Resin Composite Restorations: Effect of Energy Density on Properties And Marginal Integrity

by

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A Thesis

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ABSTRACT

Resin composite (Z250, 3M, St Paul, MN) was bonded in bulk with various degrees of cure into Class 2 preparations in extracted human molar teeth. Two control groups consisted of a highly-cured incrementally-placed group and a non-fatigued lowcure group. The restorations were subjected to 1000 thermal-cycles and 500,000 fatigue cycles from 18 to 85 Newtons using a stainless-steel sphere. Marginal integrity was evaluated using visual rating (ridit analysis) and microleakage. The degree of conversion had a significant effect on marginal defects but not on microleakage. Water had a significant dissolving effect on gingival margin integrity at very low degrees of conversion. Gingival marginal defects were maximized at 25% of maximum conversion and 3% of maximum hardness. There was no overall significant effect of thermalmechanical stressing on marginal defects or microleakage. A recommended lower limit of gingival margin acceptability in a bulk-filled resin composite restoration was created by 80% of maximum conversion, 73% of maximum hardness and approximately 70% of maximum flexural strength and modulus in the gingival marginal area.

INTRODUCTION

New light activation protocols (PAC lights and lasers) and new composites (packables) claim the advantage of shorter exposure times and bulk curing. The negative consequences of these approaches would be a composite with regions of lower properties at the base of the restoration. Laboratory research shows that many of the suggested new protocols do not produce composites with maximum depth of cure or uniform properties. However, manufacturers and some clinicians claim clinical success. Current light-curing techniques may produce adequate marginal integrity though the properties of the base of these restorations are significantly less than the properties on the surface. The question is: "Are these lower properties clinically significant?" One danger is that repeated stressing of the interface between the base of the restoration and the tooth may deteriorate the marginal seal and integrity with time as a consequence of the "undercuring". In bonded restorations, would enough occlusal forces be transferred to the cavity walls allowing for an increase in marginal integrity that did not deteriorate under fatiguing?

<u>Hypothesis.</u>— There is some minimal extent of cure required by the base of the resin composite restoration that allows it to support the rest of the filling and maintain its marginal seal under thermal and mechanical stress conditions.

Utilizing various photo-curing energy densities, a pilot study was used to determine the Knoop hardness and degree of conversion at incremental levels occlusal-gingivally throughout the depth of the resin composite in a 5 millimeter long sample created from an extracted tooth template.

Based on the pilot study, select energy densities were used to achieve variable degrees of gingival cure for resin composite restorations in prepared teeth. These restorations were subjected to thermal-mechanical stress to evaluate the effect on the gingival margins using scanning electron microscopy, stereomicroscopy and microleakage.

Additionally, bar-shaped specimens (2x2x25 mm) were tested in three-point bending to measure flexure strength and modulus. This data was correlated with the marginal breakdown data. The hypothesis tested was that marginal leakage and breakdown, as evaluated by silver nitrate and scanning electron microscopy and stereomicroscopy would increase for composites with low degree of conversion and low elastic modulus at the gingival margins.

REVIEW OF THE LITERATURE

Resin Composites.--

Introduction.-- The introduction of resin composites was one of the greatest milestones in esthetic restorative dentistry. Introduced in the early 1960's by R. L. Bowen with the development of the Bis-GMA molecule, it satisfied an initial need for an aesthetic anterior restorative material – eventually replacing silicate cement and acrylic resins.³ Bowen began his experiments to reinforce epoxy resins with filler particles. However, epoxy resins were abandoned due to slow curing and discoloration. The monomer, Bis-GMA, prepared from bisphenol A and glycidyl methacrylate was bulky, but superior to the methyl methacrylate. It provided rapid hardening, less shrinkage and stronger and more rigid resins.⁴

With anterior resin composite restorations providing satisfactory performance, most research is now focused at developing a resin composite capable of replacing amalgam. Improvements in formulations, optimization of properties and the development of new placement techniques have made esthetic direct posterior restorations more commonplace and dependable. A concern with resin composite is polymerization shrinkage. A major step in solving this problem would be the development of a polymer that experiences only enough contraction on curing to offset any expansion from water absorption.

Composition.-- A composite is a material that consists of two or more components of metals, ceramics or polymers. Dental resin composites typically contain a mixture of a soft, organic resin matrix (polymer) and hard, inorganic filler particles

(ceramic). Other components are included to improve the efficacy of the combination and initiate polymerization. The resin matrix consists of monomers, an initiator system, stabilizers and pigments. The inorganic filler consists of particles such as glass, quartz and colloidal silica. The matrix and filler are bonded together with a coupling agent. The performance of resin composites is obviously dependent on these basic components.⁴ The current improvement of these materials has primarily focused on filler technology, but the resin monomers have remained largely unmodified. The majority of resin composites are still based on variations of Bis-phenol A glycidyl methacrylate (Bis-GMA).⁵

The most common monomers used are Bis-GMA, urethane dimethacrylate (UEDMA), and triethylene glycol dimethacrylate (TEGDMA). Bis-GMA is extremely viscous at room temperature due to hydrogen bonding by hydroxyl groups (-OH). Lower viscosity is obtained by mixing Bis-GMA with dimethacrylate monomers (TEGDMA) of lower molecular weight to facilitate the addition of fillers. Addition of the diluents allows greater degree of conversion and more extensive cross-linking to occur between chains – providing a matrix that is more resistant to solvents. However, this increased conversion and crosslinking increases the polymerization shrinkage. Resin composites undergo volumetric shrinkage of 1.9 to 7.1 percent. The shrinkage in the resin matrix results from the conversion of weak intermolecular attractions to primary covalent bonds. Polymerization shrinkage and the resultant stress can contribute to gap formation at the margins of restorations. The current goal of resin composite restorative materials remains to be the improvement or elimination of contraction stress – possibly through low or non-shrinking monomers. Recent research has focused on ring opening

spiro-orthocarbonates, epoxy-based resin systems and a new class of difunctional monomers called oxybismethacrylates which exhibit cyclopolymerization.⁵

Polymerization shrinkage can also be reduced by increasing the concentration of filler particles because the overall shrinkage depends on the amount of polymer matrix present.⁴ However, the modulus of elasticity of the resin composite is also increased at high filler levels contributing to higher polymerization stress.¹¹ Inclusion of filler particles into a resin matrix drastically improves the mechanical properties of the composite material. Improvement is seen in properties such as tensile and compressive strength, modulus of elasticity, abrasion resistance, radiopacity, esthetics and handling.⁴ As a general rule, the higher the filler loading, the higher the physical properties of the resin composite. Most current resin composites have filler loaded between 50 and 86 percent by weight and 35 to 71 percent by volume.¹² Percentage filler is best expressed by volume instead of weight because the mechanical properties of composites are mainly dictated by their filler volume fraction.¹³

The type of filler directly influences radiopacity and is typically accomplished through the inclusion of elements of high atomic number. Barium and strontium are the most common elements used in filler particles to increase radiopacity.¹⁴

Various classification systems for resin composites have developed through the years based on particle size.⁴ A very traditional system includes traditional, small particle, microfilled and hybrid filler particles.¹⁵

Originally, crystalline quartz was used as a filler for resin composites because of its availability, excellent optical properties, and chemical inertness. However, it proved to be extremely hard, a challenge to grind and difficult to polish with the potential

to abrade opposing tooth structure.⁴ The softer polymer would wear away easily, exposing the hard quartz particle, only to be plucked and perpetually roughen the surface.¹⁶ These traditional quartz particles were produced by grinding or milling and typically were quite large – average 8 to 12 microns in size.⁴

Microfills were developed to provide better esthetics and polishability. These tiny particles of silica are only 0.04 microns in diameter and are literally "born in fire" through a pyrolytic process. The large surface area of these filler particles demands much more resin matrix to wet the surface. This creates extremely high viscosity that limits the percentage filler content possible. In order to maximize filler loading and minimize viscosity, prepolymerized resin and microfiller is used. The heavily filled polymerized resin is ground into 30-65 micron particles and mixed with more resin and microfiller to provide a composite that is filled 30 to 50% by volume. A smoother surface can be produced due to the smaller size of the silica particles. However, mechanical properties such as strength and stiffness are generally inferior to larger quartz or glass filled composites because of the lower filler content. 16

Small particle composites were developed to combine the esthetics of microfills with the mechanical properties of traditional composites. The average filler size ranged from 1 to 5 microns.⁴

The most common filler today is barium glass with average particle size of 0.6 to 1 micron.⁴ A small amount of microfiller is added to improve handling and reduce stickiness.¹⁶ To incorporate a maximum amount of filler into a resin matrix, a distribution of particle sizes is necessary. These so-called hybrids are potentially superior because increased filler loading improves the stress transfer between particles in the composite.⁴

The current trend is to maximize filler loading and minimize filler size.¹⁶ The table below compares and contrasts various mechanical properties of these various resin composite types:⁴

Composite Resin Mechanical Properties

Property	Traditional	Microfilled	Small Particle	Hybrid
Compressive strength (MPa)	250-300	250-300	350-400	300-350
Tensile strength (MPa)	50-65	30-50	75-90	70-90
Elastic Modulus (GPa)	8-15	3-6	15-20	7-12
CTE	25-35	50-60	19-26	30-40
Knoop Hardness	55	5-30	50-60	50-60

Other classification systems have been developed over the years due to the difficulty in categorizing the newer complicated formulations of resin composites based on a wide range of filler sizes, loading and compositions. Willems proposed a system based on volume fraction in 1992.¹⁷ More recently, a newer re-classification based on particle size was introduced. The various groups included megafill – 0.5 to 2 millimeters; macrofill – 10 to 100 microns; midifill – 1 to 10 microns; minifill – 0.1 to 1 microns; microfill – 0.01 to 0.1 microns and nanofill – 0.005 to 0.01 microns. Most new systems are minifill hybrids with a new trend toward nanofillers.¹²

It is important that the filler particle be bonded to the resin matrix via a coupling agent to improve mechanical and physical properties. The most commonly used coupling agent is an organosilane such as gamma-methacryloxypropyltrimethoxy silane. The silane reduces hydrolytic breakdown and allows stress transfer between the filler and the matrix. The silane agent is a bifunctional molecule with a methacrylate group on one end

and a silanol group on the other. The methacrylate end undergoes addition polymerization with the composite resin and the silanol end bonds to the hydroxyl groups on the filler particle via a condensation reaction.⁴

Resin composites undergo addition polymerization by chemical, heat or light energy activation. In chemical activation, a tertiary amine activator reacts with a benzoyl peroxide initiator supplied as two separate pastes to produce free radicals. The radicals attack the carbon double bonds of the polymer to propagate the reaction. In heat activation, higher temperatures activate the initiator. The most popular method of curing resin composites is with light activation. Originally, ultraviolet light was used to initiate free radicals. Today, visible light provides greater depth of cure and safety. 19,20

Light activation is achieved with blue light that is absorbed by the photoinitiator, camphoroquinone in the 400 to 500 nanometer range. The excited initiator reacts with an amine activator to form free radicals and initiate the addition polymerization.⁴ Compared to chemical curing of direct restorative resin composites, visible light activated systems provide longer working time, reduced porosity, greater color stability and reduced waste of material. A possible disadvantage with visible light materials is limited depth of penetration by the curing light.¹⁸

An inhibitor is added to resin systems to rapidly scavenge propagating radicals and prevent spontaneous polymer chain formation from heat or light. The inhibitor delays the start of the reaction, but once begun, the reaction proceeds at a normal rate. A typical inhibitor is butylated hydroxytoluene.⁴

Different metal oxides are added to resin composites in tiny amounts to provide shading and opacity to match natural tooth color.¹⁸

<u>Polymerization Chemistry.--</u> Many types of dental resin composites utilize free radical polymerization with monomers containing double carbon bonds reacting to produce longer polymer chains of single carbon bonds. The reaction takes place in three stages: initiation, propagation and termination.

Initiation involves the production of highly reactive free radicals with either the use of heat, chemicals or light.

Heat activation is typically used in denture resins to initiate the heat sensitive benzoyl peroxide molecules. These benzoyl peroxide molecules split quite easily into two free radicals at temperatures between 50 and 100 degrees Centigrade, which initiate the polymerization of methyl methacrylate monomer.⁴

Chemical activation typically involves the mixing of two reactants that initiate the polymerization of self-cure dental resins. Benzoyl peroxide is again the usual initiator, but instead of heat energy, a tertiary amine activator catalyzes the split of the benzoyl peroxide molecule.⁴

Light activation is the most common method of initiating the polymerization process in resin composites for use in restorative dentistry. Initially, ultraviolet light activated resin composites were introduced as a welcomed replacement to the chemically cured resins. Utilizing a benzoin methyl ether photoinitiator, they offered command setting, no mixing and unlimited working time. The potential for retinal and skin damage ushered in the incorporation of visible light initiators by the early 1980s.²⁰

In visible light activated polymerization, camphoroquinone is most commonly used to induce photoinitiation.¹⁸ Camphoroquinone (CQ) requires a tertiary amine co-

initiator to photocure composite in a reasonable amount of time.²¹ CQ absorbs energy in the visible region of 400 to 500 nanometers with a peak at 468 nanometers. Photons associated with this frequency range will be absorbed by camphoroquinone, raising it from the ground state to an excited, but short-lived, activated triplet state. However, if the energy is removed or diminished, the complex may de-energize and return to the ground state. If the excited triplet manages to bump into an amine co-initiator, an exciplex forms, with the subsequent transfer of an electron and then a proton to CQ to create an aminoalkyl free radical capable of initiating polymerization.^{22,23}

Propagation is a chain reaction that results in the incorporation of hundreds of monomer units into the polymeric network. The rigidity of Bis-GMA molecules reduces its ability to rotate during polymerization and to participate efficiently in the polymerization process. The complete conversion of the methacrylate double bonds is eventually hindered by the reduction of diffusion rate of free radicals and unreacted methacrylate molecules as the polymer undergoes vitrification.⁴ Therefore, only 50 to 75% of the methacrylate groups convert into polymer.²⁴ The process continues with considerable velocity until terminated either by direct coupling or by the exchange of a hydrogen atom from one growing chain to another.⁴ The extent of the reaction is important in developing physical and mechanical properties of the composite.¹⁶

Light-Curing .--

<u>Light generation</u>.-- Light is a form of electromagnetic radiation. Other forms of electromagnetic radiation include radio waves, microwaves, infrared radiation, ultraviolet rays, X-rays, and gamma rays. All of these, known collectively as the electromagnetic

spectrum, are fundamentally similar in that they move at 186,000 miles per second - the speed of light. The only difference between them is their wavelength, which is directly related to the amount of energy the waves carry. Shorter wavelengths have higher energy. The rainbow of colors that we see in visible light represents only a very small portion of the electromagnetic spectrum. On one end of the spectrum are radio waves with wavelengths billions of times longer than those of visible light. On the other end of the spectrum are gamma rays. These have wavelengths millions of times smaller than those of visible light. Visible light is the portion of the electromagnetic spectrum with wavelengths between 400 to 700 nanometers.²⁵

Quartz-tungsten-halogen lamps are integrated into most dental curing lights in use today. The principle output from these lamps is infrared energy with the generation of high heat. An infrared filter is used to reduce the heat energy to the oral structures. Next, a band-pass filter provides further restriction of visible light to the narrower spectrum of the photoinitiator. Finally, the dichroic filter, a silver coated surface passes infrared energy out the back and reflects and focuses the light forward to provide a focal area of energy at defined distance. Ultimately, 99.5% of the original radiation is eliminated. Due to the high operating temperatures, the halogen bulbs have a limited efficient lifetime. The reflector, bulb and filters can break down over time, reducing the curing effectiveness.

Less noticeable on the surface to the operator, a decreased light intensity could dramatically affect the cure in deeper areas of the restoration. Therefore, curing lights should be periodically tested with a dental radiometer.²⁷ However, recent research suggests that the accuracy of the commercial radiometers vary widely depending on the

diameter of the curing tip. Commercial radiometers have fixed apertures, while an industrial radiometer measures all light intensity from the curing tip, regardless of the diameter. A study in dental offices revealed that 46% of the halogen lights provided inadequate intensity to sufficiently cure resin composites. 29

Increased levels of light can be developed within the composite depths if the intensity of the curing source is increased. With curing procedures considered highly time consuming, the demand for expensive high energy plasma-arc and laser light sources has increased.³⁰ However, polymerization stress and contraction gaps increase as the mechanical properties are maximized and perhaps, as the rate of polymerization is increased.³¹

Plasma-arc (PAC) lights generate a high voltage pulse that creates hot plasma between two electrodes in a Xenon filled bulb.³² The irradiance of up to 2400 mW/cm² is much higher than typical halogen bulbs, but the PAC light generates very high heat with an inefficient emission spectrum similar to halogen bulbs.²⁵

Light emitted from an argon laser is very different from that emitted from the halogen or PAC lights. The photons produced are coherent and do not diverge, therefore they concentrate more photons of specific frequency into a tiny area. The laser has a power of only 250 mW, but the energy is restricted to only a few wavelengths. With very little infrared output, unwanted heat is minimized. If the 250 mW is transmitted through a 50 micron fiber it results in an irradiance of more than 12 million mW/cm² However, the tiny spot size of only 50 micrometers makes clinical application more tedious. Some systems utilize diffusers that increase the spot size, however, the irradiance will be

reduced. Other clinicians will use the tiny fiber itself, placing it directly on the tooth until it actually appears to glow.²⁵

(LED) use special semiconductors the for diodes emitting Light electroluminescence of light rather than a hot filament found in halogen lights providing a longer life span, consistent output and lower power consumption.³³ No significant ultraviolet or infrared light is emitted thereby reducing heat and minimizing the need for a noisy fan. Since the energy is clearly defined by the semiconductor, most of the light emitted is concentrated in a narrow band around 470 nanometers which is ideally suited for composite resins that use the photoinitiator camphoroquinone.³⁴ Initial claims of greater efficiency in curing resin composites have yet to be unequivocally established in vitro. 33-35 More laboratory and clinical data is needed to establish the significance of any increased efficiency and determine the overall efficacy of this potentially promising technique in light curing. 36

<u>Light-Restorative Interaction</u>.—

<u>Light transmission.</u>— Light is reflected, scattered and absorbed as it passes into the polymerizing resin composite.³⁷ There are many factors that may alter the transmission of light, such as filler type and size, composite shade, and refractive index.

Light scattering may be related to filler particle size in resin composite. Kawaguchi et.al. showed that microfilled composites had a lower transmission coefficient and depth of cure than hybrid and small particle composites.³⁸ One theory suggests that light scattering is maximized when the filler particle size is one-half the wavelength of the activation light.³⁹ One-half of the peak wavelength of the 450 to 500 nanometers

typically emitted from halogen curing lights would be 250 nanometers or 0.250 microns. However, microfill composites are, on average, only 0.04 microns in diameter. But at least one study found that microfills tend to clump in sizes that may closely approximate these critical sizes and increase light scattering and reduce depth of cure relative to hybrid composites. Another study found resin composites with larger filler particles were capable of showing good transmission coefficient and cure depth.

The content of pigment, or shade, appears to influence light scattering in resin composites. McCabe et. al. showed a strong correlation between depth of cure and light attenuation for various shades of microfilled composite.⁴⁰ However, Ferracane et. al. suggested that the depth of cure may be less dependent upon shade than upon other factors, such as translucency.⁴²

Transmission of light is facilitated through composite when the refractive index is closely matched between the filler particles and the resin matrix. Refractive index is the ratio of the velocity of propagation of an electromagnetic wave (light) in vacuum to its velocity in the medium (composite). However, this coordination produces a translucent composite. Therefore, a compromise is needed to provide some obvious esthetic qualities. Some mismatch in refractive index between the filler particle and the resin matrix is needed to provide some opacity. 43

<u>Light absorption</u>.-- There are many factors that may be altered with light absorption, such as depth of cure, degree of conversion, and biocompatibility.

Numerous studies have suggested that depth of cure is affected by light intensity, spectral distribution and exposure time. 43,44 The more intense the light intensity, the

more photons available. With more photons, more camphoroquinone molecules are raised to the excited state, react with the amine and form free radicals. At the top surface, polymerization proceeds quickly and more completely. However, deeper in the composite, attenuation of light leads to a potential gradation of cure within the depths of the material and is responsible for what has become known as "depth of cure". To compensate for this decreased potential, the duration of exposure can be increased, providing enhanced opportunity for the creation of a free radical. A study by Rueggeberg and others found that at depths greater than 2 millimeters, poorer cure may result and polymerization is very susceptible to changes in light intensity and exposure duration. Routine exposure times of 60 seconds are recommended using light-source intensities of at least 400 mW/cm². 46

The maximum degree of conversion of monomer into polymer of resin composites is between 50 and 80%. 47 However, only a small amount of unreacted monomer actually remains. Most of the remaining unreacted double carbon bonds are bonded to the polymer chain. Therefore, less than 10% of the available methacrylate is extractable, thereby significantly reducing biological effects. 48 Most of the unreacted monomers are believed to be TEGDMA molecules. 49 However, with decreased degree of conversion, there is greater residual monomer available to be leached. 48,50 And as the percentage of monomer conversion decreases, the potential for cellular toxicity increases. 51 Several *in vivo* studies have shown that complete or incompletely polymerized resin composites cause little pulpal irritation if the restorations are sealed. 52,53

In addition to biocompatibility issues, less degree of conversion may reduce the mechanical properties of resin composites. There appears to be a good correlation between increasing degree of conversion and increasing hardness,⁵⁴ fracture toughness ⁵⁵ and abrasive wear resistance.⁵⁶

<u>Restorative Response</u>.-- The prognosis of the restoration may depend on how the resin composite is placed and how it is light activated.

One disadvantage of resin composite is the polymerization shrinkage that arises during curing and the potential for subsequent marginal gap formation. Marginal gap formation may facilitate secondary caries and post-operative sensitivity especially in posterior resin composite restorations with greater technique sensitivity.

Various studies have advocated incremental instead of bulk placement and curing for resin composite to decrease the effects of polymerization shrinkage and to provide complete polymerization in deeper cavities. 2,58,64-65 Incremental placement may also reduce the configuration factor – the ratio of bonded to unbonded surfaces – and reduce the stress by making more resin available for flow. The superiority of the incremental technique has not been totally proven. Tjan and others found that when resin composite was cured in bulk from the occlusal, there was no difference in marginal gap formation at the dentin margin compared to incremental placement. Another study found no significant difference between bulk and incremental cure and its effect on fluid permeability in dentin under class 1 resin composite restorations. 68

Another approach to stress reduction is the application of various liners. Glass ionomer liners may provide some advantages when used with resin composite

restorations. The setting reaction is much slower and less polymerization stress may result.⁶⁹ Less bulk of resin composite is necessary to fill the lined restoration, thereby reducing the amount available for shrinkage. Resin-modified glass-ionomer restorative materials have been placed in areas of the preparation with dentinal margins in an "open-sandwich" technique to reduce marginal leakage.⁷⁰ However, the use of glass ionomer liners has decreased with the improvement in efficacy and ease of use of resin adhesive bonding agents.⁶³

The use of a relatively thick layer of resin adhesive to serve as a flexible intermediary between the stiff dentinal cavity wall and the shrinking restorative resin composite has been proposed. This "elastic bonding theory" proposes that the stretching of a 100 micron thick layer may provide enough elasticity to reduce the polymerization contraction stresses of the restorative resin composite to below dentin bond strengths. 11,63,71-72

Soft-start polymerization is a method recently advocated to reduce polymerization stresses. During early polymerization, the composite cross-linking network is relatively weak - allowing flow fairly easily to accommodate stresses and prevent damage to adhesive bonds. With further polymerization, contraction and flow decrease, while stiffness and stress increase. This can cause adhesive failure. The bond strength must exceed the contraction stress to provide a stable marginal adaptation. The resin-enamel bond interface may survive the shrinkage, but the weaker resin-dentin interface may not. Soft-start polymerization proposes that a slower rate of conversion will allow better flow of resin with a decrease in contraction stress. However, the final mechanical properties must be obtained in a reasonable amount of chair time. Three different types

of techniques are currently advocated. "Stepped" uses a short, low-level initial burst followed by full intensity. "Ramped" uses a gradual increase from an initial low level. And "pulsed" uses a short low level burst, a delay for polishing followed by full intensity. Although the majority of studies suggest that soft-start polymerization may be statistically beneficial *in vitro*, 74-81 further research is necessary to see if any clinically significant differences can be found.

Assessment of Polymerization.— The assessment of the effectiveness of polymerization may be done directly or indirectly. Indirect methods include surface hardness, ⁸² optical, ⁸³ and scraping. ⁸⁴ Direct methods include laser Raman spectroscopy ⁸⁵ and infrared spectroscopy. ⁸⁶ DeWald and Ferracane compared four modes of evaluating depth of cure of light-activated composites and found that the optical and scraping methods correlated well, but both over-estimated depth of cure when compared to hardness and degree of conversion values. Degree of conversion was the most sensitive testing mode for evaluating depth of cure. ⁸⁷ A good correlation can be found between hardness and relative degree of conversion for a specific composite at different time periods or under variable conditions. ⁸² Hardness testing appears to be a popular method of assessing effectiveness of polymerization because of its simplicity and reasonable correlation with other methods. ⁸⁷

Johnston et. al. suggested that the depth of cure may be defined as the level at which the hardness value is equivalent to at least 90% of the hardness at the top of the composite. 88 Others have suggested that this top-to-bottom gradient should not exceed 20% 89 and should be considered a realistic measure of depth of cure. 90 Research is

needed to determine the minimal extent of cure required by the base of the resin composite restoration that allows it to support the rest of the filling and maintain its marginal seal under thermal and mechanical stress conditions. The marginal integrity of resin composites has been routinely examined *in vitro* with marginal gap and microleakage evaluations.

Assessment of Marginal Integrity.— Ideally, evaluation of restorative margins should be repeatable, sensitive enough to discriminate between variables, recognize marginal discrepancies easily, and require little evaluator training. One of the first measures for quality assessment of restorations was the Ryge criteria. Popular as a clinical assessment, the procedure cannot detect small differences between various materials or variables. Scanning electron microscopy (SEM) provides a high magnification inspection of the marginal integrity of the restoration. Disadvantages include its potential of distortion through desiccation, lack of penetration, the destructive nature of the process and difficulty in quantifying the results. However, epoxy replicas may be used to avoid artifacts and provide observations over time. Researchers have used the SEM to measure marginal gap formation. Common *in vitro* methods to characterize marginal quality include gap or defect size, ridit analysis, ranking and microleakage.

Interfacial gaps may provide a route for bacteria to the pulp, whereas discrepancies such as marginal ditching or material defects create a site for plaque accumulation and potential secondary caries. Determining gap size involves measurement of the largest gap, 8 average gap, 9 or percent gapped. Using the

measuring capabilities of the SEM, Roulet assigned the largest gap measurement to one of five classes on the basis of gap size: 0; 0<30 microns; >30<50 microns; >50<100 microns and >100 microns<150 microns. Lutz and Krejci determined the percentage of continuous margins by digitizing the restorations using a three dimensional scanner and superimposing before and after images using a custom-made software program. 100

The ridit analysis¹⁰¹ in marginal restoration evaluation uses a numbered scale of restoration images that exhibit progressively increasing degrees of marginal defects. The procedure involves assigning to the restoration being evaluated, the number of the scale image to which it is most similar with respect to the extent of marginal defect. A ranking system described by Osborne and others consists of assigning the entire margin of each restoration in a rank of descending order from the one showing the best margin to the one showing the worst. ¹⁰²

Images may be created by enlargement of scanned computer images, 100 photographic film, 103 or SEM photographs of the entire margin. 104 Using enlarged images or photographs may be labor intensive, time consuming and produce pictures of inconsistent quality. 92 Although useful for gross marginal defects, 103 it may not be possible, in some cases, to see or estimate the width of micron-sized marginal gaps to any degree of accuracy. 92

Microleakage is defined as the clinically undetectable passage of fluids, bacteria, molecules or ions between the walls of the preparation and the restorative material. Microleakage is detected indirectly with the use of dyes, chemical tracers, radioactive isotopes, air pressure, bacteria, and artificial caries. 95

One of the most common methods of detecting leakage is the use of organic dyes or chemical tracers. There has been a wide variation in the choice, size, concentration and duration of dyes or chemical tracers. A chemical tracer, silver nitrate, is considered a severe test because of the very tiny size of the silver ion – much smaller than common bacteria. The sample is typically immersed in a fifty percent solution of silver nitrate and later developed with a photographic solution. 105

Another technique for evaluating microleakage is the use of radioisotopes. The most popular method is immersion in Ca⁴⁵. The sample is rinsed, sectioned and applied to photographic film. ¹⁰⁶

The use of bacteria to study microleakage may be the most clinically relevant. However, the relatively larger size of bacteria, 0.5 –1 micrometers, may not account for other detrimental products such as toxins. A somewhat qualitative test exists where the sectioned or shaved samples are cultured in broth to note the presence or absence of bacteria. 95

A more quantifiable method for leakage is the use of air pressure. The release of air bubbles is examined from a submerged sample, which does not need to be sectioned or destroyed. This gives an added advantage of leakage evaluation over time. 107

Bacterial cultures or an acidified-gel are used to produce artificial carious lesions. The acidified-gel technique has the advantage of expediency over bacterial induced lesions. Lesions produced are examined with polarized light and an outer and cavity wall lesion is described. 108

Many of these techniques - dyes, bacteria, chemical and radioactive tracers - necessitate the destruction of the sample, making it difficult to study the margins over

time. 109 Most studies section the tooth and restoration with a single cut through the center, allowing visualization of interfaces and dentinal tubules. Three dimensional techniques are more representative of penetration, but are tedious and extremely time consuming. The use of multiple sections is perhaps more practical. 110

Most researchers use a subjective ordinal ranking system to score the leakage. The ordinal scale has the advantage of relating relevant landmarks such as dentin-enamel junction and axial walls. However, non-parametric statistics must be used. Others have recently tried more objective techniques by measuring the microleakage directly restoration interface length and through the use of imaging software. Although more time consuming, continuous measurements allow greater precision and the use of more robust parametric statistics.

Many studies examining marginal integrity include thermocycling and load cycling in the design of the experiment. 117-119 Thermocycling is simply a way of applying stress to the restorative interface at the limits of temperature found intra-orally. 120 One researcher found a direct correlation between the degree of mismatch in the coefficient of thermal expansion between various restorative materials and tooth structure and the extent of microleakage. 121 Thermal-cycling was found to be detrimental to bond strengths of two older generation dentin adhesive agents 122-123 and the microleakage of bonded amalgam restorations. 124 However, several studies suggest that thermocycling resin composite restorations may not significantly affect microleakage. 124,125 Although resin composites have high coefficients of thermal expansion relative to tooth structure, they are relatively good insulators. Harper and others theorized with their clinical

measurement of thermal diffusion through restorations of various materials that the relatively low rate of thermal change beneath resin composite restorations might reduce the significant effect of thermal expansion. The significance of thermocycling resin composite restorations remains questionable and perhaps the procedure is unnecessary.

There appears to be little consensus on temperatures, dwell time and number of cycles among research studies involving microleakage. Investigators have used temperature ranges from 15 to 45°C, 4 to 60°C and 5 to 55°C. Dwell times have varied from 10, 15, 30, 60 and 120 seconds. Shorter dwell times are more clinically relevant, however, longer dwell times may stress the interface greater and accelerate the evaluation process. One study concluded that the depth of microleakage was correlated with the thermal conductivity of the restorative material. Composite restorations, with low thermal conductivity compared to amalgam appeared to be unaffected by differences in dwell time. Also, microleakage tests on composite restorations should be delayed 24 hours to permit the potential for water absorption. The number of thermocycles recommended has ranged from 1 to 2500. However, the stresses from thermocycling may act quickly to produce microleakage thereby reducing the need for extended cycles.

Water is absorbed as resin composite restorations are placed intraorally and may lead to hydrolytic breakdown of the silane coupling and filler particles, reducing the strength of the material. Water decreases the elastic modulus, fracture strength, and flexure fatigue limits. Water has a plasticizing effect on the matrix. Debonded fillers may act as stress concentrators, significantly increasing the number of potential crack growth sites. The breakdown can contribute to lower wear and abrasion resistance.

