ABSTRACT**

**Objective:** The objective was to investigate the way that various surface treatments could influence the bond strength of the repair of methacrylate (MC) and silorane (SIL) composites. **Materials and Methods:** A total of 160 MC and SIL cylindrical specimens were polymerized and aged in artificial saliva solution for 7 days. Depending on the following surface treatment (diamond bur or air abrasion), and the conditioning procedure (orthophosphoric acid or sodium hypochlorite), 16 groups were formed and repaired either with MC, either with SIL composite. Repaired specimens were subjected to an additional aging procedure in artificial saliva for 7 days, followed by thermo-cycling and then stressed in shear at a rate of 0.5 mm/min until failure. Failure patterns were analyzed using stereomicroscope and scanning electron microscopy. **Results:** MC composite showed statistically significant higher bond strength both as a base or repair material than SIL ($P < 0.001$). Statistically significant differences were not observed, when grinding and conditioning procedures was compared. Pretest failures were observed when aged MC-based composite was repaired with SIL-based. **Conclusions:** Type of composite seems to be the main factor influencing the bond strength of the repair. MC-based composite showed better repairability than SIL composite. Optimum repair conditions should include knowledge of the composite's composition.

**Key words:** Bond strength, methacrylate composite, repair, silorane composite, surface treatment

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European Journal of Dentistry, Vol 8 / Issue 3 / Jul-Sep 2014 353
modern dentistry. With regard to the repair of silorane (SIL)-based restorations however, too few studies have been reported. Moreover, few studies investigating interactions between MC-based composite and SIL-based restorations have been published. In addition, studies comparing the bond quality facilitated by different combinations of surface pretreatments (grinding and conditioning) have not been published.

The purpose of this in vitro study was to investigate the influence of different pretreatment procedures - grinding (by diamond bur [DB] or air abrasion [AA]) and conditioning (orthophosphoric acid [OA] or sodium hypochlorite [HY]) - and different organic matrix compositions (MC- and SIL-based composite), on the shear bond strength of repaired aged composite. The null hypotheses tested were that (a) composites with different organic matrix compositions demonstrate the same reparability, and (b) different surface pretreatment procedures have no influence on the shear bond strength of repaired composites.

MATERIALS AND METHODS

The following composite systems were tested: The Filtek Ultimate (3M ESPE, St. Paul, MN, USA) (MC), a nanofilled MC-based composite with the corresponding bonding agent Single Bond Adhesive, and the Filtek Silorane (3M ESPE, St. Paul, MN, USA) (SIL), a SIL-based composite with the corresponding bonding agent Silorane Adhesive System. The materials used, and their chemical compositions are listed in Table 1.

Standardized disc-shaped composite substrates of 4 mm diameter and 2 mm thickness were prepared for the test specimens, using cylindrical shaped teflon molds with a hole in their center, of those same dimensions. Prior to filling the holes with the resin material, the molds were set on a glass microscope plate in order to achieve a flat surface of the specimen after light-curing. Resin composite was condensed into the hole with a hand instrument, to a thickness of 2 mm, and a second microscope slide was firmly pressed onto the top of the mold to remove excess resin composite, and to create a flat surface. Polymerization of the composite specimens was achieved through the application of a light-emitting diode (LED) polymerisation device for 20 s (Heraeus Kulzer Translux Power Blue Germany 50/60 Hz 15 VA), at a light intensity of 1000 mW/cm². Light intensity was periodically verified by a radiometer after the processing of every five specimens (Bluephase meter Ivoclar, Vivadent, Schaan, Liechtenstein). The specimens were then left to age in artificial saliva for 7 days at room temperature. Artificial saliva was prepared according to Donmez et al. and was comprised of the following components (mmol/L): CaCl₂·2H₂O (0.7), MgCl₂·6H₂O (0.2), KH₂PO₄ (4.0), KCl (30.0), HEPES buffer (20.0), Na₂HPO₄ (3.0). The specimens were then removed from the artificial saliva, air-dried and randomly assigned to the test groups. The chosen combination of surface treatments (grinding procedure and conditioning procedure) for each group was performed. The two types of surface treatment tested were a grinding procedure with the use of a DB (Komet 107 μm-grit, for 10 s at high speed with water cooling) (DB) and AA (EMS Air-Flow Prep K1 Max device, 50-μm aluminum oxide for 10 s, with the device set with the nozzle at a distance of 5 mm) (AA). In addition, two different conditioning procedures prior to the application of the corresponding bonding agent and the repair composite were compared; application of