However, the expansion of resin composite associated with the water uptake could counteract some setting contraction. Studies show that microfilled composites absorb more water than hybrid composites. ¹³¹ Most resin composites require 4 days to show the majority of expansion and 7 days to reach equilibrium. ¹⁸

Researchers have proposed that mechanical loading of teeth may temporarily or permanently cause gap formation in restorations 119 and the combination of thermal and mechanical loading may simulate intra-oral conditions more accurately. 132 However. studies have shown conflicting results on load cycling, reinforcing the need for careful evaluation and interpretation due to differences in experimental designs. 100,116,133-142 Several researchers found no effect on microleakage from load cycling of resin composite restorations. 125,133-138 However, a few authors did find an effect from load cycling 116,134,140 and a couple of investigators found an effect from a simultaneous combination of thermocycling and loading. 100,139 The type of restoration, either Class 2 or Class 5, did not appear to have any effect on the conclusions reached in these studies. 100,116,133-142 Many studies used a combination of cyclic loading while the tooth soaked in dye solution to potentially help detect transient gaps formed during the loading. 116,133,139-141 Again, this combination did not appear to have any effect with some studies showing increased microleakage 116,140 and some no increase. 133,136

Different materials, bonding systems and types and location of load were used. Most studies used axial compressive loading to stress the restorations by using a stainless steel ball or rod that rests on the buccal and lingual inclines of the occlusal surface. Very few studies stress the actual restorative material itself. Very few studies and others loaded the buccal aspect of the

buccal cusp on teeth with class 5 restorations and found an effect of load cycling. ¹³⁴ While Hakimeh and others alternated the loading of the buccal and lingual inclines of the occlusal surface and found no effect on class 5 componer restorations. ¹³⁵

All studies used varying load levels and various cycles and cycling rates. 100,116,133-142 Some investigators oscillated cycles between a low of just a few Newtons to a maximum of 170 Newtons. 100,116,133-135,137,140 Others reported just a maximum load. 136-139 Number of cycles varied widely from a low of 1000 cycles 133 to a high of 1.2 million. 100 Cycling rate varied from a low of 1 Hertz 116,134,140 to a high of 5 Hertz. 137

Admittedly, however, there appears to be a wide variance in the literature as to the magnitude of actual clinical masticatory forces with an estimated range of 9 to 180 Newtons. The duration of the occlusal forces is estimated to be from 0.25 to 0.33 seconds or 3 to 4 Hertz. Dietschi et al, loaded his inlay samples with a stainless steel cusp at 80 Newtons and 1.5 Hertz for 250,000 cycles and related this to approximately one year of clinical use. 146-148

Behavior of resin composites *in vivo* is complex and is not clearly defined and is often described in terms of wear, marginal breakdown or fatigue. Fatigue is a major factor affecting the life expectancy of resin composite restorations. ¹⁴⁹ and is due to progressive fracture under repeated loading. ¹⁸ Several mechanisms may participate in fatigue-induced damage of composite materials including matrix cracking, multidirectional cracking, and filler debonding. ¹⁵⁰ Fatigue in resin composites largely depends on the materials properties, especially filler loading.

Higher filled hybrid resin composites deform little under function. Filler particles act to arrest or retard crack propagation through the matrix resulting in more localized

destruction. Under function, however, cracks can propagate, weaken the matrix and favor further crack growth resulting in a worn surface at the occlusal contact area. 149

Microfilled composites with a relatively lower filler content, lower modulus of elasticity and a weaker interface between resin matrix and prepolymerized filler particles have greater fatigue sensitivity and are more prone to mechanical deformation. Initially, while under function, cracks propagate in the subsurface area. Over the first two years, while under continued stress, the matrix is weakened as the cracks begin to connect. Eventually, after four or five years, the subsurface cracks interconnect and the restoration fails catastrophically and often, surprisingly.¹⁴⁹

There appears to be a need for more standardized testing procedures of fatigue to gain insight into the complex behavior of resin composite restorations. ¹⁴⁹ Currently, specimens may be cyclically tested in tension-tension, flexure, torsion, shear, or compression. ¹⁵⁰ Contact fatigue is used to study the wear process by cyclic loading of an indenter into the surface of the material. A study by Htang ¹⁵¹ and others found the existence of an optimal or critical filler level in resin composites in fatigue endurance.

Resin composites with low (<60% wgt) or high filler content (>80% wgt) were significantly lower in fatigue resistance. A higher susceptibility to crack development in very highly filled composites under impact stresses is probably due to the brittle nature of filler particles with fractures occurring through the filler particles as well as the matrix. The dynamic fatigue test is useful for determining crack velocity parameters. Finally, with a stair-case technique, the material is cycled at a preset stress level and cycles. If the specimen fails, a lower stress is chosen, if the specimen survives, a higher stress level is

used on the next specimen. 152

OVERVIEW

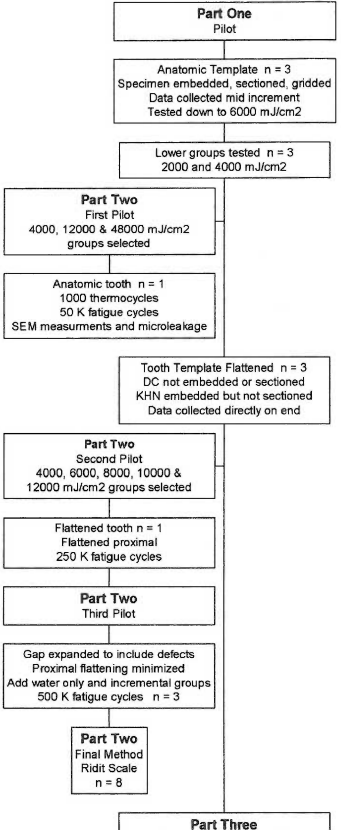
The following is an overview of this project. The study is presented in three separate parts – effects of energy density on hardness and degree of conversion, effects of degree of conversion on marginal integrity and effects of degree of conversion on flexural strength and modulus.

The first part details the materials and methods, results and discussion of an initial pilot study to determine the Knoop hardness and degree of conversion at incremental levels occlusal-gingivally throughout the depth of the resin composite in a 5 millimeter long sample created from an extracted tooth template. As can be seen in the discussion of this pilot study, there were many changes in the methods that spanned the whole project. But, the final procedure that was utilized provided the necessary information to select the various energy densities needed to achieve variable degrees of gingival cure for Z250 resin composite restorations in prepared teeth for the final method in part two of the study.

The second part details the materials and methods, results and discussion of the primary objective – to determine the minimal extent of cure required by the base of the resin composite restoration that allows it to support the rest of the filling and maintain its marginal seal under thermal and mechanical stress conditions. This section was broken up into three pilots and a final method. Again, various techniques were modified before a final version was agreed upon. Please note that the energy density groups selected changed with knowledge gained concurrently in part one of the study.

The third part details the materials and methods, results and discussion of a supplementary goal of determining mechanical properties of the composite at the marginal area at various degrees of conversion. See overview chart on next page.

Overview



Composite Bars n = 5
Flexure Strength and Modulus

PART ONE

PILOT STUDY

MATERIALS AND METHODS

Effects of Energy Density on Hardness and Degree of Conversion.-- The first objective was to determine which energy densities would produce sequentially decreasing

degrees of gingival cure and hardness in a 5 millimeter long resin composite restoration. One extracted human molar, stored in 0.525% NaOCl served as a reusable template. A tapered Class II slot cavity preparation (Figure 1) was created with a #57 carbide bur and high-speed handpiece with water coolant with the following dimensions: buccolingual (occlusal) – 4.5 mm; buccolingual (gingival) –



Figure 1: Re-usable template (anatomic)

4.0 mm; mesiodistal (gingival) 1.5 mm; occlusalgingival height of 5 mm. Composite specimens were created as outlined below. The tooth was stored in 37 degree Centigrade tap water while not in use.

A thin layer of Bis-GMA and TEGDMA (50% each by weight) containing no polymerization promoters was placed in the cavity preparation to act as a lubricant and to

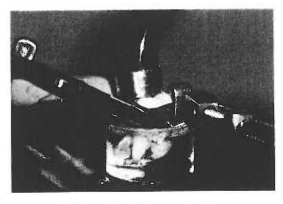
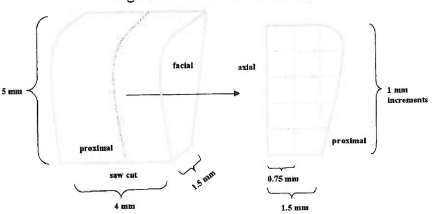


Figure 2: Mounting jig

simulate the layer of dentin bonding. A metal matrix band was placed on the tooth and held in place with a hemostat. A minifill resin composite was placed in bulk (Z250, Shade A-2, 3M, St. Paul, MN) using a resin composite syringe system (CR



tubes, Centrix, Inc, Shelton, CT) until the preparation was full. Large condensers were used to smooth the occlusal portion. The composite was cured from the occlusal with a curing light (VIP, Bisco, Schaumburg, IL) after utilizing its internal calibration feature. The light output of the curing light was confirmed using a power meter (Power Max 5200, Molectron, Portland, OR). A 10.25 millimeter curing tip was utilized and held in place with a mounting jig (Figure 2). The light tip rested on the occlusal cusp tips and a blue ink line on the occlusal surface helped to reduce variation in angulations and The distance between the light guide and the composite surface was position. approximately 0.75 millimeters. The approximate center of the light guide was over the center of the occlusal composite surface. The power density of the curing light was 600 mW/cm² and the time was varied between 10 and 100 seconds (10, 20, 40, 60, 80 and 100 seconds) to produce the various levels of cure at the gingival margin. Three specimens were made per energy density. The specimens were removed from the tooth template and stored dry throughout the entire testing process in a black film canister when not handled. The same day, the specimens were suspended in acrylic rings using a wax sprue and embedded in epoxy resin (Buehler, Lake Bluff, IL) and allowed to cure overnight in a dark box container to eliminate light exposure. The next day, the specimens were sectioned using a 0.3 millimeter thick diamond saw (Isomet, Buehler, Lake Bluff, IL) under a water coolant. The lights were dimmed in the room to reduce exposure. One saw cut was made through the specimen in the approximate center from the buccal and lingual surfaces respectfully. The test surface became the cut inner surface of the section (Figure 3) and was polished using 5 micron aluminum oxide powder (Buehler, Lake Bluff, IL) and de-ionized water on a 8 inch nylon polishing cloth (Buehler, Lake Bluff, IL) for 30 seconds.

A grid area was inscribed on the surface with a pencil to facilitate testing (Figure 4). Measurements were taken within the center of each increment at 24 hours for Knoop

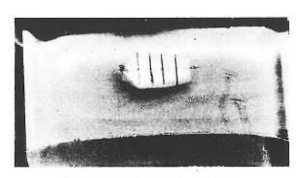


Figure 4: Gridded specimen

hardness and between 3 and 7 days for degree of conversion. The gridded specimen was tested at 2 points in the mesiodistal direction in 0.75 millimeter increments and 1 millimeter in the occlusalgingival direction for the entire

length of the specimen for Knoop hardness (KHN) (Kentron Microhardness, Torsion Balance Co., Clifton, NJ) Due to the heavy time expenditure, only the occlusal and gingival increments were tested for degree of conversion (DC) using Fourier Transform Infrared Spectroscopy (FTIR) (DS20, Analect Instruments, Irvine, CA). Tiny chips of composite - 10 to 30 microns in thickness - to be placed in the FTIR were removed from the center of each increment using the tip of a razor blade after completing the Knoop hardness test. Knoop hardness values were determined using the following formula: $KHN = L/l^2 \times Cp$ where L is the load in kilograms and l is the length of the indentation in millimeters and Cp is the constant 0.007028. A 200 gram load was used with a dwell time of 10 seconds.

The degree of conversion of the Z250 resin composite was determined using FTIR analysis. The intensities of the carbon double bond (C=C) absorbance peak at 1637.3 cm⁻¹ and the aromatic (C...C) reference peak at 1608.3 cm⁻¹ were measured with 30 scans at a resolution of 4.0 in transmission mode. The C...C peak originates from the aromatic rings in the Bis-GMA molecule and remains unchanged during the polymerization reaction. The ratio of the absorbance intensities of C=C/C...C were compared before and after polymerization using the following equation to determine the percent of reacted carbon double bonds or degree of conversion:

1 – [Abs (C=C) / Abs (C...C)] cured resin / [Abs (C=C) / Abs (C...C)] uncured resin X 100 See Appendix under Equipment for more details on FTIR and Knoop Hardness testing.

Data Analysis. Three specimens were created for each energy density level. Three hardness and two degree of conversion values were determined per axial and proximal increment examined per specimen and a mean calculated. The percentage of maximum KHN and DC was determined for the gingival regions. A two-way analysis of variance (ANOVA) was used to test the effect of energy density or location (axial vs proximal) on Knoop hardness in the 5 millimeter increment. The same tests were done for degree of conversion. Tukey's post-hoc tests were utilized to perform *a posteriori* pair-wise comparisons to determine which groups showed a significant difference (α =0.05). SPSS software (SPSS Inc, Chicago, IL) was used and the printout is provided in the Appendix under Part One. Also, paired student t-tests were used to compare axial with proximal increments for both Knoop hardness and degree of conversion. See tables in Appendix under Part 1.

RESULTS

Effects of Energy Density on Hardness and Degree of Conversion.— The mean of three hardness values per increment per specimen (Appendix Part 1) and the mean of two degree of conversion values per occlusal and gingival increment per specimen (Appendix Part 1) was determined and a percentage of maximum KHN and DC determined for the gingival regions. The highest value for KHN and DC was found to be 98.7 kg/mm² and 60.6% respectively and was therefore designated as the maximum KHN and DC. Paired t-tests revealed no significant difference between axial wall and proximal surface increments for all but a few of the paired increments found typically in the lower energy levels (Appendix, Part 1). Pooled versions of the tables combining axial with proximal values are shown below (Tables 1 and 2).

A linear regression analysis (R^2 =0.95) was performed using SPSS software relating Knoop hardness with degree of conversion at the gingival increment using pooled axial and proximal average values (Figure 5). Also, Knoop hardness (R^2 =0.96) or degree of conversion (R^2 =0.94) was plotted versus the log of energy density (Figure 6).

Table 1: Knoop Hardness in kg/mm² (anatomic template) n=6 max KHN =98.7

6000	mJ/cm ²		12000	mJ/cm ²		24000 mJ/cm ²				
	mean	st dev	•	mean	st dev		mean	st dev		
1 mm	89.4	4.8	1 mm	89.3	5.2	1 mm	90.2	5.0		
2	86.1	5.3	2	89.6	3.6	2	87.6	5.5		
3	78.5	5.7	3	84.8	2.7	3	85.8	6.1		
4	70.3	4.4	4	79.3	6.5	4	79.2	8.3		
5 mm	45.8	4.2	5 mm	64.6	9.2	5 mm	70.6	11.3		
% of max	46.4	4.3	% of max	65.3	9.4	% of max	71.5	11.6		

36000	mJ/cm ²		48000	mJ/cm ²		60000	mJ/cm ²	
	mean	st dev	mean st dev				mean	st dev
1 mm	90.2	4.6	1 mm	91.6	2.4	1 mm	90.8	2.3
2	89.9	3.8	2	91.7	3.1	2	90.7	4.5
3	90,8	4.7	3	90.3	4.2	3	92.5	5.4
4	85.7	4.2	4	83.9	5.3	4	92.5	6.2
5 mm	76.4	12.0	5 mm	80.0	6.0	5 mm	88.7	2.3
% of max	77.3	12.1	% of max	80.9	6.1	% of max	89.8	2.3

6000	mJ/cm ²		12000	mJ/cm ²		24000	mJ/cm ²	
	mean	st dev		mean	st dev		mean	st dev
1 mm	53.7	2.7	1 mm	56.1	2.4	1 mm	57.3	2.2
2			2			2		
3			3			3		
4			4			4		
5 mm	40.0	2.2	5 mm	47.2	3.0	5 mm	49.2	3.3
% of max	66.0	3.6	% of max	77.9	4.9	% of max	81.2	5.5
	,	•	49000	m I/om²		60000	m I/om²	
	mJ/cm²		48000	mJ/cm ²		60000	mJ/cm²	
3 6000	mJ/cm²	st dev		mean	st dev		mean	st dev
36000	mJ/cm²		1 mm			1 mm		st dev
3 6000	mJ/cm²	st dev	1 mm	mean	st dev	1 mm	mean	1
3 6000	mJ/cm²	st dev	1 mm	mean	st dev	1 mm	mean	1
36000 1 mm 2	mJ/cm²	st dev	1 mm	mean	st dev	1 mm	mean	1
36000 1 mm 2 3	mJ/cm²	st dev	1 mm 2 3	mean	st dev	1 mm	mean	1

Figure 5: Knoop Hardness vs degree of conversion at the gingival increment

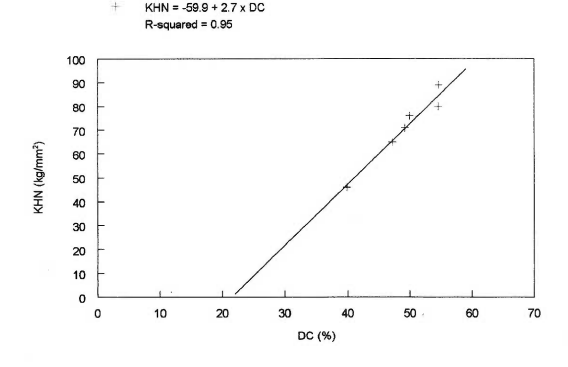
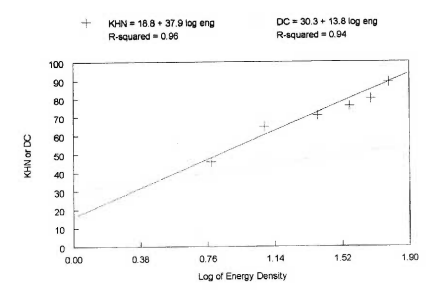


Figure 6: Knoop Hardness or degree of conversion vs log energy density at the gingival increment



A percentage of maximum Knoop hardness and degree of conversion were determined for the gingival increments. A two-way analysis of variance (ANOVA) was completed (Appendix, Part 1) using non-pooled axial and proximal average values (n=6) to test the effect of energy density or location (axial vs proximal) on Knoop hardness in the 5 millimeter increment. The same tests were done for degree of conversion. Significant differences in Knoop hardness and degree of conversion were found based on energy levels using a two-way ANOVA and Tukey's post hoc tests. (α=0.05). The percentages of maximum Knoop hardness and degree of conversion of the pooled gingival increment is shown below in Table 3. Letters (a-d) denote significant differences across rows. At least 36000 mJ/cm² was necessary to produce maximum gingival degree of conversion and Knoop hardness in a 5 millimeter deep restoration of Z250 resin composite.

Table 3: Percentages of maximum KHN and DC at the gingival increment for various energy densities n=6/group

a - d denotes significant differences across rows

	6000 mJ/cm ²	12000 mJ/cm ²	24000 mJ/cm ²	36000 mJ/cm ²	48000 mJ/cm ²	60000 mJ/cm ²
KHN	46.4 % a	65.3 % b	71,5 % b,c	77.3 % b,c,d	81.0 % c,d	89.8 % d
DC	66.0 % a	77.9 % b	81.2 % b	82.7 % b,c	90.2 % c	90.3 % c

DISCUSSION

Effects of Energy Density on Hardness and Degree of Conversion.— Although the two-way ANOVA showed no significant difference overall between axial wall and proximal surface increments at the 5 millimeter level, the individual paired T-tests did show a few differences at lower energy levels (Tables in Appendix under Part One). This variation could be due to the orientation of the light guide and the effect of tooth or band shadowing.

Regression analysis (Figure 5) revealed a significant correlation when relating Knoop hardness and degree of conversion (R² = 0.95). This agrees with Ferracane who found a similar relationship during the setting reaction of three unfilled dental resins.⁵⁴ However, the attainment of hardness relative to maximum hardness fell behind the degree of conversion of the carbon double bonds (Tables 1 and 2). This is possibly due to the late formation of networking links in the polymer chains. A greater increase in hardness relative to maximum hardness occurred with higher energy levels and subsequent increases in crosslinking.⁵⁴ Lag of hardness behind degree of conversion could also be due to the overestimation of percent cure at lower energy levels. The "overestimation" may be due to the sectioning and polishing that might cause a loss of free monomer.

Knoop hardness (R^2 = 0.96) or degree of conversion (R^2 = 0.94) at the gingival increment had a strong linear relationship to the log of the energy density. However, the KHN and DC values fell dramatically at the gingival increment below a threshold level of approximately 12000 mJ/cm².

The first objective was to determine which energy densities would produce sequentially decreasing degrees of gingival cure and hardness. However, having

examined the results of this first objective, it became obvious that the minimum energy density in the gingival increment had not been determined. Two new energy density groups were created lower than the 6000 mJ/cm². A 2000 mJ/cm² group (200mW/cm² x 10 secs) and a 4000 mJ/cm² (400mW/cm² x 10 secs) group were tested for hardness and degree of conversion. Three specimens were created for each energy density level. Three hardness and two degree of conversion values were determined per axial and proximal increment examined per specimen and a mean calculated (Tables in Appendix, Part 1). The percentage of maximum KHN and DC was determined for the gingival regions. A summarized version of each table is shown below in Table 4.

Table 4: Knoop Hardness and degree of conversion (new groups) n=6 max KHN = 98.7 max DC = 60.6

			KHN	(kg/mm²)		
2000	mJ/cm ²			4000	mJ/cm ²	
	mean	st dev			mean	st dev
1 mm	78.2	10.3		1 mm	86.4	5.2
2	83.0	8.8		2	79,7	3.6
3	71.1	10.4		3	72.0	2.7
4	47.3	8.3		4	65.2	6.5
5 mm	16.7	3.8		5 mm	43.5	9.2
% of max	17.0	3.8		% of max	44.0	7.9

			DC	(%)		
2000	mJ/cm ²			4000	mJ/cm ²	
	mean	st dev			mean	st dev
1 mm	52.7	4.6		1 mm	54.0	2.4
2				2		
3				3		
4				4		
5 mm	27.9	12.0		5 mm	36.1	6.0
% of max	46.0	8.5		% of max	59.1	6.0

Table 5: Selected groups for Part 2, 1st Pilot, based on max KHN or DC of gingival increment

% Gingival KHN/DC of max	Energy density (mJ/cm²)	Power x time (mW/cm ² x secs)
45/60	4000	400 x 10
65/80	12000	600 x 20
80/90	48000	600 x 80

The Knoop hardness of the gingival increment of the 2000 mJ/cm² group was undeterminable and soft to touch in one specimen and near the limits of the filar scale in the other two. Therefore, 4000 mJ/cm² was chosen as the lowest energy density. Table 5 shows the selected groups to be used in the first pilot of the second part of the study (Part 2) based on percent of maximum Knoop hardness or degree of conversion of the gingival increment.

Inspection of the initial data (Tables in Appendix under Part 1) raised certain questions concerning the possible inflation of the Knoop hardness and degree of conversion numbers. Various efforts were undertaken to review existing data and to implement improvements before further testing.

First, it became apparent that the variability in length and occlusal shape created from preparing an anatomic tooth could result in extreme inconsistency in degree of conversion in the gingival areas of the restoration, especially at the lower energy densities. A solution was to flatten the occlusal table to provide a uniform 5 millimeter depth and shape to the preparations of the teeth. Also, as noted earlier, variation between individual values in the axial/proximal regions, particularly for low energy conditions, suggested that the curing conditions were variable – possibly due to angulations of the

light guide or differences in surface reflectance. Flattening the tooth surface was expected to reduce the potential for light guide angulations.

Second, since the data collected per increment was only representative of the middle third of the increment, it did not represent the true values at the apical portions of that increment. At lower energy densities it became apparent that the Knoop hardness and degree of conversion values could decrease dramatically in the apical half millimeter in the gingival increment. Knowing the true conversion value of the absolute gingival marginal level would become critical in Part 2 of this study.

Third, concerns were raised over the possibility of heat generation causing an increase in conversion from embedding the specimens in epoxy prior to FTIR evaluation

and possible leaching of monomer from sectioning and polishing. Therefore, a new set of removable specimens was evaluated based on a flattened tooth template (Figure 7). A total of three specimens were created per group. These 5 millimeter long specimens were not sectioned, polished or embedded in epoxy, but were

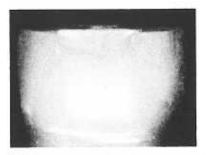


Figure 7: Flattened template

evaluated directly on the gingival and occlusal edges after lightly sanding away 0.1 millimeters to remove any unfilled resin and air inhibited layer. Testing was conducted 24 hours after dry storage at room temperature in a black film canister. Tiny chips of composite to be placed in the FTIR were removed from the occlusal or gingival area of the specimen in a darkened room. The average of three degree of conversion (DC) values per specimen was determined. Table 6 lists the occlusal and gingival degrees of conversion for the various energy densities. The percentage of maximum DC was

determined at 5 millimeters. A analysis of variance (ANOVA) was used to test the effect of energy density on DC at the gingival margin. SPSS software was used and the results are shown in the table and the printout is provided in the Appendix under Part 1.

Table 6: Occlusal and gingival degrees of conversion with percentage of maximum DC (58.2) of gingival increment using flattened template. (a - e denotes significant differences) n = 3

mJ/cm ²	occ DC	st dev	ging DC	st dev	% max
4000	54.1	1.0	14.3	5.1	24.5 a
6000	53.4	1.9	26.2	4.4	44.9 b
8000	55	1.2	33.4	0.2	57.3 b,c
12000	55.2	1.9	39.4	2.3	67.6 c,d
24000	56.2	1.9	46.5	0.9	80 d
Control 72000	57.4	0.7	56.4	1.6	97 e

It became apparent that a significant descending gradient of degree of conversion values was seen below the 12000 mJ/cm² level at the gingival margin providing greater opportunity to evaluate even more levels of conversion, especially in the critical lower levels. Therefore recommended groups for the second part of the study would eventually expand to include 4000, 6000, 8000, 10000, 12000 and 72000 (control) mJ/cm².

Finally, the Knoop hardness indenter utilized in this study displayed variability between operators and possibly overestimated actual Knoop hardness when evaluating resin composites. The poor contrast between the indentation and surrounding resin matrix may have provided a hindrance to reproducing filar length determinations. To reduce the potential for altering the hardness values in the poorly converted portions of

the specimens, a high level of polish was not attempted. A simple study found that Knoop hardness values varied a plus or minus 10 - 20 % between operators reading the same indentation. See Appendix under Part 1 for details. Care should be taken to use the same operator throughout the testing process to preserve the <u>relative</u> significance of the numbers between groups. The same study found a potential overestimation of Knoop hardness values by an average of 14. The ability to read the indentation may be enhanced by covering the composite test surface with graphite prior to testing to greatly increase contrast. Overall, the use of graphite appeared to increase the ability to read the actual lengths of the indentations and reduce the variability between operators and even within the same operator.

A new set of removable specimens was evaluated for Knoop hardness based on the same flattened tooth template (Figure 8). A total of three specimens were created per group. These 5

millimeter long specimens were mounted vertically in a square acrylic tube with epoxy resin with the

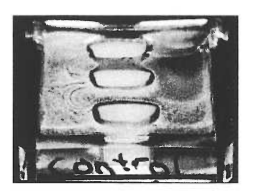


Figure 8: Vertically mounted specimens for KHN testing

occlusal and gingival portions exposed. These specimens were not sectioned, but evaluated directly on the gingival and occlusal edges after gently sanding away 0.1 millimeters to remove any unfilled resin and the air inhibited layer with 600 grit sand paper and polishing with 5 micron aluminum oxide paste for 30 seconds on a 8 inch nylon wheel. Testing was conducted 24 hours after dry storage at room temperature in a black film canister. Graphite from a mechanical pencil was rubbed on the lightly polished surface of the composite specimens before indentation. The average of three

Knoop hardness values per occlusal or gingival surface was determined. The following table lists the occlusal and gingival Knoop hardness values for selected 5 millimeter specimens. The percentage of maximum KHN was determined at 5 millimeters. A one-way analysis of variance (ANOVA) was used to test the effect of energy density on KHN at the gingival margin. SPSS software was used and the results are shown in Table 7 and the printout is provided in the Appendix (Part 1).

Table 7: Occlusal and gingival Knoop Hardness with percentage of maximum KHN (72.7 kg/mm^2) of gingival increment using flattened template (a-f denotes significant differences) n=3

mJ/cm ²	occ KHN	st dev	ging KHN	st dev	% of max
4000	69.3	2.3	2.1	3.6	2.8 a
6000	69.9	2.3	16.4	0.8	22.6 b
8000	69.1	3.1	28.5	4.5	39.2 c
12000	68	2.1	39.3	0.3	54.1 d
24000	68.8	1.7	52.9	0.3	72.8 e
Control 72000	71.7	1.1	70.2	1.2	96.5 f

A linear regression analysis ($R^2 = 0.99$) was performed using SPSS software relating Knoop hardness with degree of conversion at 5 millimeters (Figure 9 (1)). The lowest data point, representing 4000 mJ/cm² had a high standard deviation and was dropped from the analysis to obtain a better fit ($R^2 = 1.0$; Figure 9 (r)).

Also, Knoop hardness ($R^2 = 0.96$) or degree of conversion ($R^2 = 0.93$) was plotted versus the log of energy density (Figure 10). A more linear fit was seen in the lower energy densities. If the curve was extended to include higher energy densities, the curve would plateau and an exponential equation would be more appropriate.

Figure 9: Knoop Hardness vs degree of conversion (flattened template) at the gingival increment [(l) all data points; (r) dropped data point]

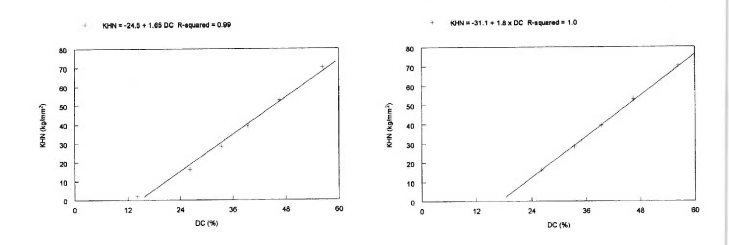
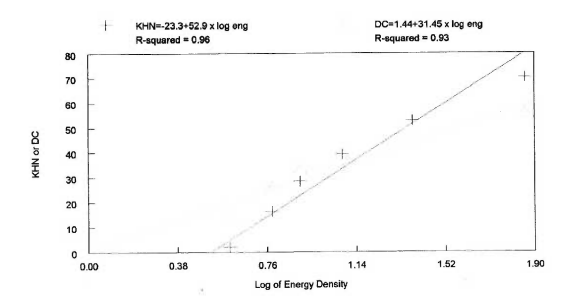


Figure 10: Knoop Hardness or degree of conversion vs log energy density (flattened template) at the gingival increment



PART TWO

MATERIALS AND METHODS

Effects of Degree of Conversion on Marginal Integrity. -- First Pilot

The first objective was to determine which energy densities would produce sequentially decreasing degrees of gingival cure and hardness. Next, selected energy densities from this data were used to bond resin composite specimens in prepared teeth to find the minimal extent of cure required by the base of the resin composite restoration that allowed it to support the rest of the filling and maintain its marginal seal under thermal and mechanical stress conditions.

Extracted human molars originally stored in 0.525% NaOCl were mounted in acrylic rings (14 mm x 22 mm) with epoxy resin (Buehler, Lake Bluff, IL). A Class II slot cavity preparation was created with a #57 carbide bur and high-speed handpiece with water coolant with the following dimensions: buccolingual (occlusal) – 4.5 mm; buccolingual (gingival) – 4.0 mm; mesiodistal (gingival) 1.5 mm; the gingival margin placed 0.5 mm into dentin. A 0.5 millimeter bevel was placed on the buccal and lingual margins. The teeth were stored in 37 degree C tap water while not in use.

A metal matrix band was placed on the tooth and held in place with a hemostat. The preparation was acid-etched for 15 seconds with 35% phosphoric acid, gently rinsed for 15 seconds with water from a three-way syringe, and lightly dried leaving the dentin moist. Two consecutive thin layers of Single Bond bonding agent (3M, St. Paul, MN) was placed in the cavity, gently thinned for 5 seconds and light cured for 10 seconds at 600 mW/cm² after internally calibrating as before (VIP, Bisco, Schaumburg, IL). A minifill composite resin was placed in bulk (Z250, Shade A-2, 3M, St. Paul, MN) using a

composite resin syringe system (CR tubes, Centrix, Inc, Shelton, CT) until the preparation was full. Large condensers were used to smooth the occlusal portion. The composite was cured from the occlusal with a VIP curing light. The light output of the curing light was assessed using a power meter (Power Max 5200, Molectron, Portland, OR). A 10.25 mm curing tip was utilized and held in place with a mounting jig. The cusp tips were flattened very slightly using a belt sander (Surfmet I, Buehler, Lake Bluff, IL) to provide a uniform distance of 0.75 millimeters from the light tip to the occlusal surface of the resin composite restoration. Various energy densities were selected based on the results of the first part of study detailed earlier for profiling the degree of conversion and Knoop hardness at the 5mm gingival levels. See Table 5.

Completed specimens were finished immediately with polishing discs (Soflex, 3M, St. Paul, MN), photographed with 2x magnification (Elite Chrome, Kodak, Rochester, NY), impressed with polyvinylsiloxane impression material (Express, 3M, St. Paul, MN), and then stored in 37 degree Centigrade water for 24 hours. The specimens received 1000 thermocycles with a 30 second dwell time at 5 to 55 degrees Centigrade in a custom-made thermocycler. After thermocycling, the specimens were photographed and impressed.

Next, the specimens were stressed in a custom-made OHSU mechanical fatigue-

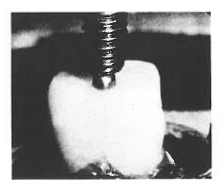


Figure 11: Fatigue cycler

cycling machine. See Appendix under Equipment for more details. A 2 millimeter stainless steel ball was cemented onto the occlusal surface of the restoration with a chemically-curing resin cement. (C&B Cement, Bisco, Schaumburg, IL). The specimens (Figure 11)

were then placed in the fatigue-cycler and stressed with a stainless steel bolt contacting the ball for 50,000 cycles under a cyclic load of 18 to 85 Newtons at 1.25 Hertz and verified using a Material Testing System machine (MTS Systems Corp, Eden Prairie, MN) by placing the load cell in the fatigue-cycler and loading with the stainless steel bolt. The specimens were constantly bathed in re-circulated 37 degree Centigrade water. After 50,000 cycles the specimens were removed from the fatigue machine and photographed and impressed as before. The estimated range of actual clinical masticatory forces range from 9 to 180 Newtons at 3 to 4 Hertz. Two hundred fifty thousand cycles has been related to approximately one year of clinical use.

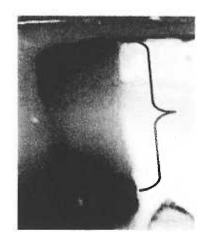
Replicas of the specimens were created using epoxy resin (Buehler, Lake Bluff, IL). The replicas were mounted on aluminum stubs, sputter-coated with 50 nanometers of gold-palladium (Hummer VII, Anatech Ltd, Alexandria, VA) and the gingival margin was examined and quantified by Scanning Electron Microscopy (SEM) (JXA-6000, JEOL, Tokyo, Japan). Examination of the marginal breakdown took place for each specimen at the pre-operative, post-thermal, and 50,000 cycle steps. Quantitation consisted of 25 equally-spaced measurements of the width of the gingival defect along the entire length of the four millimeter gingival margin. A percentage of the margin deemed defective was determined by dividing the number of the 25 locations with defects by 25 and expressed as percentage. Polaroid (T-52, Cambridge, MA) photographs at 14x magnification were taken of restorations and representative photographs were taken at 1000x magnification of select marginal defect areas. See Appendix under Equipment for more SEM details.

Marginal leakage of each specimen was determined with silver nitrate dye penetration (J.T. Baker, Palmyra, N.J) after completion of the fatigue cycling. New preparations and composite restorations were created with identical energy densities as the fatigue specimens. They were polished, left in 37 degree Centigrade water as before, but served only as pre-operative, non-fatigue specimens for marginal leakage examination.

All tooth surfaces were covered with two coats of fingernail polish to within 1.0 mm of the tooth-restoration margin. The occlusal surface of resin composite restoration was also covered with nail polish because of the concern for enhanced microleakage from the occlusal margins due to the reduction of the occlusal surface pre-operatively and the subsequent reduction in circumscribing enamel. The specimens were immersed in 3 moles per liter silver nitrate for 24 hours in a dark drawer. They were then removed, rinsed with de-ionized water, and placed in film developer (Eastman Kodak, Rochester, NY) under fluorescent lights for 24 hours. On removal from the developer, the teeth were rinsed in de-ionized water, embedded in epoxy resin (Buehler, Lake Bluff, IL) and allowed to set overnight. Three mesiodistal sections were made through the restorations

using a 0.3 millimeter thick diamond saw (Isomet, Buehler, Lake Bluff, IL) in order to assess marginal leakage along the entire length of the preparation interface.

Figure 12: Cross-section of restoration after silver nitrate stain



Length of Stain/ Length of preparation x 100%

Leakage scores were assessed on the six resultant surfaces using IP Lab software (Scanalytics, Fairfax, VA) and the extent of the dye penetration was expressed as a percentage of the entire cross-sectional length of the preparation interface. See Figure 12. The most severe dye penetration of the six surfaces was used as the score representing that specimen.

RESULTS / DISCUSSION

Effects of Degree of Conversion on Marginal Integrity. -- First Pilot

The first pilot study used the results of the initial data evaluating the effects of energy density on hardness and degree of conversion as seen in Table 1 and 2. The initial groups selected were summarized in Table 5 earlier. Only one sample from each group was created for an "n" of 1. Photos of the restorations, SEMs and gap width data per energy group can be seen in Appendix under Part 2, First Pilot. Several problems were encountered and identified during this initial pilot study.

First, as stated earlier, the variability in length and occlusal shape created from preparing an anatomic tooth could result in extreme inconsistency in gap formation in the gingival margin of the restoration, especially at the lower energy densities. Therefore, the occlusal table was flattened to provide a uniform 5 millimeter depth and shape to the preparations of the teeth. A folded piece of electrical tape 0.75 mm in thickness, placed on the occlusal surface opposite the preparation, would serve as a spacer to provide consistent light curing distance.

Second, it was noted that the gingival margin was difficult to polish to the exact margin without leaving slight overhangs of resin, especially at higher energy densities.

Therefore, the proximal area was flattened slightly to provide better access to polishing.

Third, it was obvious that the 4000 mJ/cm² group had significant degradation in the gingival marginal area. However, as stated earlier, the original hardness and degree of conversion testing only evaluated the middle third of each increment. Therefore, a new set of removable specimens was created from a flattened tooth template to determine the degree of conversion directly from the apical portion of the specimen. It then became

apparent that a significant descending gradient of degree of conversion values was seen at the gingival margin below the 12000 mJ/cm² level, providing an opportunity to evaluate even more levels of conversion (Table 6).

Fourth, a higher number of cycles on the mechanical fatigue cycling machine was used to see if greater differences could be discerned in the gingival margin, especially at higher energy densities. Mechanical cycling was increased to 250,000 from 50,000 cycles.

Second Pilot. -- Photos of the restorations, SEMs and gap width data per energy group can be seen in the Appendix under Part 2, Second Pilot. Only one sample from each group was produced for an "n" of 1. Several problems were encountered and identified during this second pilot study.

First, the definition of "gap width" needed to be expanded to include defective marginal areas instead of areas that had a true space or gap. This allowed a better assessment of all marginal areas, especially in the 4000 mJ/cm² group, which had an obvious loss of resin composite but still maintained relatively small "gaps".

Second, it was felt that the amount of proximal flattening to facilitate polishing was too severe. This could compromise the effect of enamel bonding, especially as the gingival margin was approached. Future groups would include subtle flattening only in the immediate gingival margin area.

Third, it was recommended to include two control groups. One group would be at 4000 mJ/cm² without any thermal or mechanical cycling to see the effects of water storage only on the marginal defects. The second control group would be more

traditional with 2 millimeter incremental placement of the resin composite and 40 seconds at 600 mW/cm² irradiance per increment. The first increment would only be 1 millimeter in thickness.

Fourth, a higher number of cycles on the mechanical fatigue cycling machine was attempted to see if greater differences could be discerned in the gingival margin, especially at higher energy densities. Maximum mechanical cycling was increased to 500,000, with specimens also evaluated after 100,000. In addition, specimens were created after 300,000 cycles, to be evaluated if necessary.

Third Pilot.— Representative photos of the restorations, SEMs and all of the defect width and percent defective data can be seen in the Appendix under Part 2, Third Pilot. Three specimens from each group were produced for an "n" of 3. A summarized version of the data can be seen in Table 8.

Table 8: Defect widths & percent defective margin n = 3 (width in microns)

	Pre-op			Post-Thermal			100K			500K		
	width	st dev	% def	width	st dev	% def	width	st dev	% def	width	st dev	% def
4000-W mJ/cm2	230.7	345.5	58.5	229.2	342.3	60.3	228.5	342.9	58.3	229.5	343.1	59.0
4000 mJ/cm2	466.1	247.5	100.0	474.0	246.3	100.0	479.5	248.6	100.0	488.4	245.9	100.0
6000 mJ/cm2	5.9	5.9	15.7	11.0	1.5	71.0	11.6	3.3	70.7	13.2	7.0	74.0
8000 mJ/cm2	11.8	5.0	36.7	25.1	8.6	97.0	22.2	14.0	61.0	25.2	7.6	82.3
10000 mJ/cm2	20,6	16.7	40.0	26.5	12.8	93.7	27.6	10.7	94.3	27.3	10.1	83.3
12000 mJ/cm2	26.9	3.0	88.7	30.5	4.0	100.0	26.0	5.0	98.3	24.0	12.3	84.7
24000 mJ/cm2	13.8	5.1	63.7	20.3	3.7	80.7	18.5	1.7	77.0	17.1	1.6	78.7
Control	3.4	5.3	7.8	8.0	9.1	81.3	6.3	7.0	57.3	5.5	6.0	60.3

4000-W mJ/cm² specimens placed in water only for equivalent time of stressed groups

A couple of problems were encountered and identified during this third pilot study. It became apparent after analyzing the data that the defect widths did not fully represent the true nature of the defects, especially in the 4000 mJ/cm² groups. The current technique of measuring defect width using the SEM could not account for the increase in depth of the defect. This became more significant as greater amounts of cycling or time occurred. Therefore, it was determined that the gingival margins be assessed by visual rating of the sputter-coated epoxy replicas by three examiners at 50X magnification (SMX-10, Nikon, Oak Ridge, TN). A ridit scale from 1 to 11 was created using techniques modeled after Mahler, et.al. 153 See scale in Appendix under Part 2, Final Method. This scale uses a numbered scale of restoration images that exhibit progressively increasing degrees of marginal defects. The procedure involves assigning to the restoration being evaluated, the number of the scale image to which it is most similar with respect to the extent of marginal defect. The data may be analyzed using parametric or non-parametric statistical methods. The defect and ridit data is provided in Table 9.

The tooth specimens outlining the ridit scale were available in the form of sputter-coated epoxy replicas to view at 50X magnification. The evaluators were calibrated with several practice specimens and were encouraged to rate a specimen overall with a score that may reflect a mental averaging of a combination of scores within the same specimen. For example, one half of the gingival margin could rate a "2" on the scale and the other half could rate a "6" to give a final score of "4" for that specimen. The specimens were rated blindly and randomly, with the examiners unaware of the specimen type. Any scoring discrepancy of greater than "2" between the evaluators was reassessed and the

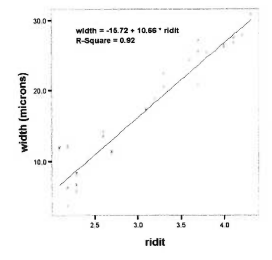
evaluators were allowed to change their score, if desired.

Table 9: Defect widths and ridit scores n = 3 (width in microns)

	DC at 5mm		Pre-op			Post-Thermal			100K			500K					
Energy Density	% of max	width	st dev	ridit	st dev	width	st dev	ridit	st dev	width	st dev	ridit	st dev	width	st dev	ridit	st de
4000-W mJ/cm2	20	230.7	345.5	4.8	2.3	229.2	342.3	5.3	2.6	228.5	342.9	5.9	3.1	229.5	343.1	6.4	3.3
4000 mJ/cm2	20	466.1	247.5	7.4	0.8	474.0	246.3	8.1	0.2	479.5	248.6	9.1	0.5	488.4	245.9	10.3	1.2
6000 mJ/cm2	45	5.9	5.9	2.2	1.3	11.0	1.5	2.7	1.6	11.6	3.3	2.1	1.0	13.2	7.0	2.6	1.1
8000 mJ/cm2	58	11.8	5.0	2.2	0.6	25.1	8.6	3.8	1.6	22.2	14.0	3.3	3.3	25.2	7.6	3.7	1.7
10000 mJ/cm2	68	20.6	16.7	3.7	2.3	26.5	12.8	4.1	1.8	27.6	10.7	4.2	1.5	27.3	10.1	4.1	1.8
12000 mJ/cm2	71	26.9	3.0	3.7	0.8	30.5	4.0	4.3	1.2	26.0	5.0	4.0	1.4	24.0	12.3	3.6	1.3
24000 mJ/cm2	84	13.8	5.1	2.9	0.2	20.3	3.7	3.0	0.7	18.5	1.7	3.2	0.2	17.1	1.6	3.4	1.0
Control (24000x3)	98	3.4	5.3	2.2	1.8	8.0	9.1	2.3	1.8	6.3	7.0	2.3	1.7	5.5	6.0	2.3	1.8

4000-W mJ/cm² specimens placed in water only for equivalent time of stressed groups

Figure 13: Width vs ridit



A linear regression analysis was performed using SPSS software relating defect width with ridit scores. When the 4000 mJ/cm² groups were left out because of the obvious lack of relationship between width and ridit scores, the correlation has an R² of 0.92. See figure 13. Including the 4000 mJ/cm² groups gives a correlation with an R² of 0.89 (not shown).

<u>Final Method.</u>— As expected, there is a high standard deviation in many of the groups and especially the 4000 mJ/cm² groups (both water only and fatigued) in both the defect analysis and microleakage study. Therefore, the number of specimens was increased to 8 per group and a ridit analysis only was conducted with 3 evaluators. The results are summarized in table 10. The raw data of the 3 evaluators is provided in the Appendix under Part 2, Final Method.

A paired t-test was completed comparing the pre-operative margins with the margins cycled 500,000 times. A significant degradation was found only in the 4000 $\,$ mJ/cm² groups (p < 0.05). However, since the degradation was seen both in the water-

KHN at 5mm 100K DC at 5mm Pre op Post thrmal 500K t-test 2-way **Energy Density** % max st dv % max st dv ridit st dv st dv | ridit | st dv | ridit st dv pre-500K ANOVA ridit 4000-W mJ/cm² 2.8 a 3.6 24.5 a 5.1 4.0 1.9 4.5 2.0 5.6 2.0 6.3 2.4 0.002 C 4000 mJ/cm² 0.035 2.8 a 3.6 24.5 a 5.1 4.7 2.7 4.8 3.0 5.4 3.2 6.0 3.8 С 6000 mJ/cm² 2.7 0.402 22.6 b 0.8 45 b 4.4 2.5 2.9 1.2 1.0 2.8 1.3 1.2 8000 mJ/cm² 39.2 c 4.5 57.3 b,c 0.2 4.3 8.0 4.2 4.2 0.113 3.3 1.4 1.1 1.0 b,c 12000 mJ/cm² 54.1 d 67.6 c,d 0.9 4.1 1.4 4.0 4.1 8.0 0.407 0.3 2.3 3.9 0.9 b,c 24000 mJ/cm² 72.8 e 0.3 80 d 0.9 3.0 0.6 3.3 0.6 3.3 0.5 3.2 0.6 0.516 a,b

1.2

2.2

8.0

2.3

0.9

2.3

0.8

0.487

Table 10: Ridit analysis n=8 3 evaluators

a - f denotes significant differences in columns

1.6

2.4

1.2

97 e

96.5 f

Control (24000x3)

4000-W mJ/cm² specimens placed in water only for equivalent time of stressed groups

only and the stressed groups, it can be concluded that the water was primarily responsible for the degradation and not the thermal-mechanical stressing.

A two-way analysis of variance (ANOVA) was completed with SPSS software to test the effect of energy density or thermal-mechanical stressing on gingival marginal defects (ridit scores). See printout in Appendix under Part 2, Final Method. Significant

differences in the ridit scores were found based on energy levels. Tukey's post hoc test was used to determine differences between individual groups (α = 0.05; Table 10). No overall significant difference was seen based on stressing (p = 0.167). Differences were seen based on energy density (p = 0.0001). There were no significant differences between the incrementally-filled control group, and the bulk-filled 24000 and 6000 mJ/cm² groups.

The results of the microleakage study are summarized in Table 11. The raw data and photos of all of the sectioned specimens used for scoring are shown in the Appendix under Part 2, Final Method. An unpaired t-test was completed comparing microleakage of pre-operative with post-operative margins (1000 thermal-cycles and 500,000 fatigue-cycles). No significant difference was seen between these two comparisons at any energy density. The microleakage scores of the groups had an overall normal distribution (p > 0.05). See SPSS printouts in Appendix under Part 2, Final Method. A two-way analysis of variance (ANOVA) was completed to test the effect of energy density or thermal-mechanical stressing on microleakage. No overall significant difference was seen based on stressing or energy level (p = 0.167 & 0.212). A one-way analysis of variance (ANOVA) was used to test the effect of energy density on microleakage pre-operatively and also post-operatively. No significant differences were found. SPSS software was used and the results are shown in Table 11. See printouts in Appendix under Part 2, Final Method.

Table 11: Microleakage scores: percent of cross-sectional length of preparation interface $\,n=8\,$

Energy Density	Avg Pre	st dv	Avg Post	st dv	Unpaired t-test	Two way anova
4000-W mJ/cm ²	53 a	31	52 a	21	0.96	а
4000 mJ/cm ²	53 a	31	56 a	27	0.86	а
6000 mJ/cm ²	58 a	28	55 a	30	0.85	а
8000 mJ/cm ²	57 a	25	47 a	24	0.44	а
12000 mJ/cm ²	59 a	18	56 a	31	0.84	а
24000 mJ/cm ²	38 a	14	30 a	5	0.15	а
Control (24000x3)	68 a	25	41 a	32	0.11	а

All "a" denotes no significant differences between groups in columns 4000-W mJ/cm2 specimens placed in <u>water only</u> for equivalent <u>time</u> of stressed groups

PART THREE

MATERIALS AND METHODS

Effects of Degree of Conversion on Flexural Strength and Modulus.--

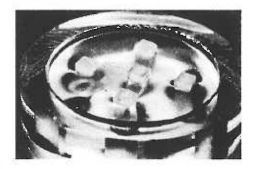
Resin composite bars were produced having the same approximate degree of conversion as the gingival margins created by various energy densities in Part 1 and 2. These bars were tested to determine their flexural strength and flexural modulus. Knoop hardness testing was used to approximate the degree of conversion because it was quicker and easier and because it correlated so well with degree of conversion in previous testing $(R^2 = 0.99; Figure 9)$.

Specimens were made in square glass rods with dimensions of 2 millimeters by 2 millimeters by 25 millimeters. The Z250 resin composite shade A-2 was light-cured in a Triad II (Dentsply, York, PA) laboratory curing unit for various time intervals to produce surface Knoop hardness levels corresponding to the marginal Knoop hardness levels found in the previous procedures. This approach was taken because it was difficult to measure the power output from the three bulbs in the Triad II unit. In addition, the light in the Triad II unit was unfiltered, and could not be directly compared to the VIP light in terms of curing efficacy or total energy density. The specimens were cured in a vertical position in the center of a rotating platform in the center of the Triad oven to produce the most uniform degree of conversion possible. The specimens were stored dry in a light-proof film canister for 48 hours, polished on each side for 10 seconds with 5 micron aluminum oxide paste on an 8 inch nylon polishing cloth, and then tested in 3-point bending (20 mm span) on a universal testing machine (Instron, Instron Engineering Corp., Canton, MA) at a crosshead speed of 0.254 mm/min. The flexure strengths were

determined from the maximum load using the equation: Flexural strength = $3Fl/2bh^2$ where "F" is the resultant force in pounds, "I" is the length of the specimen over the support beams (20mm) and "b" and "h" are the base and height width of the bar (2mm). The flexural modulus was determined from the initial slope of the force-deflection curve using the following equation: $E = L^3/4bh^3 \times F/Y$ with "F" (force in pounds) determined on a straight portion of the curve and Y as the crosshead speed per minute (0.01 inches / minute) divided by the chart speed (1 inch/min) with

"L, b and h" as before. Five specimens of each group were tested. Specimens from the flexure strength tests were evaluated for Knoop hardness in the Kentron hardness tester as before. The specimens were polished on each side for 10 seconds with 5 micron aluminum oxide paste on an 8 inch nylon

Figure 14: Center KHN measurement



polishing cloth (Buehler, Lake Bluff, IL). Graphite from a mechanical pencil was rubbed on the lightly polished surface of the composite specimens before indentation. A load of 200 grams was applied through a diamond pyramid indenter with a dwell time of 10 seconds. The length of the indention was measured and a hardness number calculated. Three measurements were made on two opposing sides of each bar and averaged. The average from three bars was calculated (n = 3). These averages were then expressed as a percentage of maximum hardness (71.8 kg/mm²) from this test and compared with the previous data of Knoop hardness values produced in the gingival margin of the 5 millimeter specimens under various energy densities. In addition to the surface hardness, the center hardness of each bar was tested by embedding the previously fractured bars

vertically in epoxy resin (Buehler, Lake Bluff, IL) in an acrylic ring and allowing them to cure overnight in a light proof container (Figure 14). The specimens were polished and rubbed with graphite before indentation. One hardness value was taken at the center of both ends of a bar and an average taken. The average of three bars was expressed as a percentage of maximum hardness (n = 3). A one-way analysis of variance (ANOVA) was used to test the effect of curing time on Knoop hardness (surface or center), elastic modulus and flexure strength. SPSS software was used and the results are shown in Table 12. See printouts in Appendix under Part 3. Detailed data can be seen in Appendix under Part 3.

RESULTS

Effects of Degree of Conversion on Flexural Strength and Modulus .--

Table 12 below shows the various hardness ratios and mechanical properties obtained at various Triad II curing times and compares them to similar hardness ratios found at the gingival margin at various energy densities with Part 1 and 2.

Table 12: Mechanical properties compared to KHN in Part 1 and 2

Part 1 n=3

n = 3

n = 5

n = 5

	KHN at 5mm			
Energy Density	% max	st dv		
4000 mJ/cm ²	2.8 a	3.6		
6000 mJ/cm ²	22.6 b	0.8		
8000 mJ/cm ²	39.2 c	4.5		
12000 mJ/cm ²	54.1 d	0.3		
24000 mJ/cm ²	72.8 e	0.3		
Control (24000x3)	96.5 f	1.2		

Time secs	KHN surf		KHN center		Flexural Mod		Flexure Strength			
	% max	st dv	% max	st dv	GPa	st dv	% max	MPa	st dv	% max
4	31.4 a	3.9	22 a	0.9	2.5 a	0.5	20.0	35.2 a	5.5	26.1
8	50 b	2.7	50.7 b	3.1	6 b	0.6	47.5	71.4 b	5.8	52.8
16	70.1 c	1.3	69.1 c	2.1	9.1 c	1.2	72.5	90.2 c	12.4	66.7
80	91.1 d	0.9	91.2 d	5.5	12.3 d	0.3	97.4	125.2 d	7.4	92.6

a - f denotes significant differences within columns

A correlation was performed using SPSS software relating the surface Knoop hardness of the flexure bars with flexural modulus (Figure 15) $R^2 = 0.100$ or flexural strength (Figure 16) $R^2 = 0.98$.

Figure 15: Surface KHN of flexural bars vs. flexural modulus

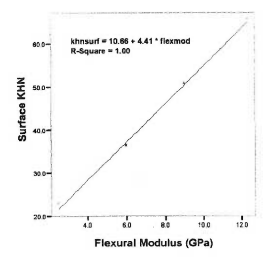
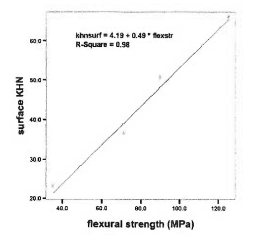


Figure 16: Surface KHN of flexural bars vs. flexural strength



It may be possible to extrapolate from these excellent correlations a good relationship between the mechanical properties of the flexural bars produced with various energy densities and the mechanical properties found at the gingival margin at various energy densities seen in Part 1 or 2 (Table 12).

OVERALL DISCUSSION

Many of the problems with the initial pilot study investigating the effect of energy density on degree of conversion and hardness in the gingival portion of the restoration were resolved with modifications in technique. These modifications consisted of standardizing the length and shape of the specimen by flattening the occlusal of the template and testing the specimens directly on the occlusal and gingival surfaces. The flattened occlusal surface reduced the potential variation in data, especially in the gingival margin area, that could occur from inconsistent lengths in specimens produced from an anatomic occlusal. It allowed greater standardization of the position and angulation of the light guide and permitted the testing of the most apical portion corresponding to the gingival margin. At lower energy densities it became apparent that the Knoop hardness and degree of conversion values decreased markedly in the apical half millimeter of the gingival increment. However, the new technique did not allow data determination within the body of the specimen.

There was a strong correlation when relating Knoop hardness and degree of conversion (R² = 0.99) with the direct technique. Once again, the attainment of hardness fell behind the degree of conversion of the carbon double bonds (Table 10) and this could be due to the late formation of networking links in the polymer chains.⁵⁴ A greater increase in hardness relative to maximum hardness occurred with higher energy levels and subsequent increases in crosslinking.⁵⁴ The theory reported earlier that the lag of hardness behind degree of conversion resulting from an overestimation of percent cure at lower energy levels from sectioning and polishing is unlikely with the new technique.

Another significant modification in technique consisted of improving the visualization of the Knoop hardness indentations with the use of graphite. It was noticed in the course of working with the hardness tester that a potentially significant variation

existed between operators or even within the same operator when reading or re-reading the same indentation. Poor contrast between the indentation and the surrounding resin may have contributed to this variation and potential underestimation of the length. A high level of polish may

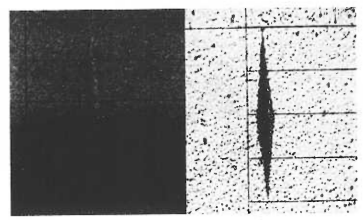


Figure 17: Knoop hardness indentation at 200x (l) Non-graphite enhanced; (r) graphite enhanced

have improved the contrast, but was not attempted in order to reduce the potential of altering the properties of the lower cured areas. Graphite from a lead pencil placed on the specimen before indentation provided dramatically improved visualization and substantially reduced the standard deviation. See figure 17. A concern was expressed over the possibility of the graphite altering the test or test surface. Therefore, graphite was also placed on the nearby epoxy mounting resin, on which the indentations were readily observed without the graphite. No differences were seen in hardness measurements with or without graphite. See data in Appendix under Part One.

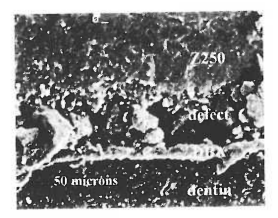
It is common in hardness testing to express the hardness values as a percentage of the top or upper surface of the composite with 80 percent typically chosen to be the critical level for adequate depth of cure. That was modified slightly in this study to reduce the potential error in comparing groups with softer top surfaces. The values were standardized by expressing the hardness and the degree of conversion numbers as a percentage of maximum. The maximum Knoop hardness value found for Z250 was originally 98.7 but changed to 72.7 after using the graphite technique that provided easier visualization of the indentation. The original data served primarily as a pilot to provide initial guidance to energy density estimation. The study was essentially repeated using modified techniques.

Many of the problems with the next part of the study investigating the effect of degree of conversion on marginal integrity in the gingival portion of a restoration were resolved with multiple modifications in technique.

The flattening of the occlusal surface became even more critical in this next phase of the project due to the variability in anatomic form of the multiple teeth used. A uniform 5 millimeter depth and shape to the preparations was produced.

In the higher energy density groups, especially the control group, it was difficult to finish and polish the gingival margins to the exact margins without leaving a slight overhang of resin composite. This became especially difficult in teeth with any sort of furcal concavities or deep contours. Many of these overhangs did not become obvious until observed under 50X magnification. Flattening the gingival marginal area with a disc just before preparation helped to create a more manageable surface. A recent study found that 51.3% of bonded Class 2 resin composite restorations had significant flash at the interproximal margins in teeth restored in a phantom head after a typical finishing and polishing procedure. 154

It was difficult when using scanning electron microscopy to consistently identify the true width or length of gaps or defects. A gap could be defined as a space or lack of restorative material whereas a defect may be defined as a loss of material or both.



It was often difficult to differentiate between the two.

Figure 18: Gingival marginal defect at 1000 X

Some areas were occupied with a thicker layer of

bonding agent, but it was problematic to determine if it was intact or fractured or simply filling a void (Figure 18). Variation in the angulation of the interface could produce variable results when attempting to measure gap width. Also, analysis at various magnifications seemed to produce variable interpretations by itself. The procedure was technically complex, expensive and very time consuming – taking up to one hour to examine and quantify a single specimen. Finally, the technique was abandoned when it was realized that the SEM could not adequately measure the depth of defects. This became significant in the gingival marginal areas of the lowest energy density groups.

It was determined that the gingival margins would best be assessed by a visual rating or ridit analysis that used a numbered scale of restoration images that exhibited progressively increasing degrees of marginal defects. This technique has been used successfully in the past by Mahler et. al. to evaluate amalgam margins. However, instead of enlarged photographic images, the sputter-coated replicas were examined using 50X stereomicroscopy by three examiners. Stereomicroscopy allowed the necessary assessment of the depth of the defects in addition to width and length. The interval nature of the ridit scale also lends itself to the use of the more powerful parametric

statistics. SEM A good correlation was found between defect widths as measured by SEM and ridit scores ($R^2 = 0.92$).

It was interesting to see no significant effect of thermal-mechanical stressing on microleakage or marginal defects. Several studies suggest that thermocycling resin composite restorations may not significantly affect microleakage in Class 2 resin composite restorations. ^{124,125,155} In a recent study, Wibowo used Z100 and Single Bond in Class 2 restorations with gingival margins in dentin. He found no significant difference between the thermal-cycled and non thermal-cycled groups using a computer imaging technique of the gingival floor after breaking away the entire resin restoration. ¹⁵⁵ The effects of thermal-cycling resin composite restorations on microleakage remains questionable and perhaps the procedure is unnecessary.

Studies have shown conflicting results of load cycling, reinforcing the need for careful evaluation and interpretation due to differences in experimental designs. 100,116,133-142 Several researchers found no effect on microleakage from load cycling of resin composite restorations. 125,133-138 However, a few authors did find an effect from load cycling. 116,134,140 A couple of investigators found an effect from a simultaneous combination of thermal-cycling and loading, 100,139,156 while another had mixed results. 157 Virtually all of the studies evaluate only microleakage with thermal-mechanical loading. However, a couple of recent studies have found conflicting results when evaluating marginal gap formation and a combination of thermal-mechanical loading. Friedl and others, 158 using testing conditions fairly similar to this study (5000 thermal cycles and 500,000 load cycles at 1.7 Hz and 72.5 N in Class 2 restorations), found no significant increase in marginal gaps using ProBond and Prisma TPH, but found a significant

increase using Scotchbond MP and Z100. Interestingly, these investigators using a measurement technique similar to this study, found an actual <u>decrease</u> in microleakage at the dentinal gingival margin in both groups after thermal-mechanical loading. They theorized that the decrease could have been due to hygroscopic expansion. However, a recent study by Lutz and Krejci utilized a three dimensional scanner and found an increase in the percentage of marginal gap formation in MOD resin composite restorations after a combination of simultaneous thermal and load cycling (3000 thermal cycles and 1,200,000 load cycles at 1.7 Hz and 49 N).

Several studies have found no correlation between gap formation and microleakage. ^{157,159-161} Similar results were found in this investigation. A linear regression analysis was performed using SPSS software relating the ridit scores and the microleakage values and found a poor correlation with an R² of 0.24. Any significant increase in marginal gap or defect would probably not contribute to an increase in microleakage, once the threshold width for dye penetration is met.

The preparation and placement of the resin composite in this study was based on techniques supported in the literature. One study found a significant reduction in microleakage by placement of a conservative bevel along the facial and lingual enamel margins. Also, the use of the syringe tip for resin composite placement has been shown to decrease the viscosity of the material and decrease voids along the preparation walls. 162,163

Various techniques have been utilized to assess dye penetration in microleakage studies. Many utilize a single section through the center of the restoration. 134-137,140,164,165 However, a three dimensional technique whereby the entire restoration is removed has

been shown to reveal more extensive dye penetration, ¹⁶⁶⁻¹⁶⁹ but it is more time consuming and does not allow good visualization of dentin tubule leakage. ¹⁶⁹ Using multiple sections, as used in this study, seems to be a practical compromise. ^{124,133,139,141,169} It was not surprising to see extensive leakage between resin composite and the dentinal gingival margin. This agrees with many studies utilizing dentin bonding agents. ^{116,124,133,136,138,169} A recent study by Hagge and others found extensive microleakage under composite using a 5th generation bonding agent with and without flowable liners. The only group not to suffer heavy leakage utilized a resin-modified glass ionomer in an open sandwich technique. ¹⁶⁴ The enhanced performance of resin-modified glass ionomer liners on dentinal gingival margins has been substantiated in several laboratory and clinical studies. ^{158,165,170}

It was interesting to find, however, no significant increase in microleakage in the groups with very low levels of gingival marginal cure. Microleakage testing using silver nitrate dyes is a very stringent test that perhaps overestimates the amount of leakage that will occur clinically.¹⁵⁷ The dye is of considerably smaller size and molecular weight than bacteria, or endotoxins.¹⁷¹ Gap formation found at the dentinal gingival margin found in this study and other studies was much larger than the silver nitrate dye.^{100,158}

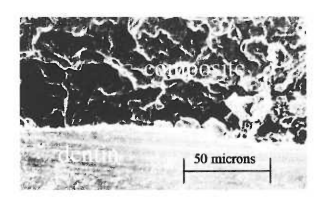


Figure 19: 500X Dentinal gingival margin 4000 mJ/cm² 500K cycles



Figure 20: 14X 5 mm composite restoration 4000 mJ/cm² 500K cycles

Perhaps any significant increase in marginal gap or defect would not contribute to an increase in microleakage, once the threshold width for dye penetration is met. Even the poorest cured group (4000 mJ/cm²) that sustained significant marginal degradation or defects over time was found in pilot studies to have relatively small marginal "gaps" at the depths of these defects (Figures 19,20). Therefore, except for the obvious loss of superficial marginal resin composite material, these lower cured groups would possibly not leak any more or less than the higher cured groups (Figure 21).