| Table 1: Materials, batch numbers, and composition |
|---------------------------------|---------------------------------|
| Adhesive & Composition | Silorane Adhesive & Composition |
| Adhesive Bond 2 Adhesive | Lot no. N120915 |
| (3M ESPE, St. Paul, MN, USA) | (3M ESPE, St. Paul, MN, USA) |
| Composition | Primer: Phosphorylated methacrylates, BisGMA, HEMA, water, ethanol, silane-treated silica filler, initiators, stabilizers |
| BisGMA, HEMA, dimethacrylates, ethanol, water, photo initiator system, methacrylate functional nanocomposite of polyacrylic and poly(titaconic acids, silica nanofiller) | Adhesive bond: Hydrophobic dimethacrylate, phosphorylated methacrylates, TEGDMA, silane-treated silica filler, initiators, stabilizers |
| Resin composite | Filtek Silorane A3 |
| Lot no. N284556 (3M ESPE, St. Paul, MN, USA) | Lot no. N284556 (3M ESPE, St. Paul, MN, USA) |
| Resin matrix composition | 3.4-epoxy cyclohexylcylohexylcyclomethylsiloxane, bis-3.4-epoxy cyclohexylcylohexylphenyl dimethacrylate, initiator, stabilizers |
| Bis-GMA, UDMA, TEGDMA, Bis-EMA | Initiating system: Camphorquinone, iodonium salt, electron donor |
| Filler size/percentage | 20 nm silica filler, 4-11 nm zirconia |
| 78.5% by weight | 0.1-2.0 μm silanized quartz |

Bis-GMA: Bisphenol-A glycidyl dimethacrylate, HEMA: 2-hydroxyethyl methacrylate, UDMA: Urethane dimethacrylate, TEGDMA: Triethylene-glycol dimethacrylate, Bis-EMA: Ethoxylated bisphenol-A dimethacrylate
Spyrou, et al.: Repair of composites

European Journal of Dentistry, Vol 8 / Issue 3 / Jul-Sep 2014

37% OA, and application of 2.5% sodium HY. For the conditioning procedure, 37% OA was placed on the specimens, left for 30 s, washed away with water, and the specimens were then air dried for 10 s. For the 2.5% NaOCl conditioning procedure, the specimens were rinsed with 5 mL NaOCl and air dried for 10 s. Thereafter, bonding procedures were performed on the aged treated composite surfaces using the corresponding bonding agent of the repair composite resin, strictly in accordance with the manufacturer’s instructions.

Following the curing of the adhesive, a teflon matrix which manufactured to fit exactly on the aged surface and to specify the exact dimensions of the repair material (3 mm in diameter and 4 mm in height) was placed over the aged specimen substrates and filled in two increments with the repair composite. Each increment was light-cured for 20 s using the same LED polymerization device and the bonded specimen was removed from the matrix with slight pressure on the top of the repair composite cylinder. This procedure resulted in cylinders of repair composite measuring 3 mm in diameter and 4 mm in height being bonded to the aged resin [Figure 1a]. All bonding procedures were carried out by the same investigator.

In the manner described above, 160 cylinders of repair composite resin specimens comprised of 16 groups of 10 specimens were prepared. The combinations of the materials and surface treatments used are shown in Table 2.

The bonded specimens were subjected to an aging procedure, undergoing storage for 7 days in artificial saliva followed by thermo-cycling (5000 cycles between 5°C and 55°C, dwell time 30 s). Shear testing was conducted using a universal testing machine (Testometric AX, M 350-10KN, Rochdale, England) at a crosshead speed of 0.5 mm/min until fracture. The specimens were fixed on a modified shear fixture, and the load was applied with the use of a chisel [Figure 1b]. The load at fracture, expressed in MPa, was calculated by dividing the peak load by the bonding area, and this value was used in statistical analyses.

The assumption of normality was tested with the Shapiro-Wilk test. Levene’s test for equality of error variances was used to test the assumption of homogeneity. According to the results of the above tests, the nonparametric Mann–Whitney U-test was used for comparisons of the interaction between base material and surface treatment. Adjustment for Type I error resulting from multiple tests was achieved via the Bonferroni method. All analyses were performed with IBM Corp. Released 2010. IBM SPSS Statistics for Windows, Version 19.0. Armonk, NY: IBM Corp., and statistical significance was set at $P < 0.05$.

Failure patterns at the resin/resin system interface were analyzed under a stereomicroscope (Olympus Co., Tokyo, Japan) at ×40 magnification, to determine the failure modes. Failure was considered to be (a) “adhesive”, if it occurred at the resin/
adhesive interface, (b) “cohesive”, if it occurred in aged composite or in new composite, and (c) “mixed”, when both the interface and the composite were involved. The bond failure patterns were not statistically analyzed. Three specimens from each group that were tested for shear bond strength were randomly selected for evaluation using scanning electron microscopy (SEM, Joel, J.S.M. - 840 Tokyo, Japan). The specimens were mounted on stubs, sputter-coated with carbon and examined by one evaluator under SEM at 19 KV.