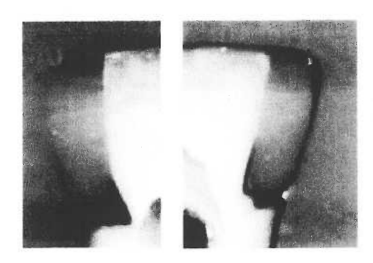


Figure 21:
Microleakage: (l) pre-op; (r) 500K cycles
4000 mJ/cm²

There was a significant loss in marginal integrity in the gingival margins with the lowest degrees of conversion after thermal-mechanical stressing. However an identical group stored only in water suffered similar degradation. This suggests that water, not the thermal-mechanical stressing contributed to the increase in marginal defects in these susceptible groups. The lesser the extent of the polymerization reaction, the more residual monomers are available to be leached. This inverse relationship ($R^2 = 0.95$) between degree of cure and percent elution was confirmed in a study by Rueggeberg and Craig. In that study, resin composite was cured through various thicknesses of cured

resin composite in order to reduce the irradiance and degree of conversion. After storing the variably cured resin composite in water for four weeks, the elution of residual monomers was determined gravimetrically.⁵⁰

There was no significant difference in marginal defects between the incrementally-filled control group and the bulk-filled 24000 and 6000 mJ/cm² groups. Although the 6000 mJ/cm² group was statistically similar to the control and 24000 mJ/cm² groups, it was on the edge of significant marginal deterioration as seen by the 4000 mJ/cm² groups and could not be recommended clinically. Any slight variation in light guide angulation or loss in power density could result in severe undercuring and marginal degradation. Also, these lower energy groups could be susceptible to solvents and enzymes not tested in this study. The lack of marginal degradation in the 6000 mJ/cm² group may have been a combination of reduced shrinkage stress and just enough degree of conversion to prevent degradation primarily from water. Therefore, a recommended lower limit of gingival margin acceptability in a bulk-filled resin composite restoration was created by 80% of maximum conversion, 73% of maximum hardness and approximately 70% of maximum flexural strength and modulus in the gingival marginal area as seen in the 24000 mJ/cm² group.

Caution should be exercised when attempting to extrapolate the results of this study to resin composites other than minifilled hybrid Z250. Direct comparisons of the sufficiency of irradiation using absolute surface hardness are not generally appropriate because the hardness is influenced by both the nature of the resin matrix and of the inorganic filler. Solution and others used a mathematical model to suggest that the depth of cure may be defined as the level at which the hardness value is equivalent to at

least 90% of the hardness at the top of the composite.⁸⁸ Others have suggested that this top-to-bottom gradient should not exceed 20% ⁸⁹ and should be considered a realistic measure of depth of cure.⁹⁰

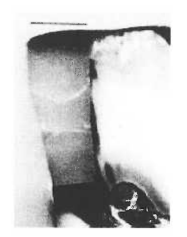
Various studies have advocated incremental instead of bulk placement and curing for resin composite to decrease the effects of polymerization shrinkage and to provide complete polymerization in deeper cavities. 2,58,64-65 Incremental placement may also reduce the configuration factor – the ratio of bonded to unbonded surfaces – and reduce the stress by making more surface resin available for flow.⁶⁶ However, other studies found no difference between bulk and incremental placement on marginal gap formation⁶⁷ or fluid permeability.⁶⁸ A finite element study by Versluis and others showed that incremental filling yields higher polymerization shrinkage stresses. ¹⁷² Polymerization contraction of each individual filling increment causes some deformation of the cavity. forcing the walls to bend and decreasing cavity volume. Less composite placed for the next filling increment results in a cavity that is volumetrically filled with less composite material than the original volume of the cavity and that results in a higher stress state. 172 Using photo-elastic material, Jedrychowski and others found that bulk resin composite placement generated the lowest shrinkage stresses compared with various other incremental techniques. 173

This study found no significant difference in gingival marginal defects between incremental cure and bulk curing at an energy density of 24000 mJ/cm². The reasons for these findings despite the great disparity in energy density between the groups may be due to a complex interaction of multiple factors. With incremental curing, the mechanical properties of the resin composite adjacent to the gingival margin and the

bond between resin composite and adhesive are maximized. These factors favoring improved gingival margin performance are balanced with the increased polymerization shrinkage that occurs with the high energy density. With bulk cure, polymerization shrinkage is lessened due to a decrease in applied energy density. Counteracting this is a decrease in the mechanical properties of the resin composite adjacent to the gingival margin. However, with the higher energy density group (24000 mJ/cm²) adequate mechanical properties at the gingival margin may result since functional forces through the restoration are likely dissipated to the tooth via adhesion. Another factor opposing gingival margin integrity in the bulk cure group may be a decreased adhesion of resin composite at the gingival margin as a result of the reduced energy density.

The incremental placement of resin composite utilized in this study was difficult to accurately place and reproduce, requiring magnification to insure uniformity among specimens. Unlike bulk placement, an obvious demarcation could be seen on occasion between sequential layers of the incremental technique. Separate silver nitrate staining along the internal restoration interface could be seen on occasion corresponding to the demarcation areas¹⁶⁴. See Figure 22.

Figure 22: Layering of increments



The 5 millimeter long posterior resin composite restorations were created with various energy densities, but these energy densities were determined based on the power density recorded at the end of the light guide. It became of interest to know just how much power density was actually transmitted through the resin composite to the gingival surfaces. Therefore, a simple study was conducted to determine what percentage of the

power density delivered to the occlusal surface actually reached the gingival margin. See details and data in Appendix under Part 3.

The 1.6 percent value of this study agrees with a recent study by Price and others that found an average of only 1.1 percent of initial light energy transmitted through 5 millimeters of various resin composites (1.6 for Z250).¹⁷⁴

It was difficult to consistently record the lower levels of power found at the gingival surface of the specimens. The Power Max reads heat energy and the ambient level of heat in the room of 1 mW tended to complicate testing at these very low levels. Also, the measurements reported with the Power Max tended to float slightly. Although not a problem at the higher levels recorded at the end of the light guide, greater variation and less reproducibility of the data was evident at these lower levels of recorded power.

Knoop hardness (R^2 = 0.96) or degree of conversion (R^2 = 0.93) at the gingival increment had a strong linear relationship to the log of the energy density (Figure 10). This agrees with Nomoto and others who reported a linear relationship between the depth of cure and the logarithm of total amount of exposure.¹⁷⁵ Cook described the intensity of light transmitted by the cured resin composite by the equation:¹⁷⁶

$$I = I_o \times 10^{-eL}$$

where I_o is the intensity of the light at the surface of the resin composite; L is the depth of cure and e is the attenuation coefficient due to the heterogeneity in resin composites altering the optical properties. Light is absorbed and scattered by matrix and fillers. When the specimen is irradiated for time t, the amount of light transmitted is defined by:

$$E = I x t = I_0 x t x 10^{-eL}$$

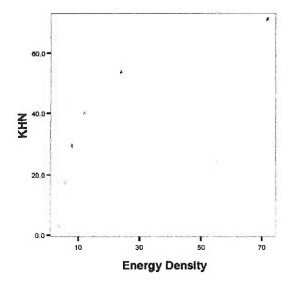
where E is defined as the minimum total amount of exposure required to activate polymerization. This equation can be expressed:

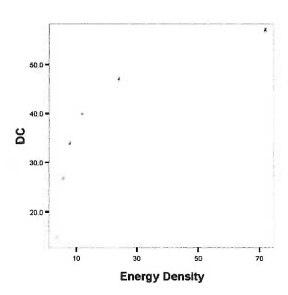
$$L = \{ \log (I_0 x t) - \log E \} / e$$

which indicates that a linear relationship exists between the depth of cure (L) and logarithm of total amount of exposure as was found in this study. 175,176

Figure 23 shows the correlation between Knoop hardness or degree of conversion and energy density without the logarithmic conversion. Notice the sudden drop in KHN or DC at lower energy densities consistent with the dramatic attenuation of energy exposure.

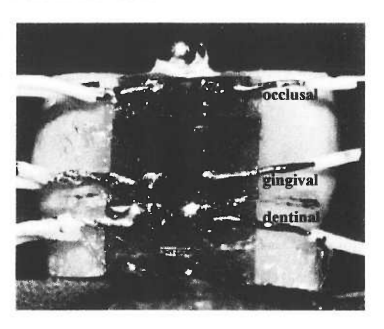
Figure 23: Knoop Hardness or degree of conversion vs energy density (flattened template) at the gingival increment





The maximum load of 85 Newtons used in this study could have been increased in attempt to produce gingival marginal degradation or until the restoration failed catastrophically. However, the clinical significance of such high forces and the likelihood of producing localized changes in the gingival area would be questionable. A simple study was conducted to determine the stress developed at the occlusal and gingival marginal area of the restoration. Three single strain

Figure 24: 3 Strain gauges at the occlusal, gingival and dentinal margins



gauges (CEA-06-032UW-120, Measurements Group, Raleigh, NC) were bonded on the resin composite (Control Group) near the occlusal, at the gingival margin, and on dentin near the gingival margin using cyanoacrylate and the tooth was subjected to the same cyclic forces as before (Figure 24). The generated strain was conditioned by a strain gauge conditioner (2100 System, Measurements Group, Raleigh, NC). Data indicates that the maximum amount of stress transmitted to the gingival marginal resin composite area is less than 12.4 Megapascals (MPa). The maximum stress near the occlusal marginal ridge was 15.5 MPa and near the gingival marginal dentin was 11.5 MPa. The bonded resin composite restoration appears to absorb most of the available stress, transferring the energy through the resin composite, the bonding agent and the supporting walls of the restoration without affecting marginal integrity. See details and data in the Appendix under Part 3.

Future studies are needed to determine the effect of energy density on caries resistance, interproximal wear resistance and post-operative sensitivity. Also, a less photosensitive material such as a microfill composite should be evaluated under similar conditions of this study to evaluate the opposite end of the spectrum of available restorative resin composites. This additional information may provide a more general description of the minimum hardness ratio necessary at the base of a resin composite restoration to maintain marginal integrity. Also, a yellowish discoloration was seen in the gingival areas of the poorly cured groups (4000 and 6000 mJ/cm²) after thermal-mechanical stressing. See the photos of the restorations in the Appendix under Part Two, 1st, 2nd and 3rd Pilot. A future study could examine the effect of water immersion on composite discoloration over time in resin composites with variable degrees of conversion using a colorimeter.

OVERALL CONCLUSIONS

Based on the limitations of this study, the following conclusions can be made concerning Z250 shade A-2 resin composite by 3M ESPE in class 2 slot preparations with gingival margins in dentin with various degrees of conversion. (see summary sheet following conclusions):

- There was a strong correlation between degree of conversion and Knoop hardness at the gingival margins. $(R^2 = 0.99)$
- There was a good correlation between degree of conversion or Knoop hardness and the log of energy density at the gingival margin. ($R^2 = 0.96 \& 0.93$)
- There was a good correlation between average defect widths and visual rating (ridit scores). ($R^2 = 0.92$)
- 4) Energy density had a significant effect on gingival marginal defects and no significant effect on microleakage.
- Water had a significant dissolving effect on the resin composite with very low degrees of conversion at the gingival margin producing defects.
- The maximum defect formed at the gingival margin with a Knoop hardness of 2.1 or 2.8% of maximum and a degree of conversion of 14.3 or 24.5% of maximum.
- There was no significant difference in gingival marginal defects between the incrementally-filled control group and the bulk-filled 24000 and 6000 mJ/cm² groups.

- A recommended lower limit of gingival margin acceptability in a bulk-filled Z250 resin composite restoration was created by 80% of maximum conversion or 73% of maximum hardness as seen in the 24000 mJ/cm² group (600 mW/cm² for 40 seconds). There was no significant difference between bulk curing and incremental filling resin composite within these limits.
- 9) There was no overall significant effect of thermal-mechanical stressing on gingival marginal defects or microleakage in class 2 resin composite restorations with gingival margins in dentin.

Summary Sheet

Ridit Analysis 3 evaluators n = 8

	KHN at 5mm		DC at 5mm		Pr	Pre op		Post thrmal		100K		500K		BY N
Energy Density	% max	st dv	% max	st dv	ridit	st dv	ridit	st dv	ridit	st dv	ridit	st dv	pre-500	ANOVA
4000-W mJ/cm ²	2.8 a	3.6	24.5 a	5.1	4.0	1.9	4.5	2.0	5.6	2	6.3	2.4	0.002	С
4000 mJ/cm ²	2.8 a	3.6	24.5 a	5.1	4.7	2.7	4.8	3.0	5.4	3.2	6.0	3.8	0.035	С
6000 mJ/cm ²	22.6 b	0.8	45 b	4.4	2.5	1.2	2.9	1.2	2.7	1.0	2.8	1.3	0.402	а
8000 mJ/cm ²	39.2 c	4.5	57.3 b,c	0.2	3.3	1.4	4.3	0.8	4.2	1.1	4.2	1.0	0.113	b,c
12000 mJ/cm ²	54.1 d	0.3	67.6 c,d	2.3	3.9	0.9	4.1	1.4	4.0	0.9	4.1	8.0	0.407	b,c
24000 mJ/cm ²	72.8 e	0.3	80 d	0.9	3.0	0.6	3.3	0.6	3.5	0.3	3.2	0.6	0.516	a,b
Control (24000x3)	96.5 f	1.2	97 e	1.6	2.4	1.2	2.2	0.8	2.3	0.9	2.3	0.8	0.487	а

Microleakage % of margin n = 8

Energy Density	Avg Pre	st dv	Avg Post	st dv	t-test	two way anova
4000-W mJ/cm ²	53 a	31	52 a	21	0.96	а
4000 mJ/cm ²	53 a	31	56 a	27	0.86	а
6000 mJ/cm ²	58 a	28	55 a	30	0.85	а
8000 mJ/cm ²	57 a	25	47 a	24	0.44	а
12000 mJ/cm ²	59 a	18	56 a	31	0.84	а
24000 mJ/cm ²	38 a	6	30 a	5	0.15	а
Control (24000x3)	68 a	25	41 a	32	0.11	а

Mechanical Properties n = 3

Part	1	n=3

	KHN at	5mm
Energy Density	% max	st dv
4000 mJ/cm ²	2.8 a	3.6
6000 mJ/cm ²	22.6 b	0.8
8000 mJ/cm ²	39.2 c	4.5
12000 mJ/cm ²	54.1 d	0.3
24000 mJ/cm ²	72 8 e	0.3

Control (24000x3) 96.5 f

Time	KHN	surf	KHN c	enter	Ela	stic N	/lod	Flexu	re Stre	ngth
secs	% max	st dv	% max	st dv	GPa	st dv	% max	MPa	st dv	% max
4	31.4 a	3.9	22 a	0.9	2.5 a	0.5	20.0	35.2 a	5.5	26.1
8	50 b	2.7	50.7 b	3.1	6 b	0.6	47.5	71.4 b	5.8	52.8
16	70.1 c	1.3	69.1 c	2.1	9.1 c	1.2	72.5	90.2 c	12.4	66.7
80	91.1 d	0.9	91.2 d	5.5	12.3 d	0.3	97.4	125.2 d	7.4	92.6

n = 5

n = 5

a-f denote significant differences in columns

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APPENDIX

PART ONE PILOT

Knoop Hardness

			t-test	4 0.63	2 0.24	5 0.59	4 0.98	7 0.04		m				t-test	2 0.95	92.7 0.38	5 0.81	3 0.61	5 0.15		8
			nal	3.2 86 2.2 87 86.5 85.2 89.4	78 75.9 74.2 83.2	74 75.9 70.6 76.5	65 0.9 65 64.8 66.2 64.4	7.6 44 8.1 40 31.5 39.9 47.7		40 31.9 40.4 48.3				nal	94.6 89.5 83.2		1 86.5	74.3	12 61 47.2 69.1 66.5		67.3
			Proximal	85.2	74.2	70.6	66.2	39.8		40.4				Proximal	89.5	88	81.9 85.1	74.8 84.1	69.		70
				86.5	75.9	75.9	64.8	31.5		31.9					94.6	90 2.5 91 91.6 88		74.8	47.2		12 62 47.8
			Avg	87			92	40					, ,	Avg	88	91	85	78	61		62
	cm ²	e .	stde	2.2	4.8	3.2	0.9	8.1	20	8.2		cm ²	a	stde	5.7	2.5	2.4	5.5	12	al	12
	4000 mJ/cm ²	Occlusal	avg	98	80	72	92	44	Gingival	77 44		12000 mJ/cm ²	Occlusal	avg	89	200	85	79	65	Gingival	65
	4	O	Avg. stdev avg stdev Avg	3.2	3.4	12	6		U			120	O	Avg. stdey avg stdey Avg	4.7	4.8	2.9	9.7	6.3		9.9
			Avg	86	82	70	65	47		53.5 48				Avg.	90	88	85	81	89		69
				83.8	83.2	80.7	75.4	52.9		53.5					94.9 90 4.7 89 5.7 89	93.8 88	87.5	87.5	71.3		72.1
			Axial	89.4 83.8 83.8 86	83.8 77.6 83.2	56.4 71.6 80.7	62.2 58.2 75.4	38.6 50.4 52.9 47		51				Axial	87.2 86.5	86	85.9 81.8 87.5	72.6 82.3 87.5	60.9 72.4 71.3 68	2	73.3
2				39.4	33.8	96.4	32.2	38.6		39.1					37.2	85.2	35.9	72.6	30.9		51.3
Section design			8	-	ا م	ر ا	4	ۍ , ,		5/max 39.1					~-	2	m	4	را ا		5/max 61.3 73.3 72.1 69
7			t-test	.05	6.0	.43	0.83	0.64		5				t-test	0.35	.15	0.23	0.04	0.15		5
5			I	74.8 66.7 86.5 0.05	6.7	57.9 64.4 84.5 0.43	0 7.0	9.6		6.6				1	2.3	88.6 84.6 72.4 0.15		2.4 0	5.3		5.7
			mal	.7 86	83.2 76.5 87.9	.4 84	49.8 39.4 50.7	mush 12.9 19.6		mush 13.1 19.9				imal	95.4 85.1 82.3	.6	69.6 76.7 80.1	63.5 65.8 72.4	45 40.4 35.3		45.6 40.9 35.7
			Proximal	99 8	2 76	9 64	8 39	th 12		sh 13				Proximal	4 85	8 84	9 76	5 65	40		6 40
						_	_	_		_	1				-	_	_		-		
			Avg. stdev avg stdev Avg	80 10.7 78 9.95 76	5.73 82.5	13.9 68.9	6.28 46.6	16.3	į	17 2.83 17 4.81 16.5				Avg stdey avg stdey Avg	87.6	90 2.17 86 8.44 81.9	5.36 75.5	4.19 70 4.62 67.2	3.5 46 4.85 40.2		52 3.56 46 4.95 40.7
	ст	la la	stdev	9.95	5.73		6.28	4.74	ਬ	4.81		cm ²	las	stde	6.9	8.44	5.36	4.62	4.85	ھ	4.95
	2000 mJ/cm ²	Occlusal	avg	78	11.9 83	71	47	17	Gingival	11		6000 mJ/cm ²	Occlusal	avg	88	86	79	2	46	Gingival	46
	8	U	stdev	10.7	11.9	73 6.96	48 10.3 47	2.76		2.83		8	Ü	stdev	2.7	2.17	60.9		3.5		3.56
			Avg.		83			17						Avg	91	90	82	73	51		52
				92.3	95.4	81.3	59.8	19.1		19.4					91.1	88	88	78	55.3		26
			Axial	77.1 71.6 92.3	83.2 95.4	68.6 81.3	41.9 41.9 59.8	mush 15.2 19.1 17 2.76 17 4.74 16.3		5/max mush 15.4 19.4				Axial	88.5	90.6	75.9	6.69			50.8
				7.1	71.6	20	6.1	nsh		, ysni					93.9	92.3 90.6	80.7	72.1	48.6 50.2		5/max 49.2 50.8
				1	2	ا ا	4	5 ⊒		naxr					100	2	က	4	5		nax 4
					- 4		1			5/n						. •					5/1

Knoop Hardness

		t-test	0.98	0.46	0.02	0.48	0.58						-test	0.01	0.3	0.42	0.44	0.78		
			99.96	89.5		82.3			85.1				_	83	94.6 89.5 81.8	91.6 97.7 85.6 0.42	85.6	87.9 88.5 88.5		89.5
		Proximal	83	9.96	8	87.5	82.8						Proximal	86.5	89.5	97.7	96	88.5		89.5
		ď	85.2	87.2 96.6 89.5	84.5	83.8 87.5 82.3	76 60.6 82.8 84.1		77 61.3 83.8				ď	84.5 86.5	94.6	9.16	89.4	87.9		88.9
	-	٩vg	8	91	06	85	92	[11				Avg	87 8	89	92	06	88		88
m ²	_	Avg. stdey avg stdey Avg	5.8	6.4	4.9	2.7	13		13	^	E	=	Avg. stdey avg stdey Avg	2.3	6.4	6.1	5.3	0.3	_	86.6 90 4.3 90 0.3 89 88.9 89.5 89.5
36000 mJ/cm ²	Occlusal	avg s	90	90	91	98	9/	Gingival			60000 mJ/cm	Occlusal	avg	91	91	93	92	83	Gingival	90
3600	ŏ	stdev	3.4	2.6	4.4	5.6	11	ଓ	11 77		900	Ŏ	stdev	2.3	2.5	4.6	5.4	4.3	ပ	4.3
	10	Avg.	06	88	92	87	77		78				Avg.	92	93	93	94	83		90
			93.2	91.1	96	88.5	85.1 77 11 76		86.1					96	90	98.7 90.6	88	85.6		9.98
		Axial	36.5	89	92.7	91.6	81		82				Axial	96.6	94.9	98.7	93.8	88	5	88
		1	90.8 86.5	85.9	87.2 92.7	80.7 91.6	64.8		9.5					92.3 96.6	93.1 94.9	8.06	98.7	93.9	8	95
		19	<u>-</u> ای	2	ကျ	4 &	2		5/max 65.6					<u>-</u>	2	က	4	2		5/max
		t-test	0.62	0.35	0.16	0.4	0.88		5/1				t-test	0.26	0.86	0.2	7.	0.94		2/
		Ī	93.8 0.		88.5 0.	86 0	83.7 0.		7:				I	1	94.9		8			1.0
		mal	_	84.5 81.4 94.3				8	.8 84.7				mal	8 92.1	89 94	96.2 95.4 87.5	82.5 88.5 82.8	77.1 78.4 84.1		79.3 85.1
		Proximal	89	5 81	1 86	8 81.4	4 63	6	2 63.8				Proximal	88		2 95	5 88	1 78		
			85.2	\vdash	72.1	64.8	64.4		9 65.2				_	3.88.6	\$ 90.8	-	_	_	1	3] 78
		Avg. stdey avg stdey Avg	91 5.77 90 4.31 89.3	6.73 86.7	8.84 82.2	79 11.2 77.4	70.4		83.9 72 11.4 71 11.7 71.2				Ava. Istdey avg Istdey Avg	94 2.55 92 2.21 89.6	3.02 91.6	93	83 7.23 84 3.38 84.6	8.33 80 3.72 79.9		3 80.8
I/cm ²	sal	stde	4.31	6.73		11.2	11.6	/al	11.7	·	J/cm²	sal	stde	2.27	3.05	88 3.49 90 4.81	3.38	3.7	Val	3.78
24000 mJ/cm ²	Occlusal	y avg	98	88	98		2 71	Gingival	11 71		48000 mJ/cm ²	Occlusal	y avo	5 92	3.16 92	8	8	88	Gingival	3 81
24	Ĭ	stde	5.77	4.2	3.4	5.53	11.2		11.4		4		stde	2.5	3.16	3.4	7.2			8.4
		Avg		88	88	20	71		72				Ava		92			8		8
			9.96	93.2	92.7	85.6	82.8	5	83.9					91.1	94.9	88	86.5	87.5		88.5
		Axial	85.1	87.2 85.1 93.2	85.9 89.5 92.7	74.8 82.3 85.6	60.6 69.2 82.8 71 11.2 71 11.6 70.4		70				Axia	96.2 93.8 91.1	88.6 92.1 94.9	85.2 91.6	88	71.1 81.8 87.5		82.8
			91.6 85.1	87.2	85.9	74.8	9.09						∢	96.2	88.6	85.2	74.8	71.1		71.9
			-	7	က	4	5		5/max 61.3					4-	2	ന '	4	ر ا		5/max 71.9 82.8 88.5 81 8.43 81 3.78 80.8
									5											5

FTIR Actual and Interpolated interpolated

		t-test	0.92	0.24	0.59	0.61	0.64					I-test	0.14	0.38	0.81	0.32	0.35		
		7			51.3 0	45 0			53.7			1	56.3 0	58.1	55.5	51 0	48 0		29.3
		Proximal	2.3 54 52.1 53.2 56.5	50.3 54.1	48.8	46.9	38 5.9 36.1 1.9 35 35.4 36.2 32.5		59.8			Proximal	59.1	$\overline{}$		51.7	52		5/max 72.7 78.4 78.3 76 3.3 77.9 6.6 79 72.7 85.9 79.3
		Pro	52.1	51 (51 4	48.4	35.4		61.1			g E	56.5	57.7 56.1	55 53.6 54.9	43.9 51.7	44		72.7
	100	Avg	54	52	20	47 4	35		28			Avg	57	57	55	49	48		62
n ²	- 19	tde/	2.3	2	1.4	1.7	1.9		4	2	=	stdey,	1.6	1	1	4.3	4	_	9.9
4000 mJ/cm ²	Occlusal	avg b	54	52.6	49.3	47.2	36.1	Gingival	59.1	12000 m 1/cm ²	Occlusal	avg		56.8	54.8	20	2 47.2	Gingival	6.77
4000	ŏ	tde	1.1	1.4 52.6	5.2 49.3	2.1 47.2	5.9	Ö	8.1 59.1	1200	ŏ	tde	3.3 56.1	2	1.2 54.8	5.2	2	Ö	3.3
		Avg. stdey avg stdey Avg	54	53	48	48	38		90			Avg. stdey avg stdey Avg	55	99	55	20	46		9/
		_		54.1	53.1	47.8	41.7		68.9				52	58.6	55.9	55	47.4		78.3
		Axial	53.7 55.4 53.3	51.7 54.1	49.2	45.3	33.4 32.2. 41.7		53.2			Axial	58.4	55.3	53.5	51.6	47.5		78.4
			53.7	54.4	42.8	49.5	33.4						54	55	55.3	44.8	44		72.7
			· ·	2	က	4	r.		5/max 57.6				, -	7	က	4	ر. د	/max	
		t-test	0.24	6.0	0.43	0.28	0.37		Ŋ			t-test	0.28	0.15	0.23	0.04	0.46		ц
				1.99			22.7		37.5				55.5	49.5		47 45.8 46.7 49.5 0.04	39.5		35.2
		Proximal	1.6	1.3	6.1	9.2	1.4	9	35.4 37.5			Proximal	51		1.4	6.7	36		59.5 65.2
		Pro	52.6 51.6 50.2	4.1 5	43.4 46.1 54.7	43.6 39.2 31.3	25 31.1 21.4 22.7	2				P	52.5	56.4 54.7	8.3 5	5.8	41		1
		- BA		54 5	48 4	38 4	25 3		41 51.4			bA		54 5	51 4	47 4	39		64 6
=		devA	2.	4.2	5.9				8.7			devlA	2.3		2.3	2	2.6		4.2
2000 mJ/cm ²	Occlusal	Avg. stdey avg stdey Avg	54.1 52 55.4 54 1.72 52.7 1.2 51	49.2 54.1 59.3 54 5.04 54 2.4 54 54.1 51.3 56.1	49	43.9 41.7 41.1 42 1.47 40.1 6.2	5 27.9 5.3	Gingival		2000 - 1/2m ²	Occlusal	Avg. stdev avg stdev Avg	54 3.01 53.7 2.3	57 0.92 55.3 3.6	51 56.1 53 2.58 52.1 2.3 51 48.3 51.4 52.8	8.6	40	Gingival	99
2000	o	lev a	72 5	40		47 4	5 2	i <u>e</u>	26	900	30	dev a	01 5	92 5	58 5	49.4 48.5 51.9 50 1.77 48.6	39 42.6 41.6 41 1.86 40	S	05
		vg.İstc	4	5.	48.5 47.9 53.3 50 2.95	2	_		8.			vg. stc	4 ω	57 0	53 2	0.	1	Ī	38 3
		<u>4</u>	5.4	3.3	3.3	1.1	29.5 36.1 26.3 31	l	3.4			4	8.7		5.1	6.	1.6		8.7
		a	2 55	1 58	9 53	7 41	1 26		6 43			a	53.5 57.8	57.2 56.1	1 50	5,	6 4	-	3 68
		Axial	1 52	54	5 47	9 41	36		7 59			Axial		1	1 5	48	42		4 70
			54	49	48	43.8	29		5/max 48.7 59.6 43.4 51 8.26 46				52	58	53.1	49	39		5/max 64.4 70.3 68.7 68 3.05 66 4.2 64 67.
			~	2	က	4	S		5/max				~	8	က	4	2		5/ma

FTIR Actual and Interpolated

	t-test	0.5	0.46	0.02	0.48	0.4					t-test	0.84	0.3	0.42	0.44	0.59		
		28.5	_	58.8	-	55.2		91.1				30.3	53.5	55.1	55.1	9.99		93.4
	Proximal	59.7 58.5	59.8 56.8	22	55.9 53.7	15.7		75.5 91.1			Proximal	59.9 60.3	56.8 53.5	30.3	59.5	55.3 56.6		91.3
	P	52	55.8	54.7	54.4	46 45.7 55.2		9/			Pro	99	58.9	57.7 60.3 55.1	56.7 59.5 55.1	53		87.5 91.3 93.4
	1vg	22	57 5	57 5	4,		[81			Avg		56	58	57	25		9
- ₋	tdev /			2.1	1.1	5.4		8.9	m ₂		tde/	2.4		5.6	2.2		_	3
36000 mJ/cm² Occlusal	avg s	57.7	56.9	57.3	55.2	20	Gingival	82.6	60000 mJ/cm ²	Occlusal	avg b	58.9	1.1 57.3 2.7	58.1	57.8	54.7	Gingival	90.3
360	Avg. stdey avg stdey Avg	59 0.9 57.7 4.1	56 1.1 56.9 2.1	9.	2.4 55.2 1.1	3.8	Ō	6.3	9009	ŏ	Avg. stdey avg stdey Avg	59 1.5 58.9 2.4 59	1.1	2	2.3	0.5	Ö	90.6 90 0.7 90.3
69	Avg.	29	99	58	26	51					Avg.		28	58	58	54		8
		59.6	57.5	59.5	56.4	51.7 54.6 51 3.8 50 5.4 49		85.4 90.1 84				58.3	57	57.2	56.1 58 2.3 57.8 2.2 57	54.5 54.9 54 0.5 54.7 1.8		90.6
	Axial	58.1	9.99	58.1	57.7 56.4	51.7	e i	85.4			Axial	60.7	59.1	_	58.6	54.5		06
	4	58	55.3 56.6	55.8	53.1	47	E .	9.7				58	58.3	57.3 60.7	60.7	54		39.2
		 -	2	w Les	4	2		5/max 77.6) —	2	က က	4	5 		5/max 89.2
	t-test	66.0	35	16	0.48	0.21		2/			t-test	0.03	98.0	7.	0.7	0.43		2/
	Ī		53.3 58.8 0.35	4	55.3 0.			7.			Σ			55.9	53.9	1		92.5
	mal	57.2 60.6	3 58	3 56	3 55	45.5 46.6 52.5		75.1 76.9 86.7			imal	56.5 60.4 59.9	.6			54.9 56		9.
	Proximal	57	53	1 55	51.3	3 46		1 76			Proximal	5 60	3 56	59.6 59.3	56	54		84.2 90.6
	_		~	17									1	w.	∞	1		5
		54	54.7	49.4	1 54	_	1	_			0	-	3 57.		5 53.8 56.4	51	1	8
		54	56 54.7	7 54 49.4 55.3 56.4 0.16	-	48		80			ev Ava	-	3 58 57.3 56.6 59.1	58	55	54 51		3 89 8
հJ/cm² usal		54	2.9 56 54.7		-	3.8 48	ival	80	nJ/cm²	usal	o stdey Aval	-	7 1.3 58 57.	2 58	1.4 55	2.6 54 51	ival	2 4.3 89 84
4000 mJ/cm² Occlusal		54	56 2.9 56 54.7		-	49.2 3.8 48	Gingival	80	48000 mJ/cm ²	Occlusal	ev avo stdey Avo	-	4 57.7 1.3 58 57.3	2 58	1.4 55	2.6 54 51	Gingival	52 90.2 4.3 89 84
24000 mJ/cm² Occlusal		54	1.78 56 2.9 56 54.7		-	49.2 3.8 48	Gingival	80	48000 mJ/cm ²	Occlusal	a Istdey ava Istdey Ava	-	1.34 57.7 1.3	2 58	1.4 55	2.6 54 51	Gingival	1 0.62 90.2 4.3 89 84
24000 mJ/cm² Occlusal	Ava. Istdev ava Istdev Avg	57 1.16 57.3 3.3 57 54	56 1.78 56 2.9 56 54.7		-	49.2 3.8 48	Gingival	80	48000 mJ/cm ²	Occlusal	Availated availated Avail	57 2.67 58 2.1 59	1.34 57.7 1.3	2 58	1.4 55	2.6 54 51	Gingival	2 91 0.62 90.2 4.3 89 84
24000 mJ/cm² Occlusal	Availstdev avg stdev Avg	57 1.16 57.3 3.3 57 54	56 1.78 56 2.9 56 54.7		-	49.2 3.8 48	Gingival	80	48000 mJ/cm ²	Occlusal	Ī	58.3 57 2.67 58 2.1 59	1.34 57.7 1.3	2 58	1.4 55	2.6 54 51	Gingival	1 92 91 0.62 90.2 4.3 89 84
24000 mJ/cm² Occlusal		58.3 57.4 57 1.16 57.3 3.3 57 54	54.9 58.3 56 1.78 56 2.9 56 54.7		-	49.2 3.8 48	Gingival	80	48000 mJ/cm ²	Occlusal	Axial Avn stdey avo stdey Avo	58 9 58 3 57 2.67 58 2.1 59	57.9 59.1 58 1.34 57.7 1.3	57.7 55.3 56 1.48 57.1 2 58	1.4 55	2.6 54 51		3 91.1 92 <u>91 0.62 90.2 4.3 89</u> 84
24000 mJ/cm² Occlusal	Availstdev avg stdev Avg	57 1.16 57.3 3.3 57 54	54.9 58.3 56 1.78 56 2.9 56 54.7	56.8 58.1 57 1.44 55.2 3.7	-	49.2 3.8 48	Gingival	80	48000 mJ/cm ²	Occlusal	Ī	58.3 57 2.67 58 2.1 59	57.9 59.1 58 1.34 57.7 1.3	57.7 55.3 56 1.48 57.1 2 58	56.1 55.5 54 3.06 54.4 1.4 55	2.6 54 51		x 90.8 91.1 92 91 0.62 90.2 4.3 89 84
24000 mJ/cm² Occlusal	Availstdev avg stdev Avg	58.3 57.4 57 1.16 57.3 3.3 57 54	54.9 58.3 56 1.78 56 2.9 56 54.7		-	49.2 3.8 48		6.2 80	48000 mJ/cm ²	Occlusal	Ī	58 9 58 3 57 2.67 58 2.1 59	57.9 59.1 58 1.34 57.7 1.3	57.7 55.3 56 1.48 57.1 2 58	1.4 55	2.6 54 51		5/max 90.8 91.1 92 91 0.62 90.2 4.3 89 84

KHN		n = 3	F	lattened	templat	e; Data collected directly on end
	4000 r	nJ/cm ²		avg	st dev	
top [68.8	71.8	67.3	69.3	2.3	
bottom	0.0	6.2	0.0	2.1	3.6	2.84 % of max
_						
	6000 1	nJ/cm ²		avg		_
top	67.7	72.2	69.8	69.9	2.3	
bottom	16.0	17.3	15.9	16.4	0.8	22.6 % of max
_						
	1 0008	nJ/cm ²		avg		
top	65.5	71.2	70.5	69.1	3.1	
bottom	25.0	33.6	26.8	28.5	4.5	39.2 % of max
	12000	mJ/cm ²		avg		_
top	66.4	67.3	70.4	68.0	2.1	
bottom	39.2	39.7	39.1	39.3	0.3	54.1 % of max
-						
	24000	mJ/cm ²		avg		
top [69.3	66.9	70.1	68.8	1.7	I
bottom	53.2	52.6	52.9	52.9	0.3	72.8 % of max
_						
	72000	mJ/cm ²	Control	avg		
top [72.7	70.6	71.9	71.7	1.1	
bottom	69.4	71.6	69.5	70.2	1.2	96.5 % of max

DC		n = 3	Flatte	ened templa	ate; Data	collecte	d directly on end
	4000 n	nJ/cm ²		avg	st dev		
top	53.0	54.9	54.5	54.1	1.0		
bottom	9.8	13.2	19.8	14.3	5.1	24.5	% of max
	6000 n	nJ/cm ²		avg		_	
top	51.7	55.4	53.2	53.4	1.9		
bottom	21.9	30.7	25.9	26.2	4.4	45	% of max
	8000 n	nJ/cm ²		avg			
top	53.7	56.1	55.2	55.0	1.2		
bottom	33.2	33.4	33.5	33.4	0.2	57.3	% of max
	12000	mJ/cm ²		avg			
top	53.3	55.2	57.1	55.2	1.9	1	
bottom	41.8	39.0	37.3	39.4	2.3	67.6	% of max
	24000	mJ/cm ²		avg			
top	54.2	56.3	58.0	56.2	1.9		
bottom	47.2	45.5	46.9	46.5	0.9	80	% of max
	72000	mJ/cm ²	Control	avg		_	
top	57.3	56.8	58.2	57.4	0.7		
bottom	54.6	57.7	57.0	56.4	1.6	97	% of max

KNOOP HARDNESS TECHNIQUE STUDY

Purpose: To determine if the pre-operative use of graphite before Knoop hardness indentation can increase the readability and reduce the variability between evaluators of filar measurements.

Materials and Methods: Three evaluators were asked to determine the length of a Knoop hardness indentation at various occlusal-gingival increments in a specimen of Z250 resin composite. The 5 millimeter long resin composite specimens were created in a flattened tooth template (Figure 7) at 12000 mJ/cm² with Shade A-2 Z250, mounted in epoxy resin, sectioned and lightly polished as before. Graphite from a mechanical pencil was placed on one half of the increment. Two indentations were created side by side per increment using a 200 gram load with a 10 second dwell time. One indentation was made in graphite and one increment without. In addition, 5 indentations were made in the nearby mounting epoxy with and without graphite and evaluated by just one evaluator.

Results:

Z250: As can be seen in the tables on the following page, there was a significant increase in the average Knoop hardness number per increment between graphite and non-graphite coated surfaces. Also, there was a significant increase in the standard deviation without the use of graphite ($\alpha = 0.05$).

Epoxy: There was no significant difference in the average Knoop hardness numbers between graphite and non-graphite coated surfaces.

Discussion:

Z250: The use of graphite significantly reduced the variability between operators. The evaluators commented on the ease of readability of the graphite-coated surfaces. There was an apparent underestimation of the indentation without the use of graphite leading to a higher Knoop hardness number. Ideally a highly polished surface should be used to reduce the potential for measurement difficulties. The smoothest surface to is created against a milar strip or glass. However, after sectioning or sanding, it is much more difficult to produce a highly reflective surface. Care was taken in this study not to produce a high polish in the concern of possibly changing the mechanical properties.

Epoxy: Graphite did not have an effect on the actual measurements because it did not affect the Knoop hardness numbers in the highly reflective epoxy resin.

Z250 Knoop Hardness 3 evaluators

12000 mJ/cm²

1st Evaluator

2nd Evaluator

3rd Evaluator

depth		
(mm)	Graphite	Non-graphite
1	73.6	81.4
2	68.2	73.6
3	70.3	73.6
4	62.7	75.5
5	57.4	59

Occlusal					
Graphite	Non-graphite				
70.6	86.5				
67.1	81				
68.2	70.3				
60.7	70.3				
56.3	66.5				

Graphite	Non-graphite
76.7	95.4
69.2	107.8
68.2	92.1
64.3	88
59.5	81

Summary

	Graphite		
	Avg	St Dev	
1	73.6	3.1	
2	68.2	1.1	
3	68.9	1.2	
4	62.6	1.8	
5	57.7	1.6	

Non-graphite					
Avg	St Dev				
87.8	7.1				
87.5	18.0				
78.7	11.8				
77.9	9.1				
68.8	11.2				

	t-tests
0.014158	
0.000854	variance

Epoxy Resin

Knoop Hardness

Graphite	Non-graphite
11.8	12
12.9	12.9
12.3	12.3
12.9	13.1
12.6	12.6
12.5	12.6
0.5	0.4

avg st dev 0.78755

t-test

2-Way ANOVA Energy or Location on DC

Tests of Between-Subjects Effects

Dependent Variable: FTIR

Source	Type III Sum of Squares	df	Mean Square	F	Sig.
Corrected Model	2529.834 ^a	11	229.985	9.493	.000
Intercept	238290.422	1	238290.422	9835.648	.000
AXPROX	19.803	1	19.803	.817	.375
ENERGY	2455.989	5	491.198	20.275	.000
AXPROX * ENERGY	54.042	5	10.808	.446	.812
Error	581.453	24	24.227		
Total	241401.710	36		ĺ	
Corrected Total	3111.287	35		1	

a. R Squared = .813 (Adjusted R Squared = .727)

Estimated Marginal Means

2. ENERGY

Dependent Variable: FTIR

			95% Confide	ence Interval
ENERGY	Mean	Std. Error	Lower Bound	Upper Bound
6	65.967	2.009	61.819	70.114
12	77.883	2.009	73.736	82.031
24	81.150	2.009	77.003	85.297
36	82.617	2.009	78.469	86.764
48	90.200	2.009	86.053	94.347
60	90.333	2.009	86.186	94.481

Post Hoc Tests ENERGY

Multiple Comparisons

Dependent Variable: FTIR

Tukey HSD

	1. 1					
		Mean			050/ 054	
//\ ENEDOV	(I) ENEDOV	Difference	0.15	<u>.</u>	95% Confide	
(I) ENERGY	(J) ENERGY	(I-J)	Std. Error	Sig.	Lower Bound	Upper Bound
6	12	-11.917*	2.842	.004	-20.703	-3.130
	24	-15.183*	2.842	.000	-23.970	-6.397
	36	-16.650*	2.842	.000	-25.437	-7.863
	48	-24.233*	2.842	.000	-33.020	-15.447
	60	-24.367*	2.842	.000	-33.153	-15.580
12	6	11.917*	2.842	.004	3.130	20.703
	24	-3.267	2.842	.856	-12.053	5.520
	36	-4.733	2.842	.566	-13.520	4.053
	48	-12.317*	2.842	.003	-21.103	-3.530
	60	-12.450*	2.842	.002	-21.237	-3.663
24	6	15.183*	2.842	.000	6.397	23.970
	12	3.267	2.842	.856	-5.520	12.053
	36	-1.467	2.842	.995	-10.253	7.320
ę	48	- 9.050*	2.842	.041	-17.837	263
	60	-9.183*	2.842	.037	-17.970	397
36	6	16.650*	2.842	.000	7.863	25.437
	12	4.733	2.842	.566	-4.053	13.520
	24	1.467	2.842	.995	-7.320	10.253
	48	-7.583	2.842	.119	-16.370	1.203
	60	-7.717	2.842	.109	-16.503	1.070
48	6	24.233*	2.842	.000	15.447	33.020
	12	12.317*	2.842	.003	3.530	21.103
	24	9.050*	2.842	.041	.263	17.837
	36	7.583	2.842	.119	-1.203	16.370
	60	133	2.842	1.000	-8.920	8.653
60	6	24.367*	2.842	.000	15.580	33.153
	12	12.450*	2.842	.002	3.663	21.237
	24	9.183*	2.842	.037	.397	17.970
	36	7.717	2.842	.109	-1.070	16.503
	48	.133	2.842	1.000	-8.653	8.920

Based on observed means.

Homogeneous Subsets

^{*.} The mean difference is significant at the .05 level.

FTIR

Tukey HSD^{a,b}

1		Subset			
ENERGY	N	1	2	3	
6	6	65.967			
12	6		77.883		
24	6		81.150		
36	6		82.617	82.617	
48	6			90.200	
60	6			90.333	
Sig.	į.	1.000	.566	.109	

Means for groups in homogeneous subsets are displayed. Based on Type III Sum of Squares
The error term is Mean Square(Error) = 24.227.

- a. Uses Harmonic Mean Sample Size = 6.000.
- b. Alpha = .05.

2-Way ANOVA Energy or Location on KHN

Tests of Between-Subjects Effects

Dependent Variable: KNOOP

Source	Type III Sum of Squares	df	Mean Square	F	Sig.
Corrected Model	7023.334 ^a	11	638.485	8.560	.000
Intercept	185890.323	1	185890.323	2492.310	.000
AXPROX	113.423	1	113.423	1.521	.229
ENERGY	6751.429	5	1350.286	18.104	.000
AXPROX * ENERGY	158.482	5	31.696	.425	.827
Error	1790.053	24	74.586		
Total	194703.710	36			
Corrected Total	8813.388	35			

a. R Squared = .797 (Adjusted R Squared = .704)

Estimated Marginal Means

2. ENERGY

Dependent Variable: KNOOP

			95% Confide	ence Interval
ENERGY	Mean	Std. Error	Lower Bound	Upper Bound
6	46.367	3.526	39.090	53.643
12	65.300	3.526	58.023	72.577
24	71.483	3.526	64.207	78.760
36	77.317	3.526	70.040	84.593
48	80.933	3.526	73.657	88.210
60	89.750	3.526	82.473	97.027

Post Hoc Tests ENERGY

Multiple Comparisons

Dependent Variable: KNOOP

Tukey HSD

		Mean Difference			95% Confide	ence Interval
(I) ENERGY	(J) ENERGY	(l-J)	Std. Error	Sig.	Lower Bound	Upper Bound
6	12	-18.933*	4.986	.010	-34.350	-3.516
	24	-25.117*	4.986	.000	-40.534	-9.700
	36	-30.950*	4.986	.000	-46.367	-15.533
	48	-34.567*	4.986	.000	-49.984	-19.150
	60	-43.383*	4.986	.000	-58.800	-27.966
12	6	18.933*	4.986	.010	3.516	34.350
	24	-6.183	4.986	.813	-21.600	9.234
	36	-12.017	4.986	.192	-27.434	3.400
	48	-15.633*	4.986	.046	-31.050	216
	60	-24.450*	4.986	.001	-39.867	-9.033
24	6	25.117*	4.986	.000	9.700	40.534
	12	6.183	4.986	.813	-9.234	21,600
	36	-5.833	4.986	.846	-21.250	9.584
	48	-9.450	4.986	.429	-24.867	5.967
	60	-18.267*	4.986	.014	-33.684	-2.850
36	6	30.950*	4.986	.000	15.533	46.367
	12	12.017	4.986	.192	-3.400	27.434
	24	5.833	4.986	.846	-9.584	21.250
	48	-3.617	4.986	.977	-19.034	11.800
	60	-12.433	4.986	.166	-27.850	2.984
48	6	34.567*	4.986	.000	19.150	49.984
	12	15.633*	4.986	.046	.216	31.050
	24	9.450	4.986	.429	-5.967	24.867
	36	3.617	4.986	.977	-11.800	19.034
	60	-8.817	4.986	.503	-24.234	6.600
60	6	43.383*	4.986	.000	27.966	58.800
	12	24.450*	4.986	.001	9.033	39.867
	24	18.267*	4.986	.014	2.850	33.684
	36	12.433	4.986	.166	-2.984	27.850
	48	8.817	4.986	.503	-6.600	24.234

Based on observed means.

Homogeneous Subsets

^{*.} The mean difference is significant at the .05 level.

KNOOP

Tukey $\mathsf{HSD}^{\mathsf{a},\mathsf{b}}$

			et		
ENERGY	N	1	2	3	4
6	6	46.367			
12	6		65.300		
24	6		71.483	71.483	
36	6		77.317	77.317	77.317
48	6			80.933	80.933
60	6				89.750
Sig.		1.000	.192	.429	.166

Means for groups in homogeneous subsets are displayed. Based on Type III Sum of Squares
The error term is Mean Square(Error) = 74.586.

- a. Uses Harmonic Mean Sample Size = 6.000.
- b. Alpha = .05.

1-Way ANOVA; Energy vs KHN and Energy vs DC

Between-Subjects Factors

		N
ENERGY	12	3
}	24	3
	4	3
	6	3
l	8	3
	con	3

Descriptive Statistics

Dependent Variable: KHN

ENERGY	Mean	Std. Deviation	N
12	39.333	.321	3
24	52.900	.300	3
4	2.067	3.580	3
6	16.400	.781	3
8	28.467	4.536	3
con	70.167	1.242	3
Total	34.889	23.287	18

Tests of Between-Subjects Effects

Dependent Variable: KHN

Source	Type III Sum of Squares	df	Mean Square	F	Sig.
Corrected Model	9147.171 ^a	5	1829.434	307.181	.000
Intercept	21910.222	1	21910.222	3678.955	.000
ENERGY	9147.171	5	1829.434	307.181	.000
Error	71.467	12	5.956		
Total	31128.860	18			ľ
Corrected Total	9218.638	17			

a. R Squared = .992 (Adjusted R Squared = .989)

Post Hoc Tests ENERGY

Multiple Comparisons

Dependent Variable: KHN

Tukey HSD

		Mean				
		Difference			95% Confide	
(I) ENERGY	(J) ENERGY	(l-J)	Std. Error	Sig.	Lower Bound	Upper Bound
12	24	-13.567*	1.993	.000	-20.260	-6.874
	4	37.267*	1.993	.000	30.574	43.960
	6	22.933*	1.993	.000	16.240	29.626
	8	10.867*	1.993	.002	4.174	17.560
	con	-30.833*	1.993	.000	-37.526	-24.140
24	12	13.567*	1.993	.000	6.874	20.260
	4	50.833*	1.993	.000	44.140	57.526
	6	36.500*	1.993	.000	29.807	43.193
	8	24.433*	1.993	.000	17.740	31.126
	con	-17.267*	1.993	.000	-23.960	-10.574
4	12	-37.267*	1.993	.000	-43.960	-30.574
	24	-50.833*	1.993	.000	-57.526	-44.140
ļ	6	-14.333*	1.993	.000	-21.026	-7.640
	8	-26.400*	1.993	.000	-33.093	-19.707
	con	-68.100*	1.993	.000	-74.793	-61.407
6	12	-22.933*	1.993	.000	-29.626	-16.240
	24	-36.500*	1.993	.000	-43.193	-29.807
	4	14.333*	1.993	.000	7.640	21.026
	8	-12.067*	1.993	.001	-18.760	-5.374
	con	-53.767*	1.993	.000	-60.460	-47.074
8	12	-10.867*	1.993	.002	-17.560	-4.174
	24	-24.433*	1.993	.000	-31.126	-17.740
	4	26.400*	1.993	.000	19.707	33.093
	6	12.067*	1.993	.001	5.374	18.760
	con	-4 1.700*	1.993	.000	-48.393	-35.007
con	12	30.833*	1.993	.000	24.140	37.526
	24	17.267*	1.993	.000	10.574	23.960
	4	68.100*	1.993	.000	61.407	74.793
	6	53.767*	1.993	.000	47.074	60.460
	8	41.700*	1.993	.000	35.007	48.393

Based on observed means.

Homogeneous Subsets

^{*.} The mean difference is significant at the .05 level.

KHN

Tukey HSD^{a,b}

ENERGY	N	1	2	3	4	5	6
4	3	2.067					
6	3		16.400		h .		
8	3			28.467		i i	
12	3				39.333		
24	3					52.900	
con	3						70.167
Sig.		1.000	1.000	1.000	1.000	1.000	1.000

Means for groups in homogeneous subsets are displayed. Based on Type III Sum of Squares
The error term is Mean Square(Error) = 5.956.

- a. Uses Harmonic Mean Sample Size = 3.000.
- b. Alpha = .05.

Univariate Analysis of Variance

Between-Subjects Factors

		N
ENERGY	12	3
	24	3
	4	3
	6	3
	8	3
	соп	3

Descriptive Statistics

Dependent Variable: DC

ENERGY	Mean	Std. Deviation	N
12	39.367	2.272	3
24	46.533	.907	3
4	14.267	5.085	3
6	26.167	4.406	3
8	33.367	.153	3
con	56.433	1.626	3
Total	36.022	14.256	18

Tests of Between-Subjects Effects

Dependent Variable: DC

Source	Type III Sum of Squares	df	Mean Square	F	Sig.
Corrected Model	3347.311 ^a	5	669.462	74.495	.000
Intercept	23356.809	1	23356.809	2599.051	.000
ENERGY	3347.311	5	669.462	74.495	.000
Error	107.840	12	8.987		
Total	26811.960	18			
Corrected Total	3455.151	17			

a. R Squared = .969 (Adjusted R Squared = .956)

Post Hoc Tests ENERGY

Multiple Comparisons

Dependent Variable: DC

Tukey HSD

		Mean Difference			95% Confide	ence Interval
(I) ENERG		(I-J)	Std. Error	Sig.	Lower Bound	Upper Bound
12	24	-7.167	2.448	.102	-15.388	1.055
	4	25.100*	2.448	.000	16.878	33.322
	6	13.200*	2.448	.002	4.978	21.422
	8	6.000	2.448	.213	-2.222	14.222
	con	-17.067*	2.448	.000	-25.288	-8.845
24	12	7.167	2.448	.102	-1.055	15.388
	4	32.267*	2.448	.000	24.045	40.488
	6	20.367*	2.448	.000	12.145	28.588
	8	13.167*	2.448	.002	4.945	21.388
	con	-9.900*	2.448	.016	-18.122	-1.678
4	12	-25.100*	2.448	.000	-33.322	-16.878
	24	-32.267*	2.448	.000	-40.488	-24.045
	6	-11.900*	2.448	.004	-20.122	-3.678
	8	-19.100*	2.448	.000	-27.322	-10.878
	con	-42.167*	2.448	.000	-50.388	-33.945
6	12	-13.200*	2.448	.002	-21.422	-4.978
	24	-20.367*	2.448	.000	-28.588	-12,145
	4	11.900*	2.448	.004	3.678	20.122
	8	-7.200	2.448	.100	-15.422	1.022
	con	-30.267*	2.448	.000	-38.488	-22.045
8	12	-6.000	2.448	.213	-14.222	2.222
	24	-13.167*	2.448	.002	-21.388	-4.945
	4	19.100*	2.448	.000	10.878	27.322
	6	7.200	2.448	.100	-1.022	15.422
	con	-23.067*	2.448	.000	-31.288	-14.845
con	12	17.067*	2.448	.000	8.845	25.288
	24	9.900*	2.448	.016	1.678	18.122
	4	42.167*	2.448	.000	33.945	50.388
	6	30.267*	2.448	.000	22.045	38.488
	8	23.067*	2.448	.000	14.845	31.288

Based on observed means.

Homogeneous Subsets

 $^{^{\}star}.$ The mean difference is significant at the .05 level.

DC

Tukey HSD^{a,b}

				Subset		
ENERGY	N	1	2	3	4	5
4	3	14.267				
6	3		26.167			
8	3		33.367	33.367		
12	3			39.367	39.367	
24	3				46.533	
con	3					56.433
Sig.		1.000	.100	.213	.102	1.000

Means for groups in homogeneous subsets are displayed. Based on Type III Sum of Squares
The error term is Mean Square(Error) = 8.987.

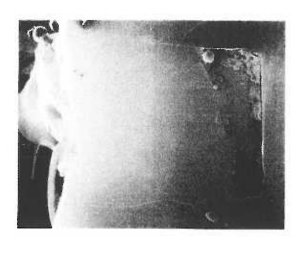
- a. Uses Harmonic Mean Sample Size = 3.000.
- b. Alpha = .05.

PART TWO FIRST PILOT

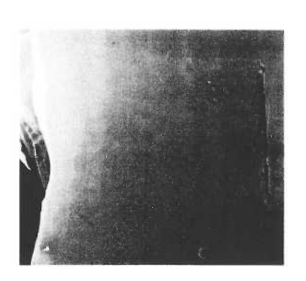
First Pilot Study

- 5 mm prep - dentin cervical margin

- 4000 mJ/cm^2 - 60% of max cure

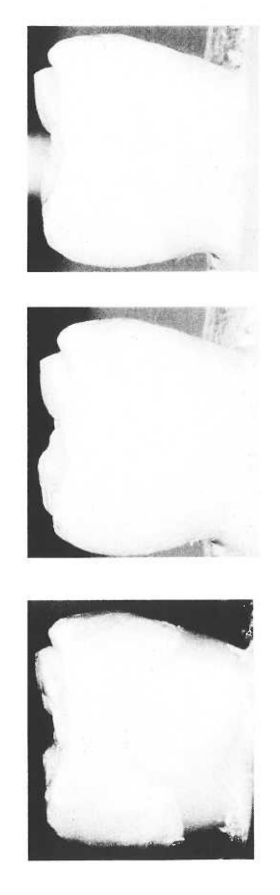






- 5 mm prep - dentin cervical margin

 $-4000 \text{ mJ/cm}^2 - 60\% \text{ of max cure}$



Post-thermal

Post-fatigue



- 5 mm prep

- dentin cervical margin

 $-4000 \,\mathrm{mJ/cm^2}$

- 60% of max cure

	Pre-op	Post-thermal	Post-fatigue
_	papple	49.24	16.17
7	0	19.08	34.26
<i>с</i>	0	23.49	0
4	23	31.8	0
2	15.76	40.62	0
9	0	17.13	0
7	3.43	14.19	0
80	0	13.9	0
6	0	15.19	3.43
10	0	22.51	0
=	0	75.37	0
12	0	2.45	0
5	0	87.09	0
14	pubble	propple	0
15	propple	propple	0
16	prophe	elqqnq	0
17	0	propple	0
18	0	piqqnq	0
19	0	11.74	0
20	0	13.7	2.16
21	0	16.15	9.3
22	0	34.3	11.74
23	0	17.13	0
24	0	14.68	0
25	0	17.15	0
avg.	2.0	26.8	3.1
std dev	5.9	21.6	7.7
% dapped	10	100	24

- 5 mm prep - dentin margins

- 12000 mJ/cm^2 - 80% of max cure



Pre-operative 12X

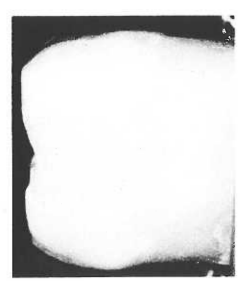
Post-thermal 12X

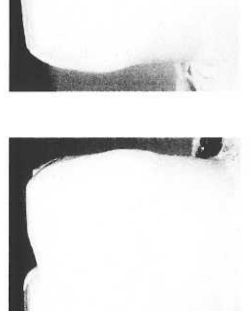
Post-fatigue 12X

- 5 mm prep - dentin margins

- 12000 mJ/cm^2 - 80% of max cure







	Pre-op	Post-thermal	Post-fatigue
-	7.4	2.5	10.2
7	8.9	7.6	7.0
က	6.3	5.1	2.6
4	2.7	2.3	2.2
ις	0.0	0.0	6.3
9	7.9	0.0	8.9
	10.6	11.8	6.8
80	17.5	8.9	0.0
တ	4.5	0.7	1.9
5	4.	0.0	1.3
7	5.1	2.5	8.9
12	5.5	1.7	10.1
13	8.2	10.8	9.5
4	10.0	4.4	12.0
15	0.0	1.3	1.3
16	1.5	0.0	10.8
17	2.7	6.3	4.6
18	0.0	3.8	0.0
10 0	3.2	1.9	2.2
20	1.0	0.0	4.4
21	12.7	3.2	14.8
22	2.1	4.4	3.9
23	3.4	6.0	5.7
24	1.4	0.0	0.0
25	1.4	4.4	2.6
avg.	5.1	3.4	5.5
std dev	4.6	3.4	4.2
% dapped	88.0	76.0	88.0

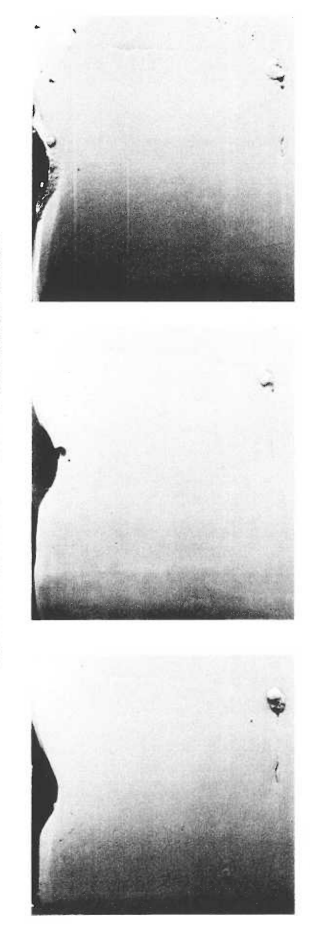
- 5 mm prep - dentin margins

 $-12000 \,\mathrm{mJ/cm^2}$

- 80% of max cure

- 5 mm prep - dentin margins

- 48000 mJ/cm² - 90% of max cure



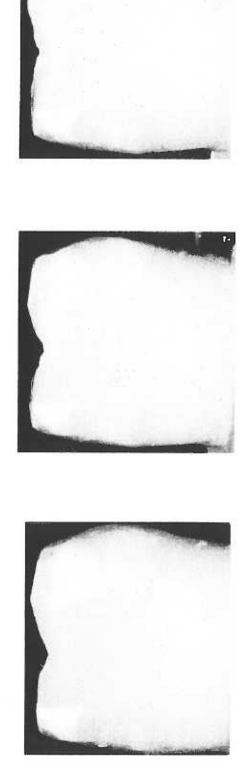
Post-thermal 12X

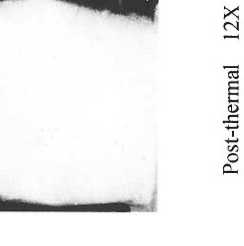
Post-fatigue 12X

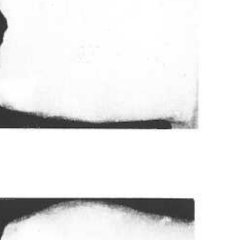


- 5 mm prep - dentin margins

- 48000 mJ/cm² - 90% of max cure







Pre-operative 12X

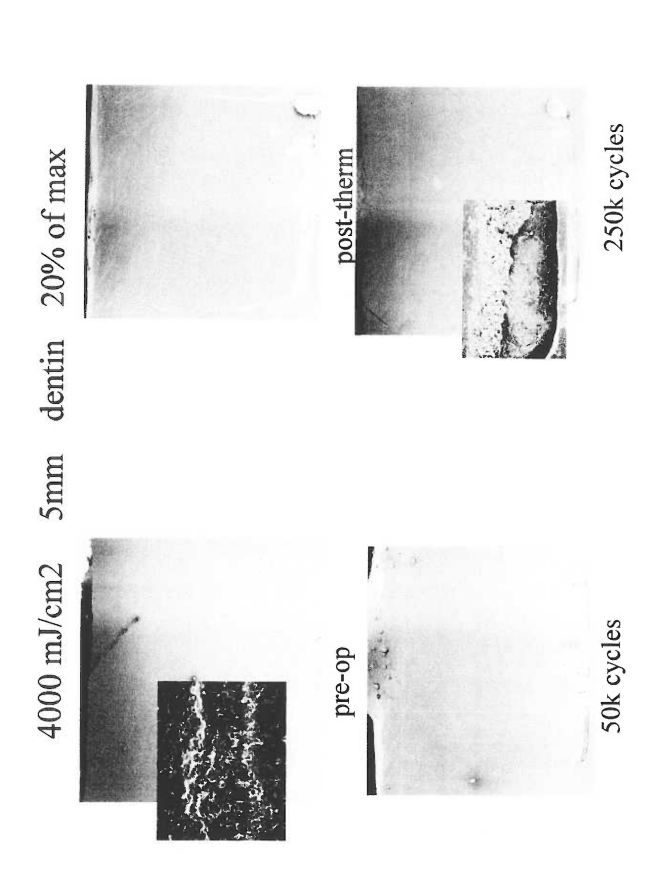
Post-fatigue 12X

	Pre-op	Post-thermal	Post-fatigue
_	7.5	2.8	4.8
2	pubble	propple	elqqnq
8	pubble	pripple	papple
4	pubble	papple	elqqnq
r.c	5.1	59.6	56.3
9	27.9	19.6	24.2
	0.0	3.2	7.0
80	5.1	17.0	0.0
တ	8.2	7.0	6.7
10	5.7	4.4	7.6
=	13.9	13.8	13.7
12	24.0	34.9	11.4
13	3.8	20.3	22.8
14	14.5	22.2	19.0
15	0.0	7.0	12.7
16	1.9	4.4	0.0
17	0.0	4.4	3.8
18	1.9	0.0	3.8
19	0.0	3.8	0.0
8	4.4	0.7	9.7
21	0.0	6.0	0.0
22	1.3	0.0	0.0
23	0.0	0.0	0.0
24	0.0	6.7	0.0
25	0.0	3.8	6.3
avg	5.7	10.5	9.4
std dev	7.9	14.3	12.8
bennen %	63.0	86.0	0.89

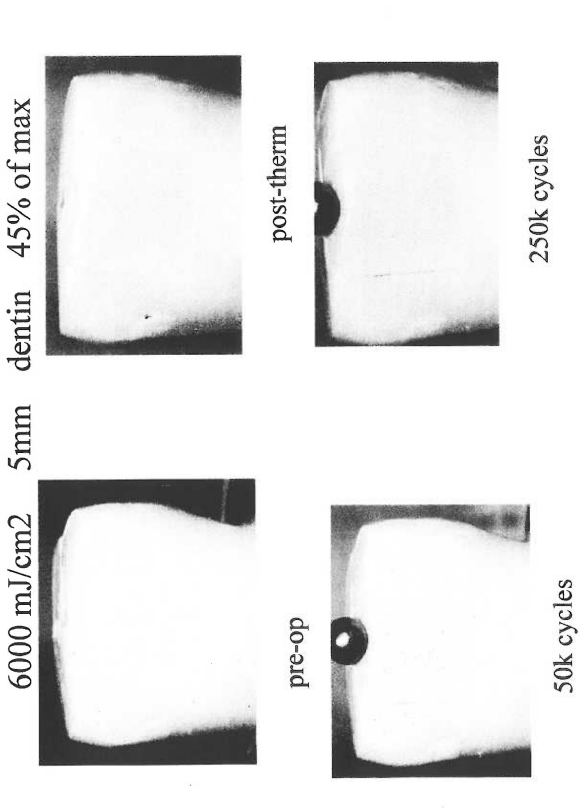
- 5 mm prep
- dentin margins
- 48000 mJ/cm²
- 90% of max cure