RESULTS

Initial repair was not possible for some groups and resulted in many pretest failures, particularly for aged MC with the application of SIL using the SIL adhesive. Failure was observed for all kinds of surface treatment tested. Table 2 shows the pretest failure groups and groups with successful bonds. The pretest failures were not statistically analyzed. Higher bond strength values were achieved when aged MC was repaired with the application of new MC with DB grinding, conditioning with OA, and Single Bond Adhesive was used as the bonding agent (69.93 ± 13.98 MPa). In contrast, lower bond strength values were evident when aged SIL was repaired with new MC with DB grinding, conditioning with OA, and bonding with Single Bond Adhesive (25.97 ± 8.67 MPa) [Figure 2]. When comparing base composite resins, ignoring all other factors (surface treatment), statistically significant differences were observed in bond strength between MC and SIL (Mann–Whitney U-test, P < 0.001).

When comparing grinding procedure surface treatments, ignoring all other factors, nonstatistically significant differences were observed between DB and AA (Mann–Whitney U-test, P = 0.761). In addition, no statistically significant differences were observed between OA and sodium HY when conditioning procedures were compared, ignoring all other factors (Mann–Whitney U-test, P = 0.499).

No statistically significant differences were observed when comparing the interaction between surface treatments (Mann–Whitney U-test with Bonferroni adjustment, P > 0.05), ignoring all other factors. Collectively, the above results indicate that the combination of composite based materials (MC-based composite and SIL-based composite) used is the most significant factor influencing the bond strength in this experimental context, as none of the other factors had a significant influence on the bond quality of the repaired composite resins.

No statistically significant differences were observed between any combination of surface treatments when MC was used as the base resin (Mann–Whitney U-test with Bonferroni adjustment, P > 0.05).

When the base resin was SIL, statistically significant differences (Mann–Whitney U-test with Bonferroni adjustment) were observed between DB + OA and DB + HY (P = 0.012), and DB + OA and AA + OA (P = 0.012), while no other comparisons yielded statistically significant differences (P > 0.05).

Table 3 shows the failure mode results of the specimens that were analyzed under a stereomicroscope after shear bond tests. In almost all groups, most fracture modes were mixed, but in the SIL-SIL-DB-OA group, most were adhesive.

![Figure 2: Shear bond strength of repair composite resins](image-url)
DISCUSSION

The first hypothesis that composites with different organic matrix compositions demonstrate the same reparability has to be rejected since at this study composites with different matrix compositions demonstrated repair bond strength variations.

This study shows that MC-based composite can be repaired with a MC-based composite and the corresponding bonding agent, and adequate bonding could be achieved in every combination of surface treatment that was investigated in this study [Figure 3a and b]. Conversely, MC-based composite could not be effectively repaired with SIL-based composite and the corresponding bonding agent, since no combination of surface treatment investigated in this study established an adequate bond.

Artificial aging of MC-based composites creates degradation processes which cause increased water sorption.[4] Owing to their hydrophobic siloxane component, SIL-based composites are more hydrophobic than MC-based composites.[26] This may influence the reparability of MC-based composites with SIL-based composite and the corresponding bonding agent.

In contrast, SIL-based composite could be effectively repaired with SIL- and MC-based composite and the corresponding bonding agent [Figure 4a and b]. SIL-based composite is more resistant to water sorption and disintegration, and this leads to smaller changes in material properties resulting from aging.[24,27] SIL-based composite could only be used effectively as a repair composite for aged SIL with the surface treatment combination of DB and OA. Although, the poor shear bond strength results along with the high rate of adhesive failures that the failure mode analysis showed for this group are evidence of the weakness of bonding in this context. This result is concordant with a study reported by Ivanovas et al.,[15] who concluded that SIL-based composite can be effectively repaired with MC-based composite. The results of this study show that the MC resins that were repaired with the same composite reached the highest possible bond strength. The failure pattern analysis affirms the above results, as a large number of mixed and cohesive fractures were observed.

The reparability of MC-based composite and SIL-based composite differs depending on whether the material is used as a base material or as a repair material. MC-based composite yielded good results when used as a base material, as well as a repair material. In contrast, SIL-based composite yielded good results when used as a base material, but was evidently a poor repair material. These results are concordant with those of a study reported by Baur and Ilie[22] who concluded that the properties of materials can differ strongly depending on whether they are used as a base material, or a repair material.