PART TWO SECOND PILOT

4000 mJ/cm2 5mm dentin 20% of max post-therm 250k cycles 50k cycles pre-op



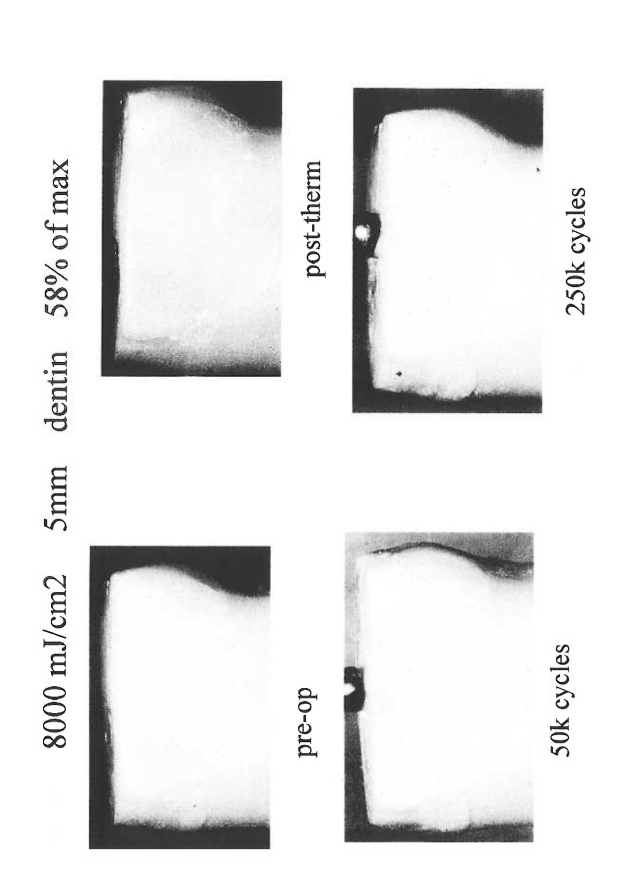
dentin 20% of max	50K cyc 250K cyc	bubble 80.1	17.9 40.8	14.1 5.1	20.9 3.0	19.6 9.7	24.1 0.0	21.7 0.0	21.4 0.0	0.0 0.0	6.3 0.0	14.1 9.3	0.0 4.6	27.2 35.5	27.2 0.0	0.0 0.0	0.0 0.0	0.0 0.0	0.0 0.0	17.1 6.2	15.7 9.5	15.5 10.1	18.2 6.1	19.3 5.2	13.5 24.5	22.6 17.7	14.0 10.7	007
5mm denti	Post-thermal 50K	bubble buk	bubble 17	22.8	0.0	20.7 19	16.0 24	13.1 21	5.9 21	2.5	16.2 6	0.0	0.0	0.5	9.7 27	0.0	0.0	0 2.0	4.4		22.3 15	22.8 15	16.0 18	0.0	0.0	0.0	7.5	76
)00 mJ/cm2	Pre-op	1 bubble	0.0	3 0.0	0.0	0.0	0.0	0.0	0.0	0.0	10 0.0	11 0.0	12 0.0	13 0.0	14 0.0	15 0.0	16 0.0	17 0.0	18 3.8	19 0.0	20 0.0	21 0.0	22 0.0	23 0.0	24 0.0	25 0.0		std dev 0.8

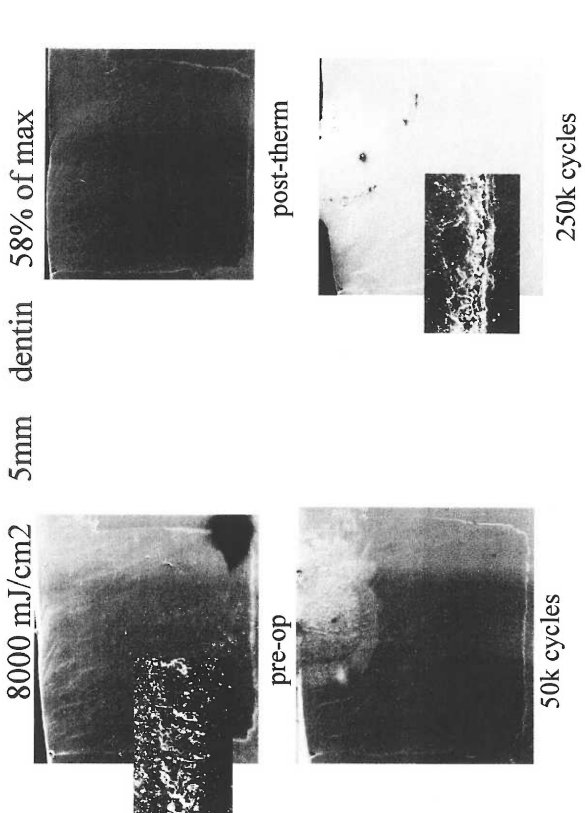


250k cycles 6000 mJ/cm2 5mm dentin 45% of max postpre-op 50k cycles

6000 mJ/cm2 5mm dentin 45% of max

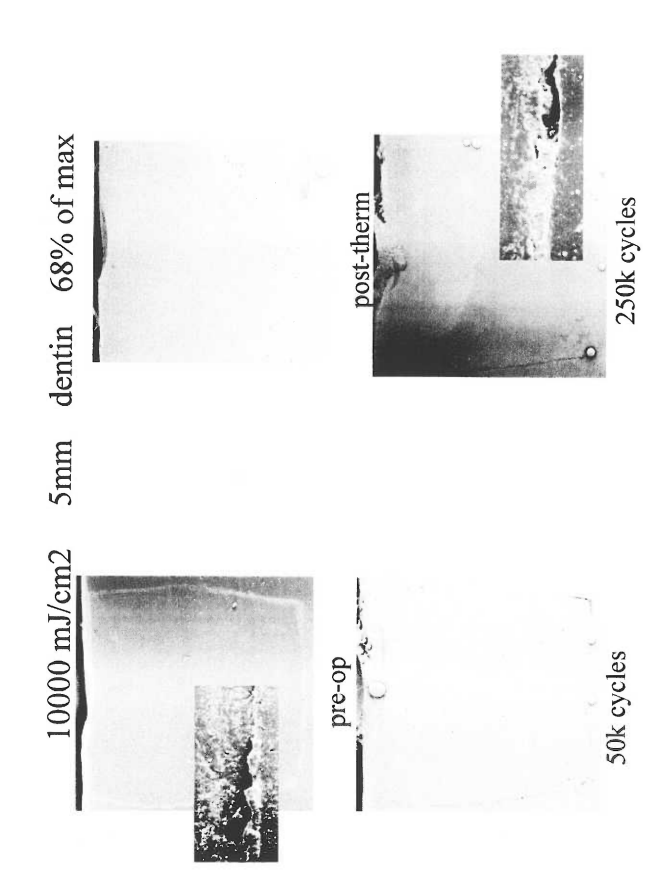
	Pre-op	Post-thermal	50K cyc	250K cyc
-	0.0	0.0	0.0	0.0
N	0.0	0.0	0.0	0.0
က	0.0	0.0	0.0	0.0
4	0.0	0.0	8.2	12.4
2	0.0	0.0	0.0	0.0
9	0.0	0.0	0.0	0.0
	0.0	0.0	0.0	0.0
80	1.0	0.0	0.0	0.0
თ	3.6	3.6	0.0	8.4
0	0.0	3.2	0.0	2.9
1-	0.0	0.0	0.0	0.0
12	0.0	0.0	0.0	0.0
5	0.0	0.0	3.5	7.3
4	0.0	0.0	0.0	0.0
15	0.0	0.0	2.9	0.0
16	0.0	0.0	2.9	0.0
17	0.0	0.0	6.3	0.0
18	3.8	0.0	0.0	0.0
19	1.0	4.8	11.4	0.0
50	0.0	6.0	0.0	0.0
21	0.0	0.0	0.0	0.0
22	0.0	0.0	0.0	0.0
23	0.0	0.0	0.0	5.4
24	0.0	0.0	0.0	4.4
25	0.0	0.0	25.3	15.6
avg.	0.4	0.7	2.5	2.1
std dev	1.0	1.7	5.7	4.2
% dapped	16.0	16.0	28.0	28.0





8000 mJ/cm2	cm2	5mm d	dentin	58% of max
	Pre-op	Post-thermal	50K cyc	250K cyc
-	0.0	0.0		3.8
0	0.0	0.0	0.0	0.0
ო	0.0	0.0	0.0	0.0
4	0.0	0.0	2.2	7.3
2	0.0	0.0	0.0	8.9
9	0.0	0.0	8.6	0.0
7	0.0	0.0	0.0	0.0
80	0.0	0.0	0.0	0.0
6	0.0	0.0	0.0	0.0
10	0.0	0.0	0.0	0.0
-	0.0	6.7	0.0	2.5
12	1.5	0.0	0.0	0.0
13	0.0	0.0	0.0	0.0
4	0.0	0.0	0.0	0.0
15	0.0	0.0	0.0	0.0
16	0.0	0.0	0.0	0.0
17	0.0	0.0	0.0	4.7
18	0.0	0.0	0.0	0.0
19	0.0	0.0	0.0	0.0
20	0.0	0.0	0.0	0.0
21	0.0	0.0	0.0	0.0
22	0.0	0.0	0.0	0.0
23	0.0	0.0	0.0	0.0
24	0.0	0.0	0.0	0.0
25	0.0	0.0	0.0	0.0
avg.	0.1	0.3	0.5	1.0
std dev	0.3	1.3	1.8	2.2
% ganned	0 1	0,	0	007

10000 mJ/cm2 5mm dentin 68% of max post-therm 250k cycles 50k cycles pre-op

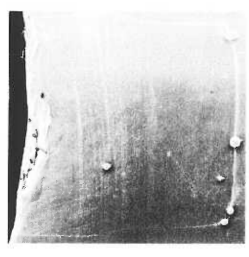


68% of max
dentin
5mm
mJ/cm2
100001

	71110/			
	Pre-op	Post-thermal	50K cyc	250K cyc
-	7.2	16.5	089	68.0
7	0.0	0.0	53.8	53.8
ო	8.2	3.8	0.0	0.0
4	0.0	0.0	0.0	9.9
2	0.0	0.0	5.1	0.0
ဖ	0.0	2.2	prople	0.0
	0.0	3.5	0.0	0.0
ω	0.0	0.0	0.0	0.0
တ	0.0	9.2	0.0	0.0
0	1.8	0.0	0.0	5.5
7	0.0	0.0	0.0	1.1
12	25.7	9.5	0.0	0.0
13	0.0	3.2	0.0	0.0
4	0.0	0.0	0.0	0.0
5	0.0	0.0	0.0	0.0
9	0.0	0.0	0.0	2.1
17	8.6	5.7	0.0	7.4
18	0.0	4.8	0.0	0.0
19	0.0	0.0	0.0	4.3
50	0.0	0.0	0.0	0.0
21	0.0	0.0	3.9	0.0
75	0.0	11.4	0.0	3.5
73	0.0	0.0	0.0	0.0
24	0.0	0.0	18.2	0.0
25	0.0	11.8	20.9	20.9
avg.	2.1	3.2	4.4	6.9
std dev	5.6	4.7	12.2	17.0
gabbed %	20.0	44.0	24.0	40.0

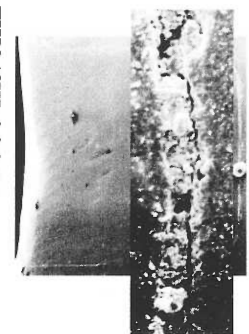
12000 mJ/cm2 5mm dentin 71% of max 250k cycles post-therm 50k cycles pre-op

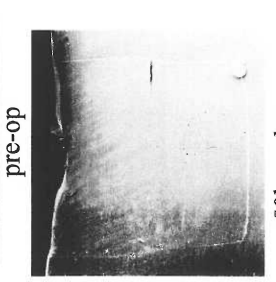
12000 mJ/cm2 5mm dentin 71% of max



post-therm

250k cycles



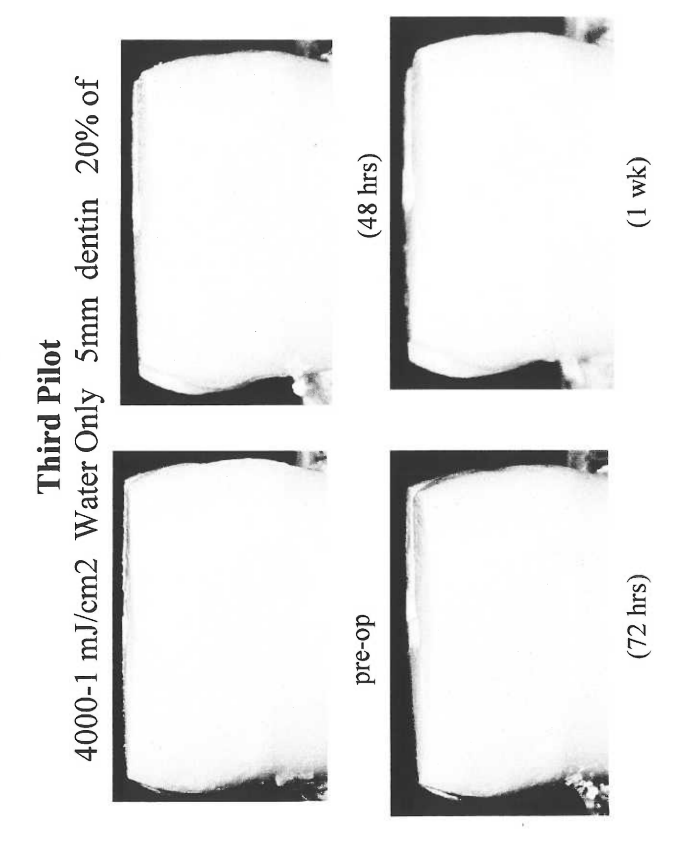


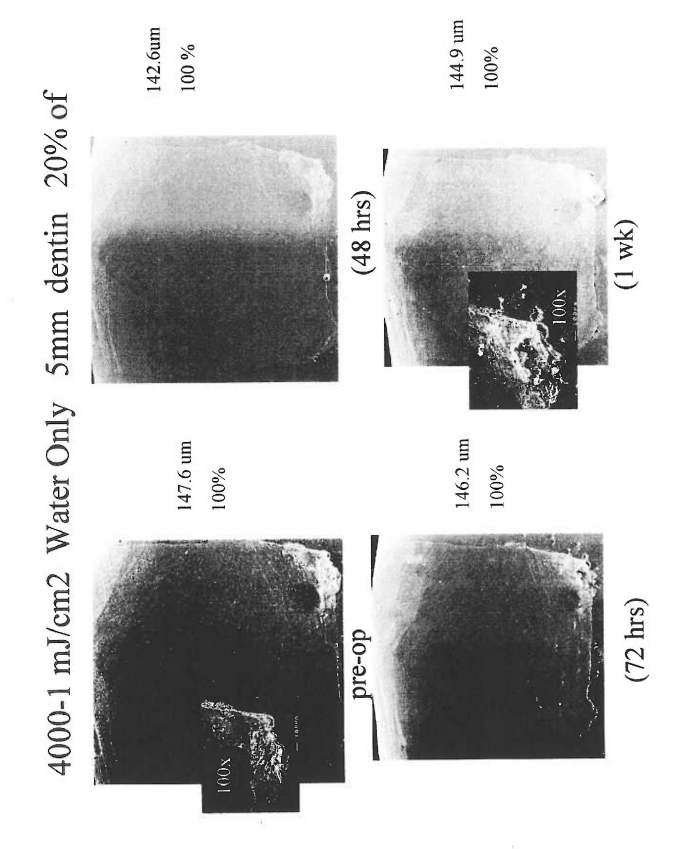
50k cycles

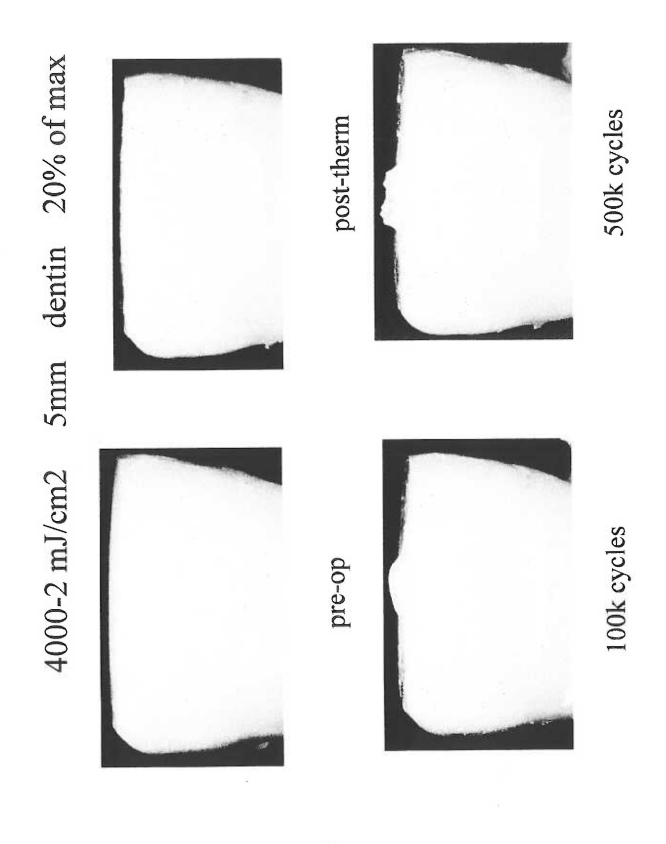
of max	
71% of	250K cyc
lentin	50K cyc
5mm c	Post-thermal
mJ/cm2	Pre-op
12000	

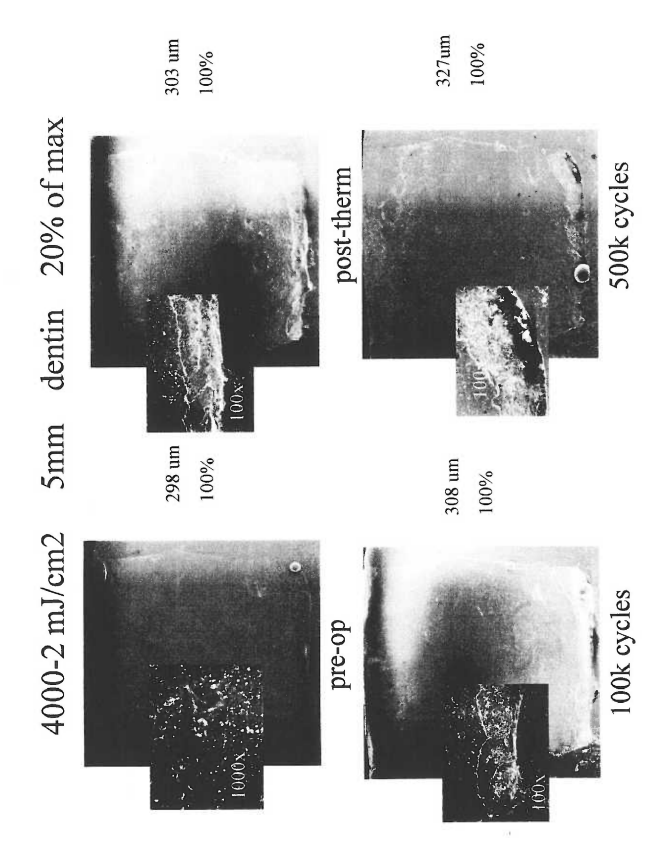
	Pre-op	Post-thermal	50K cyc	250K cyc
_	3.5	0.0	0.0	0.0
7	8.2	0.0	pubble	0.0
က	2.1	0.0	0.0	0.0
4	0.0	0.0	0.0	0.0
2	0.0	0.0	5.1	0.0
9	0.0	0.0	propple	0.0
7	0.0	0.0	0.0	0.0
ω	2.1	0.0	0.0	0.0
6	0.0	0.0	0.0	1.2
0	1.7	0.0	0.0	0.0
7	0.0	0.0	0.0	0.0
12	0.0	0.0	0.0	0.0
13	3.6	0.0	0.0	0.0
14	1.2	0.0	0.0	0.0
15	0.0	0.0	0.0	0.0
16	2.2	0.0	0.0	0.0
17	pripple	0.0	0.0	0.0
18	pripple	0.0	0.0	0.0
19	16.1	0.0	0.0	0.0
20	4.1	0.0	0.0	0.0
21	0.0	0.0	0.0	0.0
22	0.0	0.0	0.0	0.0
23	0.0	3.8	0.0	0.0
24	17.4	0.0	0.0	0.0
25	1.4	0.0	0.0	elqqnq
avg.	2.8	0.2	0.2	0.1
std dev	4.8	0.8	1.1	0.2
% gapped %	48.0	4.0	4.0	4.0

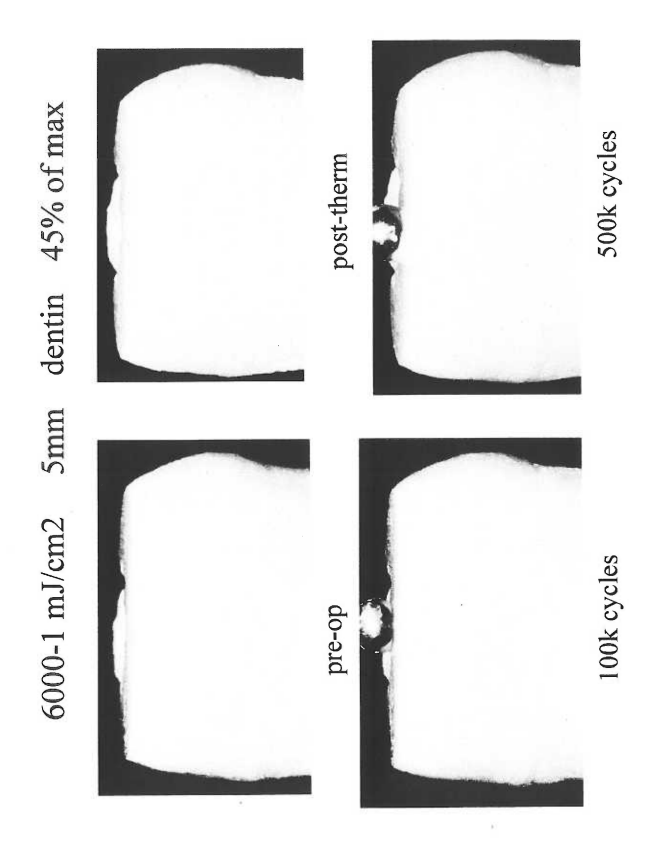
PART TWO THIRD PILOT

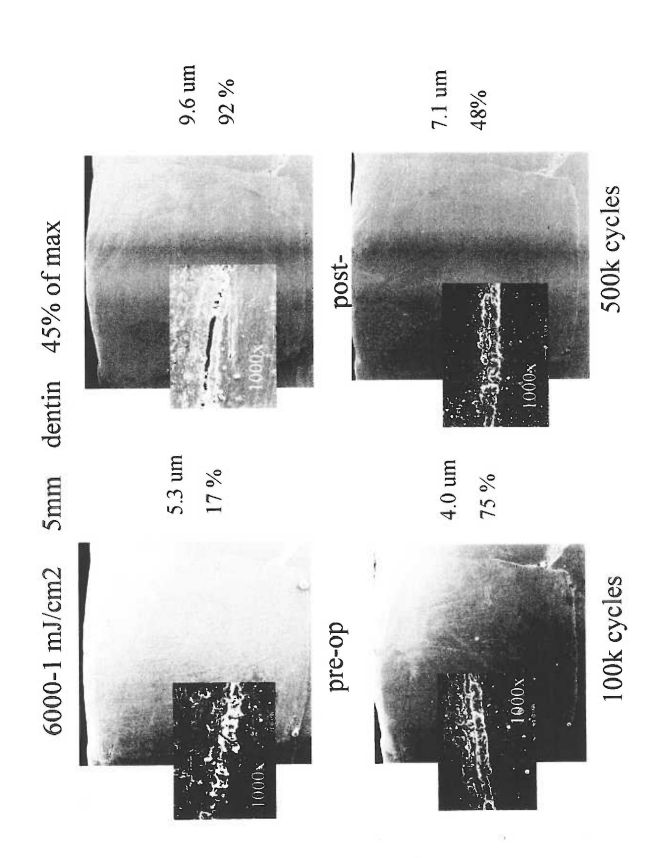


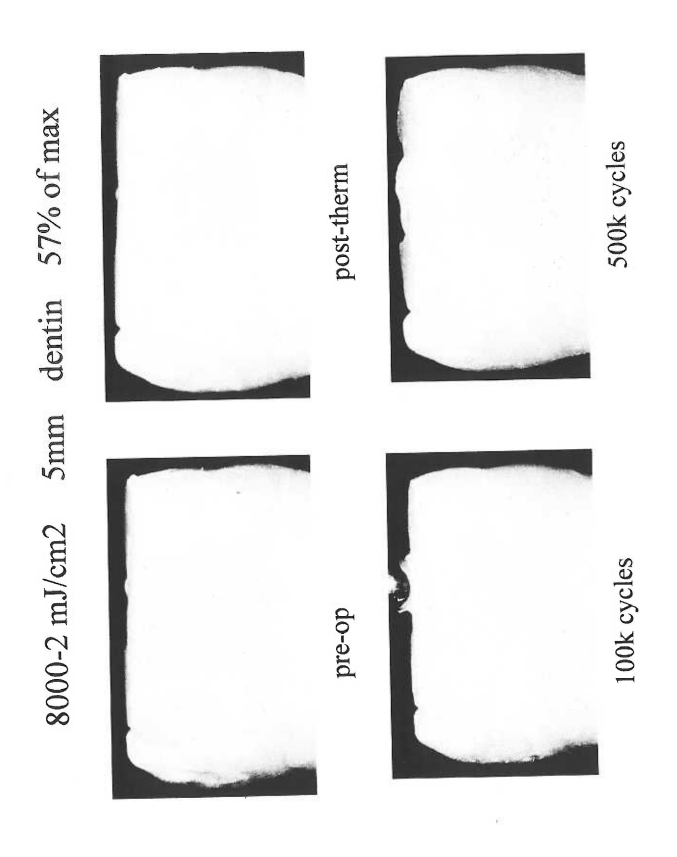


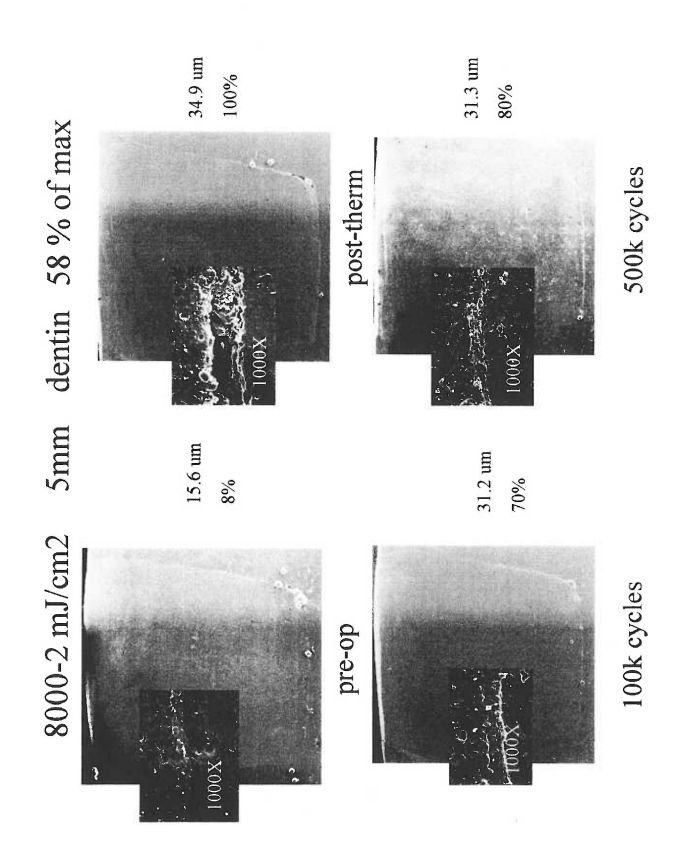


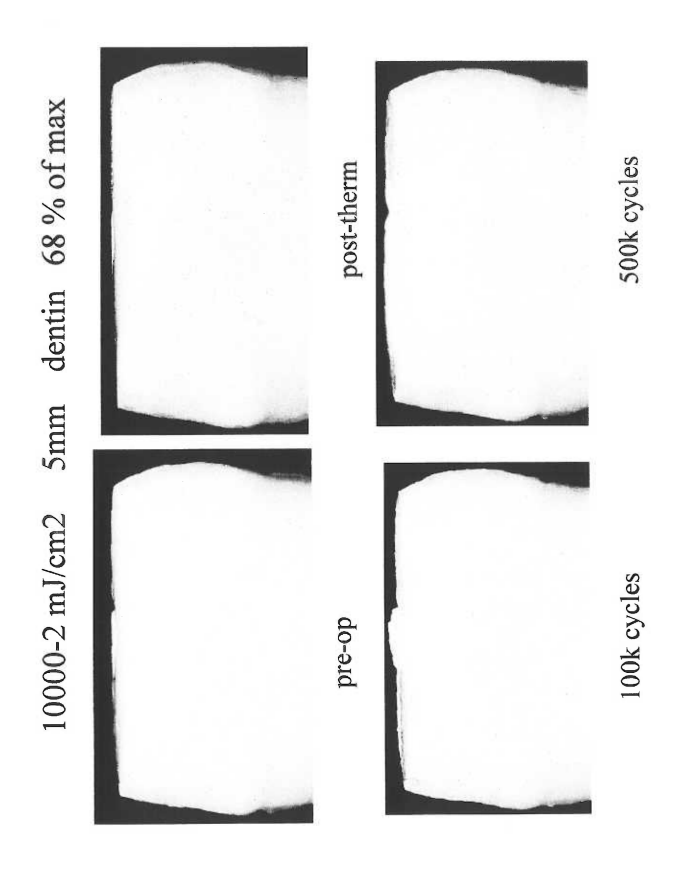


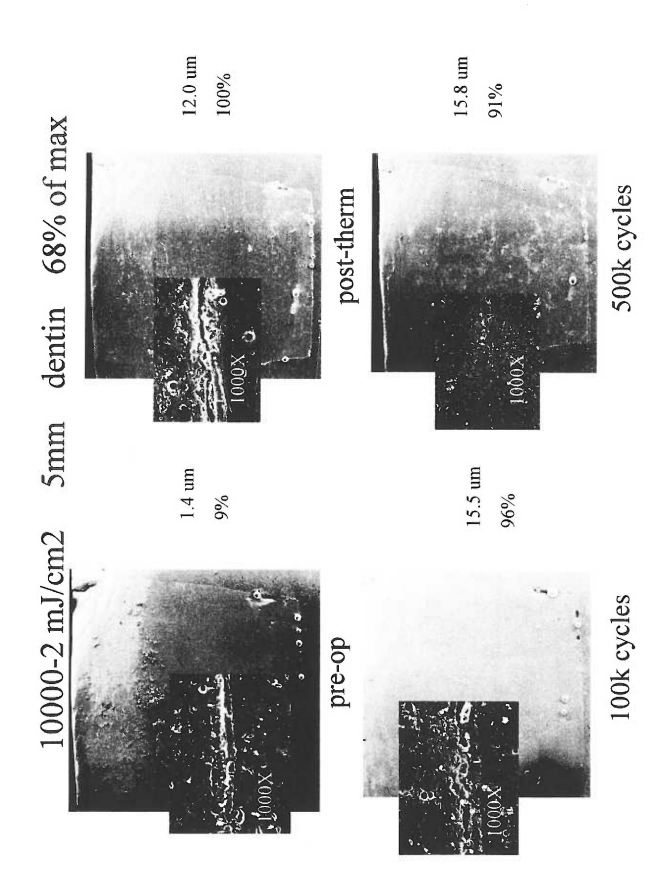


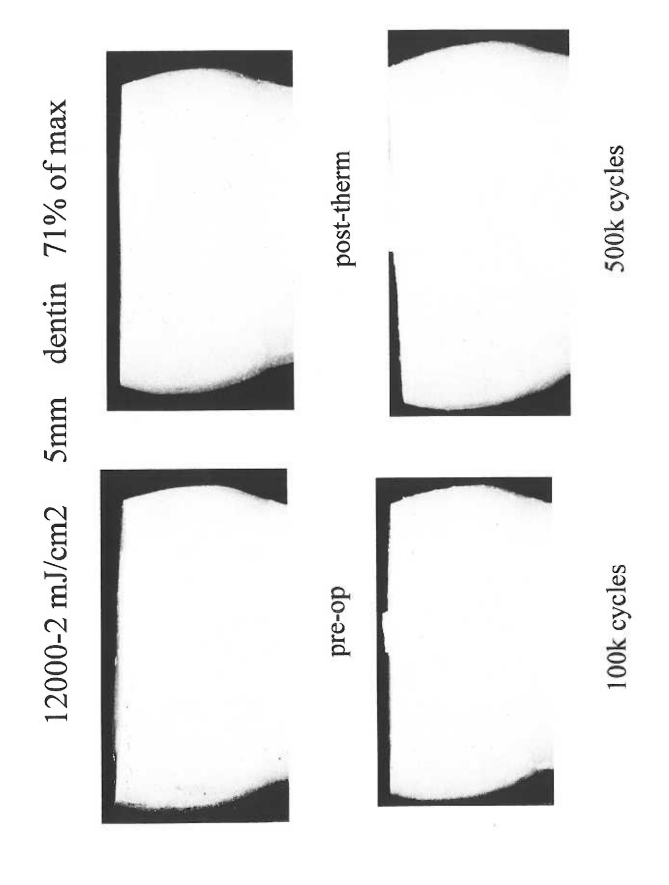


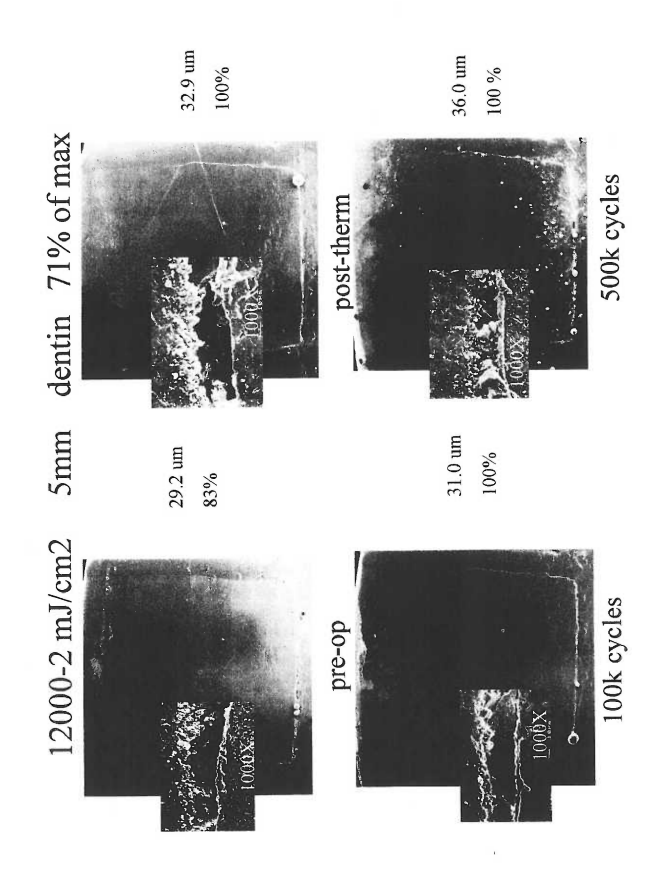


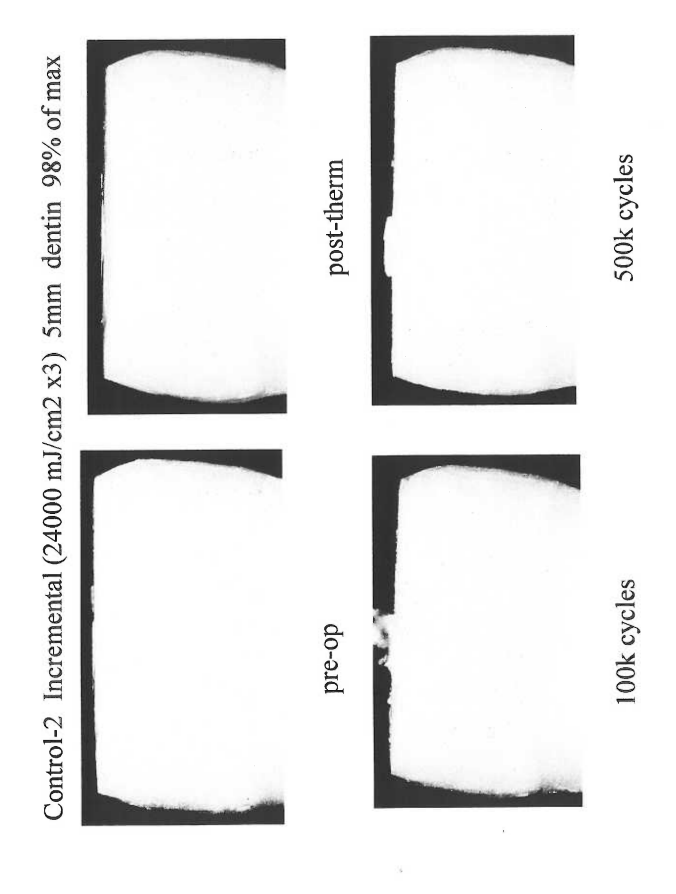












3.1 um 5.1um %68 %09 Control-2 Incremental (24000 mJ/cm2 x3) 5mm dentin 98% of max 500k cycles post-therm 2.6 um 4.8 um 10% 81% pre-op

100k cycles

		1 wk	1213.0	1131.2	1111.3	1075.6	1044.2	1023.0	998.0	933.0	900.0	847.0	800.0	750.0	723.2	650.2	638.2	586.5	530.6	480.2	452.3	444.6	436.9	410.5	410.2	410.9	400.0	736.0	273.9	100
		72 hrs	1150.3	1143.2	1110.0	1095.3	1035.2	1023.3	989.3	953.2	906.5	863.2	786.3	760.0	723.6	665.3	624.3	593.5	526.5	490.2	438.5	435.5	430.5	420.3	415.2	405.3	375.2	734.4	273.5	9
	4W-4	48 hrs	1160.0	1130.0	1100.0	1083.0	1043.0	1022.0	1000.0	947.0	899.0	853.0	801.3	760.0	715.9	0.699	623.0	583.2	536.1	491.7	440.0	436.0	429.5	422.3	416.8	410.9	400.0	734.9	270.4	8
		Pre-op	1275.0	1129.5	1112.0	1079.3	1040.6	1020.3	1010.6	946.3	9.006	846.9	800.2	770.5	713.5	9.029	628.3	590.3	544.6	497.2	435.6	440.3	428.6	420.3	418.2	409.3	385.3	740.6	279.6	90
	Í	¥	ဖွ	5								0	0	0	0		0	0	0	0	0	0	0	0.0	0	0	0	5.9	22.4	8
		5 1 WK	Н	105.5	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0		0.0	Н	-			_
		72 hrs	41.0	98.2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	5.6	21.0	80
(suo	4W-3	48 hrs	40.0	100.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	5.6	21.2	εο
Third Pilot Defect Width (microns)		Pre-op	48.2	104.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	6.1	22.6	80
ct Wid		1 wk	133.0	64.0	00	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	150.0	120.0	110.0	100.0	0.0	31.1	52.7	28
t Defe		72 hrs 1	_	54.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	H	105.1	145.6	pubble 1	121.9	110.0	0.0		51.1	22
Pilo		72	12	Ć.							L								Ľ	Ľ		1	-	E P	1.7		_		4,	H
Third	4W-2	48 hrs	132.0	63.1	0.0	0.0	0.0	0.0	0.0	4.8	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	94.2	162.0	120.1	119.2	109.6	pubble	33.5	55.2	33
		Pre-op	168.0	57.6	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	eldand	120.3	110.0	100.0	pubble	28.5	52.0	5 8
		1 wk	515.0	480.0	430.0	312.0	235.0	210.0	193.0	210.0	200.0	20.1	26.4	63.1	70.4	53.4	54.1	414	53.2	49.2	54.4	40.1	39.8	19.0	95.1	90.1	68.2	6.44	147.8	90
		72 hrs	520.0	495.0	428.0	317.0	245.0	207.0	183.0	215.0	204.0	12.7	28.9	60.3	63.4	50.4	51.4	42.3	50.9	49.1	40,3	48.4	40.2	21.2	100.1	90.2	90.1	146.2	150,3	8
	4W-1	48 hrs	518.0	490.0	422.0	300.0	230.0	210.0	191.0	183.0	175.0	10.1	25.2	50,3	78.4	60.3	51.7	40.2	53.6	52.1	53.4	45.8	40.1	20.2	93.4	90.7	80.3	142.6	147.1	5
		Pre-op	480.0	516.0	427.0	323.0	237.0	193.0	191.0	234.0	206.0	25.5	28.5	76.0	53.8	47.5	57.1	43.2	47.8	53.4	57.2	48.1	40.5	20.1	115.2	97.0	72.0	147.6	147.8	100
	4000 mJ/cm2		-	2	ı m	4	ß	ω		ω	, o	. 0	-	. 21	<u>(</u>	4	5	, L	12	. 60	0 0	2 8	72	2	1 8	24	25	ava	std dev	% def

	342.3 342.9 343.1	60.3 58.3 59.0	47.0 48.7 48.0	
430.1	345.5	58.5	48.5	

		500K	235.8	235.8	225.5	198.5	215.4	237.7	281.4	234.6	289.9	285.6	345.2	333.4	375.2	398.2	415.6	455.7	456.6	486.5	489.7	499.6	475.5	500.0	512.3	498.5	489.6	366.9	113.8	100
		100K			200.1	189.6	225.6	247.6	280.0	286.5	284.6	296.4	330.2	345.4	386.5	400.0	421.5	467.2	444.2	496.8	478.8	500.0	506.3	487.5	475.6	454.6	463.5	365.4	109.5	9
	4-4	Post Therm	200.5	236.5	210.5	175.5	235.6	248.9	286.5	277.2	272.3	289.4	325.6	333.3	378.5	410.6	411.5	452.3	438.2	487.2	489.6	495.3	489.5	475.2	469.5	444.7	450.3	362.8	109.1	100
		Pre-op	195.3	200.3	212.4	196.3	215.8	235.6	270.6	265.3	278.2	298.6	315.2	345.6	389.5	395.8	412.3	423.6	445.5	475.3	478.9	489.2	475.0	468.8	456.8	435.6	412.6	351.5	105.8	100
		500K												1.35																
		100K	broke											100			10000													
crons)	4-3	Post Therm																												
ath (mic		Pre-op		0.00000		2000							1					3												
ect Wid		500K	501.0	465.2	425.3	445.3	358.0	375.2	365.8	351.2	382.5	414.7	398.5	385.2	275.2	265.3	297.9	275.3	255.3	233.5	241.3	215.2	210.2	200.2	235.3	345.9	252.6	326.8	87.9	100
ot Def		100K	507.4	456.3	413.2	458.3	386.5	396.5	356.2	345.2	330.2	272.4	292.3	278.2	288.5	255.3	250.0	256.3	263.0	247.5	244.3	203.0	213.3	213.2	223.3	275.3	285.5	308.4	84.8	100
Third Pilot Defect Width (microns	4-2	Post Therm	394.0	386.5	385.6	398.9	405.2	386.5	375.6	352.3	333.9	303.9	281.2	293.2	291.5	261.3	237.5	235.6	265.3	253.4	255.8	237.5	232.1	241.3	215.3	289.6	258.0	302.8	63.8	001
		Pre-op	prophe	395.6	387.5	402.3	375.2	372.4	364.2	343.5	325.4	300.4	275.6	289.3	288.5	254.8	245.4	232.5	256.4	247.8	254.2	244.3	238.5	245.6	256.3	275.6	250.4	296.7	57.7	100
		500K	765.3	745.2	753.6	745.2	744.3	756.2	762.3	801.2	802.3	768.3	785.0	765.0	803.2	795.3	789.3	765.3	745.8	730.9	745.2	755.6	789.3	785.3	802.3	807.3	775.3	771.4	23.1	5
		100K	763.2	728.6	748.0	732.5	736.9	745.8	758.2	790.5	783.5	745.3	786.3	775.6	801.6	802.6	795.3	756.8	735.2	732.5	735.5	754.2	745.2	6.077	798.6	806.3	789.3	764.7	26.1	8
	4-1	Post Therm	749.3	730.2	744.5	742.3	720.1	737.5	755.6	780.3	770.3	751.3	770.3	780.8	790.3	778.4	764.3	745.2	743.2	728.4	735.5	750.0	740.5	760.1	789.3	798.3	751.2	756.3	21.1	100
		Pre-op	750.0	728.3	738.0	752.3	718.9	732.4	746.1	775.3	762.9	750.0	7.797	777.4	775.3	762.3	754.3	730.2	742.8	733.2	715.8	745.8	738.2	756.2	775.3	780.2	745.2	750.2	18.5	100
	4000 mJ/cm2		-	2	m	4	S	9	7	80	o	5	F	5	5	4	5	16	17	18	61	20	27	22	23	24	72	avg.	std dev	% def

Third Pilot Defect Width (microns)

								200			
둟	Post Therm	100K	500K	Pre-op	Post Therm	100K	500K	Pre-op	Post Therm	100K	500k
121	22.3	22.4	22.2	7.4	6.6	11.0	6.1	95.0	85.0	90.0	89.0
۲	12.4	12.3	11.3	0.0	0.0	0.0	7.6	115.0	100.0	110.0	107.0
1	15.0	pubble	6.5	0.0	0.0	14.9	14.6	0.0	0.0	0.0	0.0
_	6.7	8.8	12.5	0.0	17.7	14.9	15.5	0.0	0.0	0.0	14.9
ľ	0.0	3.5	1.4	0.0	14.6	14.6	15.2	0.0	0.0	0.0	12.1
°	0.0	0.0	0.0	0.0	10.1	10.8	11.4	0.0	7.3	0.0	0.0
2	2.9	10.1	0.0	0.0	15.2	12.0	16.1	9.7	10.5	0.0	12.
۲	13.5	2.4	0.0	0.0	13.3	11.1	11.1	0.0	0.0	8.4	33.5
e	3.2	4.0	0.0	0.0	16.1	14.6	12.0	flash	flash	flash	flash
9	6.3	0.0	0.0	0.0	8.9	13.9	10.8	flash	flash	flash	flash
	4.1	1.5	0.0	0.0	0.0	9.5	12.3	flash	flash	flash	flast
6,	3.5	10.5	0.0	0.0	13.3	8.2	8.6	flash	flash	flash	flash
ľ	6.7	8.3	4.7	0.0	14.6	11.4	15.2	flash	flash	flash	flast
-	1.9	2.9	0.0	0:0	11.7	8.6	9.5	flash	flash	flash	flast
	5.9	1.6	3.5	0.0	10.8	10.7	9.5	0.0	0.0	0.0	0.0
(,,	3.5	1.0	4.3	0.0	14.9	11.7	12.0	2.2	0.0	10.4	12.0
	2.5	1.5	0.0	0.0	10.7	8.6	6.7	10.4	10.5	20.2	13.6
(2.8	2.2	0.0	0.0	10.7	10.1	12.7	0.0	pubble	pubble	20.6
-	11.7	6.0	7.7	0.0	8.2	12.0	12.3	0.0	0.0	5.4	0.9
	4.4	1.9	0.0	0.0	9.7	6.9	11.4	0.0	14.2	12.2	16.5
,	4.1	0.0	0.0	0.0	6.3	9.2	9.4	0.0	0.0	pubble	32.7
ľ	2.5	0.5	0.0	0.0	25.9	24.8	23.1	0.0	0.0	0.0	0.0
-	14.5	14.0	10.7	0.0	9.8	7.9	8.4	0.0	0.0	0.0	13.5
2	22.4	20.8	22.7	0.0	11.2	10.4	10.4	0.0	0.0	0.0	11.1
7	70.07	71.3	68.0	0.0	10.7	11.5	10.3	0.0	0.0	0.0	0.0
0,	9.6	8.4	7.1	0.3	10.8	11.3	11.7	12.1	12.6	15.1	20
	14.1	14.9	14.3	1.5	5.7	4.2	3.6	33.0	29.5	32.7	29.1
	92	75	84	4	88	96	100	56	33	41	7.

5.9	11.0	11.6	13.
5.9	1.5	3.3	7.0
15.7	71.0	7.07	74.
+.	33.0	27.8	26.

8 deve

_
icrons)
E
Width
Defect
Pilot
Third

	8000 mJ/cm2		, ide			8-2				200	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
	Pre-op	Post Therm	100X	500K	Pre-op	Post Therm	100K	200K	Pre-op	Post Therm	10K	8
	0	16.3	23.7	12.2	90.2	152.0	172.0	160.1	0	18.3	5.8	9.4
7	0.0	0.0	6.3	0.0	150.0	161.6	142.2	162.8	0.0	60.2	pubble	72.4
е,	0.0	4.1	0.9	10.2	121.0	133.0	128.0	130.0	pubble	elqqnq	72.6	82.3
4	0.0	0.0	0.0	11.1	0.0	6.7	0.0	0.0	10.3	27.4	17.9	15.9
2	0.0	3.8	30.4	41.4	0.0	12.7	5.1	6.7	7.3	6.7	0.0	0.0
9	0.0	6.9	30.4	57.9	0.0	18.0	0.0	14.9	5.4	13.0	0.0	0.0
_	0.0	5.4	10.2	6.1	0.0	12.7	0.0	9.2	4.8	10.1	0.0	0.0
ω	0.0	10.2	28.0	16.5	elqqnq	17.7	pubble	15.5	0:0	9.5	0.0	13.6
. О	80.0	76.3	64.0	51.0	0.0	13.3	0.0	13.0	5.1	8.9	0.0	0.0
0	67.7	67.0	60.1	pubble	0.0	17.1	0.0	0.0	0.0	12.7	0.0	0.0
11	82.3	91.2	90.5	93.6	0.0	13.3	10.5	15.2	9.8	16.5	0.0	9.8
12	0.0	3.2	90.6	86.1	0.0	18.0	12.7	13.3	0.0	12.0	0.0	8.2
13	0.0	8.6	8.2	14.2	0.0	9.8	16.1	13.9	12.7	13.3	0.0	11.7
14	0.0	13.0	11.7	6.8	12.7	12.7	0.0	21.2	6.3	12.7	0.0	7.9
15	0.0	15.2	7.9	12.2	0.0	23.8	17.7	13.3	11.7	12.0	0.0	0.0
91	0.0	9.6	18.0	10.8	0.0	24.1	0.0	0.0	8.2	12.7	0.0	17.2
17	0.0	15.5	24.4	12.1	0.0	27.9	əjqqnq	22.2	9.2	11.4	0.0	17.2
18	0.0	13.3	12.7	10.2	0.0	46.9	10.5	20.0	8.9	10.1	0.0	16.5
19	20.6	2.1	9.5	10.4	0.0	34.8	44.0	52.8	7.0	10.1	0.0	12.1
8	24.4	pripple	22.8	14.9	0.0	37.0	47.5	30.7	0.0	17.7	0.0	22.2
21	pripple	13.0	43.0	65.8	0.0	32.0	33.2	26.6	17.1	17.7	0.0	17.4
22	əlqqnq	17.1	67.4	55.6	0.0	12.1	32.0	0.0	6.7	12.1	0.0	7.6
23	5.7	pripple	pubble	18.7	0.0	5.1	24.7	0.0	6.1	67.1	0.0	0.0
24	3.2	pubble	4.2	4.5	0.0	16.9	8.2	26.6	propple	89.0	44.3	60.4
25	31.2	32.8	35.4	39,3	0.0	15.6	13.3	14.0	propple	elddud	pubble	pupple
avg.	13.7	19.3	29.4	27.6	15.6	34.9	31.2	31.3	6.2	20.9	6.1	16.7
std dev	26.5	25.3	26.7	27.0	41.5	44.3	48.6	46.9	4.7	21.2	17.5	22.5
% de	25	16	98	96	89	100	70	80	89	100	17	71

12.7	40.3	5.2	30.1
82.3	61.0	97.0	36.7
7.6	14.0	8.6	5.0
25.2	22.2	25.1	11.8

(microns)
Width
Defect
Pilot
Third

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3.5	TOUGO MA/CINZ	L				7-01			[201		
2012	Pre-op	Post Therm	100K	300K	Pre-op	Post Iherm	100K	SOUX	Pre-op	Post Therm	100K	500K
-	propple	67.8	36.2	6.04	30.8	22.7	22.1	18.9	142.5	166.5	164.8	162.8
7	26.6	26.4	16.7	0.0	0.0	10.5	10.8	7.3	147.0	65.0	47.0	45.0
ო	11.7	23.6	19.6	17.5	0.0	8.9	10.1	2.2	pripple	150.8	propple	pubble
4	9.5	39.2	30.1	29.5	0.0	83.0	87.4	85.6	140.5	elqqnq	142.6	126.6
5	flash	flash	flash	flash	0.0	pripple	18.1	28.2	140.0	110.2	130.9	109.2
ဖ	flash	flash	flash	flash	0.0	1.1	63.3	69.0	71.3	84.5	72.8	70.4
_	9.9	15.8	23.2	10.1	pubble	elqqnq	elqqnq	pnpple	45.2	25.7	38.1	24.5
00	0.0	10.7	10.7	9.8	0.0	8.2	10.1	37.3	0.0	24.1	21.2	20.7
ത	9.9	4.1	7.2	9.2	0.0	4.4	3.8	0.0	0.0	32.3	33.9	pubble
9	0.0	0.0	13.6	9.8	0.0	11.7	2.0	pripple	0.0	17.1	propple	36.1
=	flash	flash	3.2	flash	propple	propple	9.2	12.4	0.0	20.6	25.3	17.4
12	flash	flash	flash	flash	0.0	5.4	5.1	7.6	0.0	19.0	25.6	21.5
13	flash	flash	flash	flash	0.0	5.4	5.3	8.2	0.0	14.9	0.0	9.2
4	flash	flash	flash	flash	2.1	3.5	6.4	5.4	0.0	25.1	23.4	15.5
15	flash	flash	flash	flash	0.0	2.1	11.7	6.3	0.0	14.8	0.0	0.0
16	flash	flash	flash	7.3	0.0	5.1	6.3	5.4	0.0	0.0	0.0	0.0
17	23.1	1.1	8.2	38.6	0.0	5.1	5.0	4.3	0.0	1.8	0.9	0.0
18	57.6	28.1	52.5	47.2	0.0	5.2	5.1	1.2	0.0	2.0	0.5	0.0
6	59.5	43.9	43.2	53.5	0.0	5.0	3.5	3.1	0.0	0.0	2.0	0.0
20	8.99	51.5	54.4	62.8	0.0	4.1	2.6	1.9	0.0	6.9	7.0	0.0
73	63.9	pubble	66.4	8.99	0.0	2.2	0.0	2.1	0.0	9.6	0.0	0.0
22	63,4	pubble	pripple	70.2	0.0	4.1	2.5	1.3	0.0	0.0	4.5	0.0
23	plqqnq	63.2	pubble	84.5	0.0	pubble	2.4	1.7	0.0	15.5	11.7	16.1
24	plqqnq	pubble	elddud	pubble	0.0	4.2	28.2	0.0	0.0	51.6	51.9	33.2
55	49.4	61.7	94.6	elqqnq	0.0	50.9	50.0	53.0	0.0	10.8	11.4	10.4
avg.	31.8	31.2	32.0	34.8	1.4	12.0	15.5	15.8	28.6	36.2	35.5	31.2
std dev	26.7	23.7	26.0	27.0	6.4	19.5	21.7	23.6	54.7	46.7	48.2	44.7
% def	98	93	100	94	6	100	96	91	25	88	87	65

26.5 12.8 93.7 6.0 20.6 16.7 40.0 40.6

27.3 10.1 83.3 15.9

27.6 10.7 94.3 6.7

	12000 mJ/cm2	2 12-1		1		12-2				12-3		
	Pre-op	Post Therm	100K	500K	Pre-op	Post Therm	100K	500K	Pre-op	Post Therm	100X	20
_	52.5	56.2	58.5	56.6	62.6	66.4	60.5	65.6	38.7	81.6	33.7	23.
2	33.5	38.5	51.2	40.7	42.1	elqqnq	43.9	55.8	15.2	10.8	6.1	ŏ
	50.6	37.2	42.2	38.6	44.0	53.8	51.3	69.0	11.7	16.5	12.4	0.0
4	28.4	44.6	28.4	27.0	<u>1</u> .00	36.1	30.4	32.6	10.8	26.0	19.0	20.
· w	bubble	34.4	pubble	pubble	93.6	35.8	37.0	44.7	elqqnq	36.4	21.6	16.
9	32.5	31.9	30.1	26.8	28.6	31.3	34.8	41.5	33.6	39.9	pubble	23
~	26.8	15.7	28.7	28.2	22.1	17.1	20.6	22.8	40.2	47.5	26.9	15.
. 00	21.7	22.7	27.5	32.5	30.2	38.4	29.2	35.8	63.3	50.3	pubble	19.
0	33.8	22.1	32.2	37.9	37.7	50.6	39.6	32.6	pripple	43.7	40.5	4
. 6	19.2	36.3	27.5	27.8	40.8	27.2	30.7	35.2	pripple	48.1	50.9	23
	26.2	28.1	17.7	36.3	22.2	31.3	32.9	pubble	47.0	51.3	36.1	ŏ
: 2	6.7	23.0	22.4	10.1	30.8	30.1	27.2	25.9	44.2	37.1	29.1	Ö.
. 5	flash	pubble	flash	flash	24.6	28.8	14.8	22.8	31.4	17.4	14.5	ŏ.
4	flash	pubble	flash	flash	0.0	29.4	36.1	30.1	15.8	15.2	3.2	ö
, fc	flash	flash	flash	flash	31.3	42.1	pubble	32.0	pripple	32.0	29.4	œ.
, c	flash	flash	flash	flash	29.9	32.9	30.1	papple	20.6	32.9	17.5	13
2 1	00	2.4	9.5	4.5	24.3	25.6	27.1	28.5	40.2	25.0	əlqqnq	10
- α	96	3.1	4.5	4.7	pubble	23.7	24.2	23.2	21.5	23.7	11.1	0
<u> </u>	flash	11.1	flash	flash	16.5	22.5	22.1	30.7	20.3	14.2	9.5	ö
2 2	15.7	20.5	8.1	8.5	0.0	22.5	25.8	20.6	11.4	14.6	15.2	Ö
7	30.4	19.0	13.2	8.5	0.0	25.1	23.9	24.4	21.2	27.5	19.3	ö
2	41.1	24.0	31.7	24.6	0.0	51.9	43.7	23.9	10.8	propple	10.1	Ö
1 8	pubble	35.4	33.1	pubble	15.6	36.4	propple	44.6	propple	propple	8.4	17
24	00	32.3	28.2	17.2	61.4	22.5	15.4	49.4	27.9	15.2	pubble	gq
. بر د	0.0	5.0	0.0	11.6	33.6	8.0	12.3	elqqnq	34.9	47.9	26.2	32
ave.	23.5	25.9	26.0	24.6	29.2	32.9	31.0	36.0	28.0	32.8	21.0	=
td dev	16.6	13.9	15.1	14.6	22.1	12.9	11.6	13.8	14.5	17.1	12.4	12
% def	83	100	992	100	83	100	100	100	100	190	100	اث

3.0 3.0 88.7 9.8

26.0 5.0 98.3 2.9

30.5 4.0 0.0 0.0

(microns)
-
Width
Defect V
Pilot
Third

	500K	9.4	41.5	51.6	8.2	10.7	26.9	35.1	31.7	26.3	18.7	31.9	37.7	9.5	17.1	8.2	10.2	8.7	10.2	8.4	10.1	10.1	8.6	8.3	17.1	12.2	18.8	12.7	-
	100K	elddud	pripple	45.8	0.0	12.0	42.7	38.0	elddud	32.6	19.3	40.5	27.9	7.9	7.6	8.8	10.1	14.2	12.9	16.1	15.2	pubble	9.8	13.7	pripple	pubble	19.7	13.7	
24-3	Post Therm	5.8	45.6	45.6	9.2	9.5	31.4	37.4	38.0	32.0	20.0	40.2	26.3	10.8	7.1	9.4	22.2	11.4	8.9	8.9	5.1	10.2	15.5	15.5	52.0	propple	21.6	14.9	
	Pre-op	6.2	45.1	48.1	10.8	10.1	48.5	elddud	49.4	34.5	19.6	37.1	26.6	10.5	8.0	8.2	11.1	10.4	8.3	0.0	0.9	10.1	8.2	12.0	27.9	8.0	19.4	15.9	
	500K	32.8	45.6	1.5	0	0	5.8	0	9.2	16.2	10.8	4.7	11.4	7.9	3.2	8.6	27.8	17.1	9.8	1.2	54.7	pubble	33.6	65.1	pubble	pubble	16.7	18.7	
	100K	28.2	51.9	0.0	0.0	0.0	5.7	12.0	10.1	10.8	pubble	15.2	3.5	9.8	6.9	12.0	29.8	24.1	19.3	6.0	29.8	38.6	34.5	pubble	41.5	50.1	19.1	16.1	
24-2	Post Therm	27.8	27.2	0.0	0.0	0.0	0.0	9.5	9.5	19.9	72.1	pipple	21.8	18.1	16.5	32.3	36.1	29.1	12.9	10.8	19.6	20.2	43.0	82.3	elqqnq	eldqnq	23.1	21.3	
	Pre-op	0	17.7	0.0	0.0	16.2	0.0	7.6	10.5	16.2	10.8	9,5	8.9	9.2	9.5	26.9	23.4	24.4	8.2	14.6	34.8	pubble	12.4	24.1	12.4	propple	12.9	9.5	
	500K	105.7	0.0	41.5	0.0	16.1	0.0	0.0	0.0	11.1	5.7	0.0	4.1	0.0	0.0	0.0	0.0	0.0	39.3	0.0	40.8	12.7	32.1	31.7	pubble	36.4	15.7	24.8	
	100K	103.2	0.0	0.0	0.0	0.0	0.0	25.9	29.2	0:0	0.0	0.0	0.0	0.0	0.0	0.0	8.9	pubble	26.3	22.5	51.3	17.1	13.6	27.2	44.2	27.1	16.5	242	
24-1	Post Therm	102.2	0.0	0.0	16.5	4.8	5.4	13.0	5.7	16.5	0.0	0.0	0.0	00	0.0	0.0	5.7	14.3	29.5	411	50.9	14.6	56.0	8.6	46.8	0.0	16.1	24.3	
24000 mJ/cm2	Pre-op	1001	0.0	0.0	0.0	pubble	0.0	0.0	0.0	0.0	0.0	00	00	00	000	0.0	00	00	0.0	00	42.4	0.0	71.0	200	7.0	00	26	25.4	10.7
2		<u>_</u>	. 0	(7)	4	L vo			. «	0 0	, ç	-	5	1 4	5 4	. 10	9 4	1 2	. 82	0 0	2 8	7	3 :	1 8	3 8	, K	2 5%) of the	מכר כמי

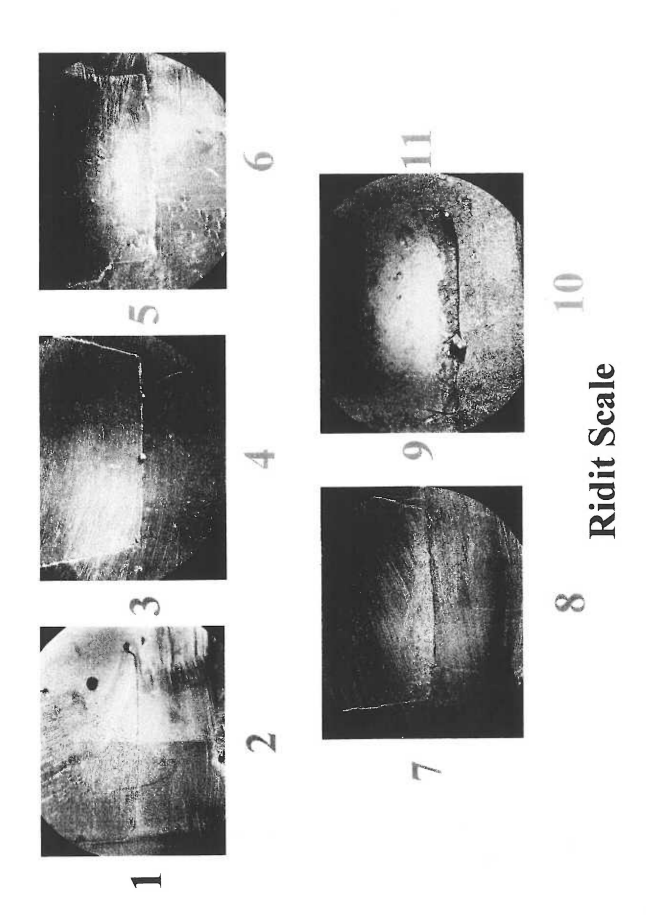
13.8	20.3	18.5	17.1
5.1	3.7	1.7	1.6
63.7	80.7	0.77	7.87
44.4	16.0	23.6	25.8