Table 3: Failure mode analysis of fractured surfaces

| Failure mode | Group          | Adhesive (%) | Cohesive (%) | Mixed (%) |
|--------------|----------------|--------------|--------------|-----------|
| MC-MC-DB-OA | 2 (20)         | 1 (10)       | 7 (70)       |
| MC-MC-DB-HY | 3 (30)         | 1 (10)       | 6 (60)       |
| MC-MC-AA-OA | 4 (40)         | 1 (10)       | 5 (50)       |
| MC-MC-AA-HY | 3 (30)         | 1 (10)       | 6 (60)       |
| SIL-MC-DB-OA| 5 (50)         | 1 (10)       | 4 (40)       |
| SIL-MC-DB-HY| 4 (40)         | 0 (0)        | 6 (60)       |
| SIL-MC-AA-OA| 4 (40)         | 1 (10)       | 5 (50)       |
| SIL-MC-AA-HY| 3 (30)         | 0 (0)        | 7 (70)       |
| SIL-SIL-DB-OA| 7 (70)       | 1 (10)       | 2 (20)       |

OA: Orthophosphoric acid 37%; HY: Sodium Hypochlorite 2.5%, AA: Air abrasion, DB: Diamond bur, SIL: Silorane matrix composite, MC: Methacrylate matrix composite

Figure 3: Scanning electron microscopy images of a specimen where aged methacrylate (MC) has been repaired with MC after the aged surface was treated with air abrasion and conditioned with orthophosphoric acid. (a) Adhesive failure with mixed components. The area marked inside the square is the magnified area in “b” (×20). (b) Detailed view of the fractured area. Inorganic fillers originating on the repair resin are detectable on the aged resin surface (×500)

Figure 4: Scanning electron microscopy images of a specimen where aged silorane was repaired with methacrylate after the aged surface was treated with diamond bur and conditioned with hypochlorite. (a) Mixed fracture. Distinguishable are the repair composite (R), the adhesive failure area (A) and the base composite (B). The area marked inside the square is the magnified area in “b” (×20). (b) Magnification of the failure line in the base composite area (×500)
Some sources propose the additional use of silane as an intermediate agent, in order to effectively repair a SIL-based composite. On the contrary, a new research on microtensile bond strength of SIL-based composites repaired with MC-based composites showed that silane did not contribute to an increase in the tensile strength. In this study, a silane solution was not employed prior to the adhesive, in order to focus on the influence of grinding and conditioning of the repair surface on the bond. Implementation of silane enhances the wetting of the surface for the bonding agent, which then infiltrates easily into the irregularities created by the surface roughening, and the silane-coated composite surface becomes more reactive to the MC groups of the repair resin.

With regard to surface treatment, it should be noted that both DB and AA procedures are used to increase the micromechanical retention of the new material onto the aged composite. In this study, there were no statistically significant differences between the two grading procedures tested. These results are concordant with the results of Rathke et al. and Yesilyurt et al., who investigated the influence of different methods of mechanical roughening on bond quality between new and aged composite.

A treated composite either with DB or with AA infiltrates on its surface organic and inorganic particles originated on the environment. The conditioning procedure using OA does not aim to decalcification of the resin surface similar with what happens in the hard dental tissues surfaces. It is used in an order to obtain a cleansing effect by removing inorganic particles. On the contrary, sodium HY is a known organic solvent, which is used in many dental applications in order to remove the organic particles from surfaces. Statistical analysis in this study has shown that there were no statistically significant differences between the two types of conditioning procedures.

The statistical analysis of the results of this study showed that the best two combinations of surface treatment were DB with OA, and AA with OA.

Therefore, the second hypothesis should be rejected up to a point, and it is concluded that within the limitations of this study the surface treatment has a minor influence on the shear bond strength of repaired composites. This is concordant with Loomans et al., who attested that none of the common surface treatments can be recommended as a universally applicable repair technique, since their effectiveness is material-dependent.

The authors are in agreement with the conclusions of a recent systematic review that repair of restorations is a valuable method of improving their quality, and can yield acceptable results. However, methodologically sound, randomized, controlled, long-term clinical trials are required, in order to facilitate evidence-based recommendations.

CONCLUSIONS

Within the limitations of this study, it can be concluded that:

I. Material is the major contributory factor for the successful repair of a composite resin restoration.
II. Knowing the composition of the aged composite should allow the operator to achieve reliable bond strength with the new composite resin.
III. MC-based composites showed better reparability than SIL-based composites.
IV. Regarding the surface treatment, no statistical differences in repair bond strength were evident.

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