		200K	6.2	6.4	5.4	7.3	6.2	4.6	2.7	4.2	4.2	1,1	2.7	3.7	1.8	1.7	6:	1.5	1.1	1.3	3.5	4.4	6.8	3.4	5.2	4.2	8.5	1.4	2.0	9
		ğ	5.6	6.2	4.7	6.5	6.7	4.8	2.7	4.9	4.5	2.8	2.9	3.5	1.5	1.8	1.6	1.5	1.8	1.4	2.9	4.5	5.5	3.4	8.7	5.4	9.6	4.2	2.3	5
	4	Post Therm	5.2	6.0	4.7	6.0	6.1	4.9	3.5	4.9	3.5	2.8	2.6	3.0	1.6	1.8	1.9	1.8	1,3	1.6	4.9	3.8	3.7	2.6	6.4	5.2	8.4	3.9	1.9	9
		Pre-op	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0
		500K	11.3	6.7	6.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	5.7	0.0	0.0	0.0	0.0	0.0	63.3	57.0	41.2	0.0	0.0	80.7	86.4	14.3	27.4	36
		100K	4.9	5.1	9.7	0.0	11.7	8.3	0.0	0.0	0.0	1.8	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.09	54.4	55.8	34.5	80.0	85.1	16.4	28.0	84
crons)	c-3	Post Therm	14.9	9.8	11.7	10.1	2.9	3.2	5.4	7.9	8.5	11.1	3.0	4.7	3.5	7.3	4.4	8.6	4.4	7.2	44.9	44.7	42.3	42.7	70.9	66.5	96.3	21.5	25.8	100
oth (mi		Pre-op	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	14.9	42.8	46.2	propple	31.2	47.6	84.1	11.1	22.4	21
Sct W		500K	47.0	pubble	1.6	0.0	3.2	flash	flash	flash	flash	0.5	9.0	0.1	4.0	1.0	1.	4.	0.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	3.5	3.1	10.4	8
Defe	7,0	100K	47.0	2.9	0.0	3.2	4.0	flash	flash	flash	flash	11.1	5.3	0.0	5.1	2.9	0.0	2.9	5.4	2.2	2.2	0.0	1.9	9.0	0.5	1.0	2.3	4.8	2.7	26
Third Pilot Defect Width (microns)	c-2	Post Therm	47.0	pubble	2.2	2.5	4.3	flash	flash	flash	flash	2.1	piddud	2.9	2.3	1.8	2.9	4.4	9.1	2.0	1.5	2.4	0.0	1.7	10.7	0.0	4.5	5.1	10.4	88
-	The state of the s	Pre-op	47.0	4.4	0.0	0.0	0.0	flash	flash	flash	flash	0.0	0.0	0.0	pubble	0.0	0.0	0.0	0.0	0.0		0.0	0.0	0.0	0.0			2.6		10
		500K	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	9.	0.3	00	0.2	00	2.4	0.0	0.0	2.3	0.0	00	90	0.	45
		100K	flash	flash	flash	Н	⊢	flash	\vdash	⊢	⊢	+	+	╁	H	╁	0.0	00	00	000	00	0.0	00	0.0	0.0	00	00	0.0	0.0	0
	r	Post Therm	flash	flash	flash	r	t	t	T	t	flash	flash	flash	flash	flash	flash	13	0.0	00	0.0	00	0.0	80	0.3	<u>r</u>	00	00	15	3.1	98
		Pre-op	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	flash	00	000	000	000	00	000	00	00	C	00	000	00	0.0	0
	72000 mJ/cm2		,-	2	m	4	- IC	. 6		. α		. 6		. 5	i 6	2 4	. to	, c	5 5	- 00	<u>.</u>	2 6	3 8	. 8	3 1	2 6	, K	3 5	std dev	% def

3.4	8.0	6.3	5,5
5.3	9.1	7.0	0.0
7.8	81.3	57.3	60.3
0.0	30.6	43.8	28.3

			•													-
	Pre-op	Pre-op Post-thm	100 4	200K	Pre-op	Post-thm	100 ¥	Š	Pre-op	8	ğ	% %	Pre-op	Post-thm	ğ	ğ
4000-1-W mJ/cm	1	7	7	7	9	7	9	9	4	9	9	9	5.7	6.7	6.3	6.3
2	4	2	S	9	6	4	ည	ဖ	2	4	4	9	4.0	4.3	4.7	6.0
m	2	2	6	6	2	2	9	8	7	7	2	2	2.0	2.0	2.7	2.7
4	-	8	ę	F	8	80	10	10	7	00	10	=	7.3	8.0	10.0	10.7
. 6	O'u	r.	60	88	48	5.3	6.0	6.3	4.5	5.0	5.5	6.3	4.8	5.3	5.9	6.4
st dev													2.3	2.6	3.1	3.3
4000-1 mJ/cm*	80	8	6	11	80	8	6	11	6	80	6	11	8.3	8.0	9.0	11.0
2	7	6	10	1:	ω	80	6	11	7	8	10	11	7.3	8.3	9.7	11.0
m	fell	out			[e]	ont			fell	ont						
4	7	8	6	6	9	8	8	6	7	8	თ	တ	6.7	8.0	8.7	0.6
avg	7.3	8.3	9.3	10.3	7.3	8.0	8.7	10.3	7.7	8.0	9.3	10.3	7.4	8.1	9.1	10.3
st dev													9.0	0.2	0.5	1.2
6000-1 mJ/cm2	-	2	2	2	-	2	1	2	2	2	2	2	1.3	2.0	1.7	2.0
2	-	2	7	2	-	7	1.5	2	2.5	2	2	2	1.5	2.0	1.8	2.0
l m	4	4	8	4	4	S	၉	4	က	3	2.5	3.5	3.7	4.0	2.8	3.8
avo	2.0	2.7	2.3	2.7	2.0	3.0	1.8	2.7	2.5	2.3	2.2	2.5	2.2	2.7	2.1	2.6
st dev													1.3	1.6	1.0	1.1
8000-1 m.l/cm2	3	S	ဖ	5	3	9	5	9	2	4	9	4	2.7	5.0	5.7	5.0
2	2	3	3	8	2	2	2	2	2	2	2	2	2.0	2.3	2.3	2.3
ı m	-	က	2	4	-	က	2	4	4	4	2	3.5	2.0	4.0	2.0	3.8
ava	2.0	4.3	3.7	4.0	2.0	3.7	3.0	4.0	2.7	3.3	3.3	3.2	2.2	3.8	3.3	3.7
st dev													9.0	1.6	2.0	1.7
12000-1 mJ/cm2	4	4	4	4	6	2	4	5	8	3.5	4	3.5	3.3	4.2	4.0	4.2
2	r0	3	S	5	4	4	4	4	4	4	2	4	4.3	4.3	4.7	4.3
1 (*)	4	2	4	3	9	4	3	2	က	4	3	2	3.3	4.3	3.3	2.3
DAG	4.3	4.7	4.3	4.0	3.3	4.3	3.7	3.7	33	3.8	4.0	3.2	3.7	4.3	4.0	3.6
st dev													9.0	0.1	0.7	1.1
24000-1 mJ/cm2	2	3	3	8	2	3	3	4	2	3	3	4	3.0	3.0	3.0	3.7
2	2	2	3	4	4	က	8	4	2	2	4	က	2.7	2.3	3.3	3.7
6	2	9	8	2	4	4	4	4	က	4	3	ဗ	3.0	3.7	3.3	3.0
D'VE	2.0	2.7	3.0	3.0	4.3	3.3	3.3	4.0	2.3	3.0	3.3	3.3	2.9	3.0	3.2	3.4
stoley													0.2	0.7	0.2	4.0
Control-1	2		-	-	-	,	-	-	-	1	1	,	1.3	1.0	1.0	1.0
2	-	-	,-	-	2	2	2	2	2	2	2	2	1.7	1.7	1.7	1.7
ı m	5	ഗ	2	5	2	22	4.5	വ	4.5	2	5	5	4.8	5.0	4.8	5.0
4	-	-	-	-	-	1.5	1.5	1.5	-	2	2	2	1.0	1.5	1.5	ر ئ
DXE	23	2.0	2.0	2.0	23	2.4	2.3	2.4	2.1	2.7	2.5	2.5	2.2	2.3	2.3	2.3
0 00													α.	9,	4.7	0 4

PART TWO FINAL METHOD



Ridit Analysis n = 8 3 evaluators

	T-tests	pre-500k								0.0022083											0.0348542											0.4015076				
	20 20 20	7.0	5.0	3.0	11.0	4.3	6.3	7.3	0.0	6.3	2.4	0.15	10.7		9.0	-	6.4	1.3	4.3	5.3	0.9	3.8	2.3	2.7	2.0	2.7		2.3	4.3	1.3	1.7	2.8	1.3	5,0	3.7	0.0
	10k	7.0	4.3	2.7	9.3	4.0	5.3	6.3	5.7	5.6	2.0	93	9.3		8.7	23	4.3	1.3	3.3	4.7	5.4	3.2	2.3	3.0	4.7	5.0		3.0	3.3	1.7	1.7	2.7	1.0	5.7	3.0	3.7
Avg. Total	Post-thm	6.3	3.7	2.0	8.0	2.3	5.3	4.0	4.3	4.5	2.0	8.3	8.0	ont	8.0	1.3	4.0	1.0	3.7	4.3	4.8	3.0	3.3	3.0	5.0	2.0	ont	2.7	4.0	1.7	1.7	2.9	1.2	2.0	4.0	2.0
24	Pre-op	6.3	3.3	2.0	7.0	2.0	5.0	3.0	3.3	4.0	1.9	8.7	7.0	fe]	7.0	ر. دن	4.7	1.3	3.0	4.7	4.7	2.7	1.3	2.0	4.7	2.0	重	3.7	3.3	1.7	1.7	2.5	1.2	2.7	1.7	1.7
	500k	1 2	4	2	1.1	4	7	7	7	6.1		=	9		တ	-	က	-	4	9	5.6		2	က	4	2		2	ဗ		2	2.4		2	3	4
	100k	7	4	2	6	4	7	5	7	5.6		6	တ		8	2	4	1	3	4	5.0		2	4	4	2		4	3	2	2	2.9		9	3	6
	Post-thm	ဖ	3	2	7	3	9	2	3	4.0		6	8		80	-	4	1	4	4	4.9		က	3	4	2		3	4	2	2	2.9		5	က	4
	Pre-op	9	3	2	7	2	9	-	3	3.8		6	7		7	-	4	1	3	4	4.5		-	-	4	1		4	၉	2	-	2.1		2	-	-
	500k	7	2	4	=	2	2	7	5	6.1		11	11		o	2	4	2	5	ß	6.1		3	8	9	4		က	5	2	7	3.5		2	2	7
	100k	7	4	9	6	2	က	7	4	5.3		10	6		ത	3	3	2	က	ß	5.5		3	3	9	2		2	4	2	2	3.0		5	3	2
	Post-thm		3	2	6	3	4	8	2	4.4		8	7		89	2	2	-	3	4	4.4		5	4	9	2		2	4	2	2	3.4		2	9	9
	Pre-op	9	8	2	7	2	6	2	2	3.8		6	7		7	2	4	2	က	S	4.9		2	3	ß	က		4	က	2	က	3.1		က	2	7
	500k	7	g	8	=	4	7	8	9	6.5		11	1,		တ	2	9	-	4	2	6.1		6	10	2	2		2	5		-	25		2	9	4
	100k	+-	<u>ر</u>	0 80	10	3	9	7	9	5.9		6	10		6	2	ဖ	-	4	2	5.8		2	10	4	2		3	3	1		23		9	9	3
	Post-thm		· uc	2	8	-	9	7	2	5.1		8	6	ont	8	-	9	-	4	2	5.3	2.5	0	10	1 14.	2	out	6	4			25		5	3	5
	Pre-op	1	4	2	7	, ,	1 6	0	0	45		8	_	lej.	7	-	9	-	6	ر د	AB		,	- '	1 m	2	ā	8	4	-	-	2.4	1.1	3	2	2
		4000-1-W m.l/cm²		۷ ۳	> 4	٠ ٠	യ	۸ ر	- a	200	Stoley	4000-1.mJ/cm ²	2	ı e	9 4	LC.	· (C		- α	ο σ		Day to	St uev	ממסבו וווס/כוווד	W 60	0 4	· v) (C) h	- α	οσ		500	SOOD-1 m.l/cm2	2	r

					0.1134822										0,4070838										0.5164896										0,4869154	
4.3	5.7	2.3	4.0	3.7	4.2	1.0	4.0	5.0	3.7	2.7	4.7	5.0	4.0	3.7	4.1	8.0	3.7	3.7	3.0	3.7	3.0	2.0	3.0	3,3	3.2	9.0	2.3	2.0	4.0	1.7	2.3	1.3	2.3	2.0	2.3	9.0
4.3	5.7	2.7	4.3	4.3	4.2	1.1	4.0	4.7	4.0	2.0	4.3	5.0	4.0	4.3	4.0	6.0	3.0	3.3	3.3	3.7	3.3	2.3	3.7	4.0	3.3	0.5	2.3	2.3	4.3	1.7	2.3	1.3	1.7	2.0	2.3	6.0
4.0	5.0	2.7	4.3	4.7	4.3	0.8	4.7	5.3	4.7	1.0	4.7	5.0	3.7	3.7	4.1	4.1	3.0	2.3	3.7	4.3	3.3	3.0	3.3	3.7	3.3	9.0	2.3	2.3	4.0	1.3	2.0	1.7	2.0	1.7	2.2	0.8
4.0	6.0	2.7	3.7	4.0	3.3	1.4	3.7	5.0	3.7	2.0	4.0	4.7	3.7	4.7	3.9	6.0	3.0	2.7	3.0	4.3	2.7	3.0	2.3	3.0	3.0	9.0	1.7	2.3	5.3	1.7	2.7	2.0	2.0	1.7	2.4	1.2
4	2	2	4	3	3.8		4	2	4	3	5	2	4	က	4.1		4	ဧ	ဧ	4	၉	2	က	9	3.1		2	2	4	2	8	1	2	2	2.3	
4	2	2	ဧ	4	3.8		4	4	4	2	4	2	4	4	3.9		9	4	3	4	3	2	3	4	3.3		2	3	4	2	3	1	2	3	2.5	
က	4	3	ဗ	4	3.6		2	9	4	1	9	ဥ	3	2	3.6		က	2	4	လ	4	က	4	4	3.6		2	2	4	2	2	2	က	2	2.4	
8	2	2	3	4	2.6		က	5	ဧ	2	3	ဧ	8	4	3.3		2	2	က	4	3	က	2	2	2.6		1	2	2	2	က	1	2	,	2.1	
ß	9	ဧ	2	2	5.1		4	2	4	4	2	2	S	4	4.5		4	4	4	က	4	2	3	4	3.5		3	2	3	2	2	-	က	2	2.3	
2	9	4	ဖ	2	4.9		4	2	4	3	2	2	2	2	4.5		က	က	4	4	4	က	2	4	3.8		9	2	4	2	7		2	2	2.3	
5	2	3	2	9	5.1		2	2	ည	1	2	2	က	2	4.8		က	ဧ	4	4	က	က	က	4	3,4		ო	က	က	1	7	-	2	2	2.1	
2	7	4	ည	4	4.0		4	2	4	က	2	9	ഹ	ស	4.6		2	4	4	2	က	က	က	2	0.4		2	က	9	2	က	2	7	2	2.8	
4	9	2	8	8	3.8		4	2	3	-	4	S	ო	4	3.6		3	4	2	4	2	2	3	6	2.9		2	2	ည	,	2	2	2	2	2.3	
4	9	2	4	4	4.0		4	5	4	1	4	သ	က	4	3.8		3	9	3	3	3	2	3	4	3.0		2	2	5	1	2	2	1	1	2.0	
4	9	2	2	4	4.3		4	ည	ιc	-	4	က	က	4	3.9		3	2	8	4	3	က	3	3	3.0		2	2	ည	1	2	2	-	-	2.0	
4	9	2	က	4	3.3		4	3	4	-	4	က	က	က	3.9		2	2	2	4	2	က	2	2	2.4		2	2	ഹ	-	2	8	2	2	2.4	
4	വ	မ	7	œ	avg	st dev	12000-1 mJ/cm2	2	က	4	2	9	7	80	avg	st dev	24000-1 mJ/cm2	2	က	4	ιΩ	9	7	60	avg	st dev	Control-1	2	e	4	ιΩ	9	7	89	avg	st dev

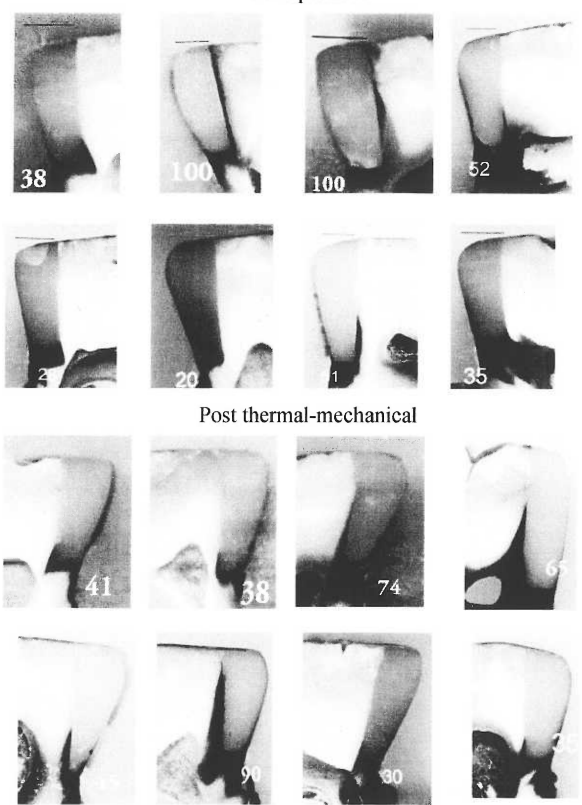
Microleakage Raw Scores and Summary

% pre	% post_
38	41
100	38
100	74
52	65
28	45
20	90
51	30
35	35
38	52
100	38
broke	broke
100	100
52	52
28	45
20	38
51	95
35	25
20	44
76	100
100	100
42	64
	broke
89	33
32	23
59	42
43	33
	100
	39
	39
	31
	61
	43
	24
	40
	43
	100
	100
	70
	28
	21
	34
	51
	38 100 100 52 28 20 51 35 38 100 broke 100 52 28 20 51 35 20 76 100 42 broke

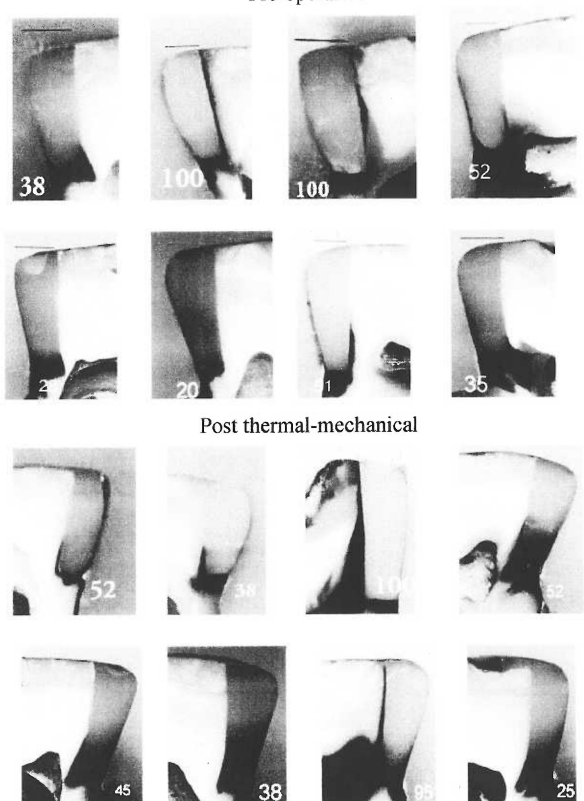
Group	Avg Pre	St dev	Avg Post	St Dev	T-test	ANOVA
4w	53	31	52	21	0.96	а
4	53	31	56	27	0.86	а
6	58	28	55	30	0.85	а
8	57	25	47	24	0.44	а
12	59	18	56	31	0.84	а
24	38	14	30	5	0.15	а
С	68	25	41	32	0.11	а

Group	% pre	% post
24-1	30	27
24-2	15	37
24-3	36	27
24-4	36	30
24-5	49	26
24-6	38	24
24-7	39	34
24-8	64	37
C-1	100	13
C-2	65	39
C-3	68	100
C-4	62	100
C-5	19	14
C-6	52	20
C-7	91	20
C-8	85	21

Microleakage 4000 mJ/cm² (water only) Pre-operative

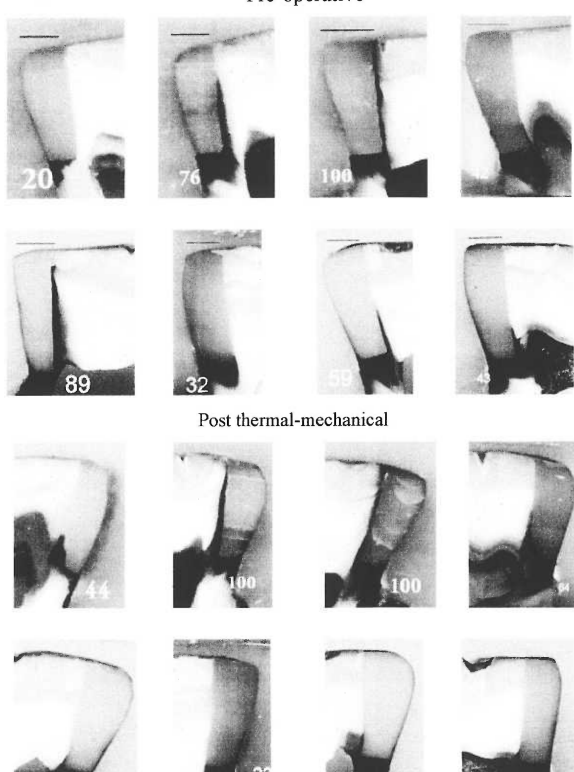


Microleakage 4000 mJ/cm² Pre-operative

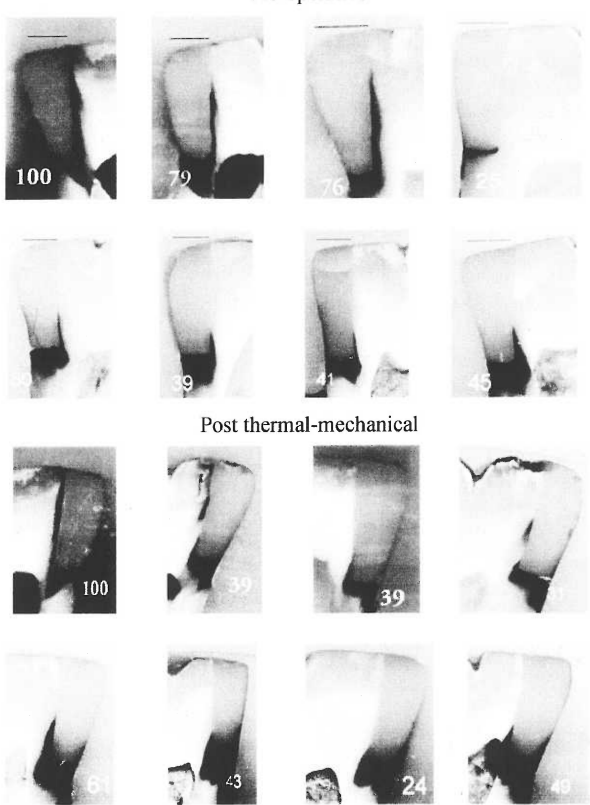


Microleakage 6000 mJ/cm²

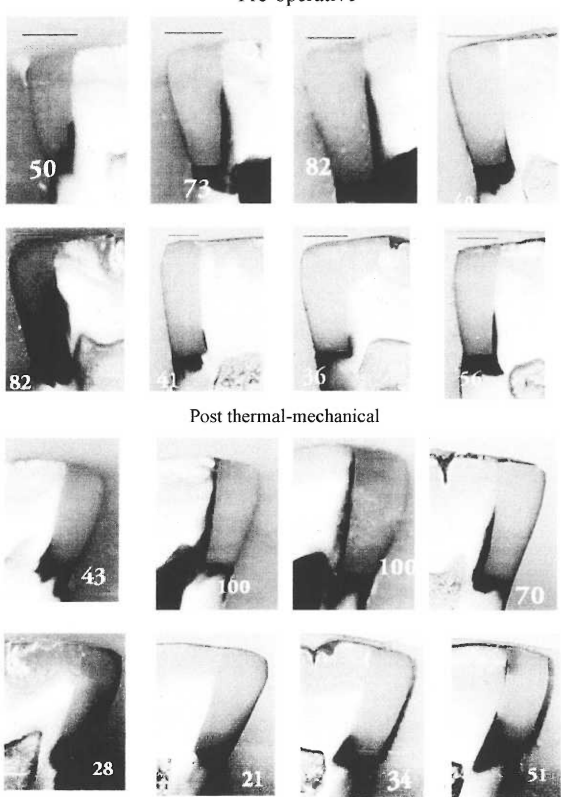
Pre-operative



Microleakage 8000 mJ/cm² Pre-operative

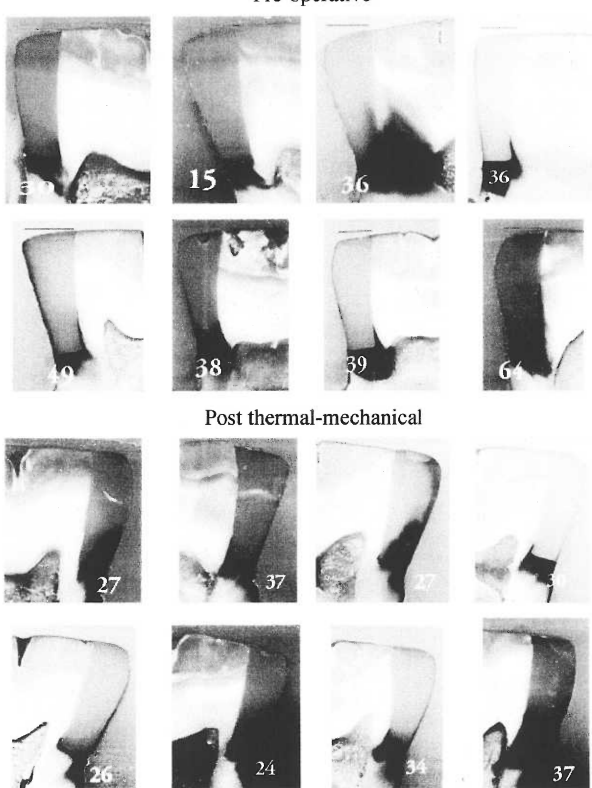


Microleakage 12000 mJ/cm² Pre-operative

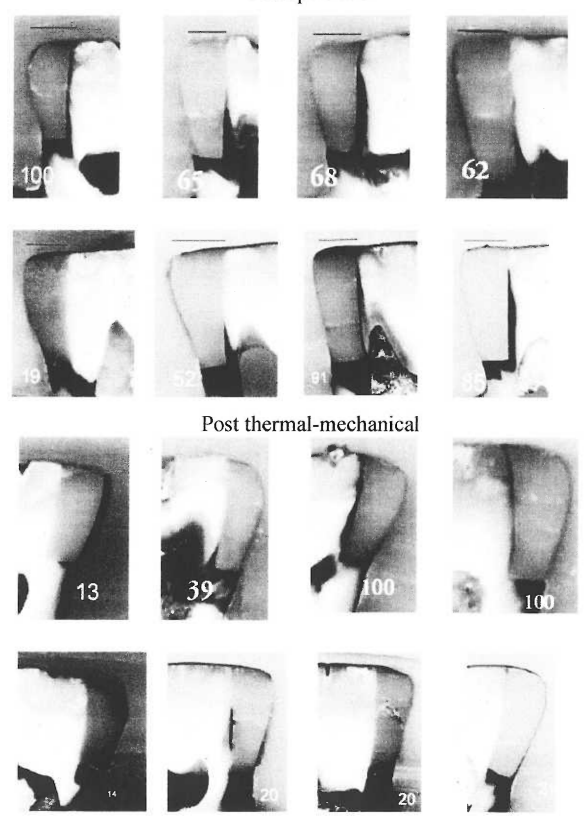


Microleakage 24000 mJ/cm²

Pre-operative



Microleakage 72000 mJ/cm² (Control) Pre-operative



Ridit 2 - WAY ANOVA

Descriptive Statistics

Dependent Variable: RIDIT

ENERGY	STRESS	Mean	Std. Deviation	N
12	100k	4.037	.899	8
	500k	4.100	.782	8
	postherm	4.100	1.376	8
	preop	3.938	.943	8
25	Total	4.044	.976	32
24	100k	3.325	.520	8
	500k	3.175	.575	8
	postherm	3.325	.597	8
	preop	3.000	.581	8
	Total	3.206	.558	32
4	100k	5.400	3.248	8
	500k	5.950	3.834	8
	postherm	4.825	2.964	8
	preop	4.713	2.741	8
	Total	5.222	3.104	32
4w	100k	5.575	2.032	8
	500k	6.238	2.397	8
	postherm	4.488	2.005	8
1	preop	3.988	1.900	8
	Total	5.072	2.182	32
6	100k	2.713	1.013	8
	500k	2.788	1.257	8
	postherm	2.925	1.166	8
	preop	2.550	1.202	8
	Total	2.744	1.114	32
8	100k	4.212	1.100	8
	500k	4.213	1.048	8
	postherm	4.338	.789	8
	preop	3.313	1.428	8
	Total	4.019	1.139	32
Con	100k	2.238	.909	8
	500k	2.238	.793	8
	postherm	2.163	.814	8
	preop	2.425	1.213	8
	Total	2.266	.905	32
Total	100k	3.929	1.950	56
	500k	4.100	2.278	56
	postherm	3.737	1.749	56
	preop	3.418	1.677	56
	Total	3.796	1.931	224

Tests of Between-Subjects Effects

Dependent Variable: RIDIT

Source	Type III Sum of Squares	df	Mean Square	F	Sig.
Corrected Model	282.080 ^a	27	10.447	3.725	.000
Intercept	3227.724	1	3227.724	1150.942	.000
ENERGY	242.210	6	40.368	14.395	.000
STRESS	14.359	3	4.786	1.707	.167
ENERGY * STRESS	25.512	18	1.417	.505	.954
Error	549.666	196	2.804		
Total	4059.470	224			
Corrected Total	831.746	223			

a. R Squared = .339 (Adjusted R Squared = .248)

Post Hoc Tests ENERGY

Multiple Comparisons

Dependent Variable: RIDIT

Tukey HSD

		Mean			95% Confide	ance Interval
(I) ENERGY	(J) ENERGY	Difference (I-J)	Std. Error	Sig.	Lower Bound	Upper Bound
12	24	.838	.419	.414	397	2.072
	4	-1.178	.419	.073	-2.412	5.622E-02
	4w	-1.028	.419	.176	-2.262	.206
	6	1.300*	.419	.031	6.565E-02	2.534
	8	2.500E-02	.419	1.000	-1.209	1.259
	Con	1.778*	.419	.000	.544	3.012
24	12	838	.419	.414	-2.072	.397
	4	-2.016*	.419	.000	-3.250	781
	4w	-1.866*	.419	.000	-3.100	631
	6	.463	.419	.927	772	1.697
	8	813	.419	.453	-2.047	.422
	Con	.941	.419	.271	294	2.175
4	12	1.178	.419	.073	-5.622E-02	2.412
	24	2.016*	.419	.000	.781	3.250
	4w	.150	.419	1.000	-1.084	1.384
	6	2.478*	.419	.000	1.244	3.712
	8	1.203	.419	.062	-3.122E-02	2.437
	Con	2.956*	.419	.000	1.722	4.191
4w	12	1.028	.419	.176	206	2.262
	24	1.866*	.419	.000	.631	3.100
	4	150	.419	1.000	-1.384	1.084
	6	2.328*	.419	.000	1.094	3.562
	8	1.053	.419	.154	181	2.287
	Con	2.806*	.419	.000	1.572	4.041
6	12	-1.300*	.419	.031	-2.534	-6.565E-02
	24	463	.419	.927	-1.697	.772
	4	-2.478*	.419	.000	-3.712	-1.244
	4w	-2.328*	.419	.000	-3.562	-1.094
	8	-1.275*	.419	.038	-2.509	-4.065E-02
	Con	.478	.419	.915	756	1.712
8	12	-2.500E-02	.419	1.000	-1.259	1.209
	24	.813	.419	.453	422	2.047
	4	-1.203	.419	.062	-2.437	3.122E-02
	4w	-1.053	.419	.154	-2.287	.181
	6	1.275*	.419	.038	4.065E-02	2.509
	Con	1.753*	.419	.001	.519	2.987
Con	12	-1.778*	.419	.000	-3.012	544
	24	941	.419	.271	-2.175	.294
	4	-2.956*	.419	.000	-4.191	-1.722
	4w	-2.806*	.419	.000	-4.041	-1.572
	6	478	.419	.915	-1.712	.756
	8	-1.753*	.419	.001	-2.987	<u>5</u> 19

Based on observed means.

^{*.} The mean difference is significant at the .05 level.

Homogeneous Subsets

RIDIT

Tukey HSD^{a,b}

			Subset	
ENERGY	N	1	2	3
Con	32	2.266		
6	32	2.744		
24	32	3.206	3.206	
8	32		4.019	4.019
12	32		4.044	4.044
4w	32			5.072
4	32			5.222
Sig.		.271	.414	.062

Means for groups in homogeneous subsets are displayed. Based on Type III Sum of Squares
The error term is Mean Square(Error) = 2.804.

- a. Uses Harmonic Mean Sample Size = 32.000.
- b. Alpha = .05.

STRESS

Multiple Comparisons

Dependent Variable: RIDIT

Tukey HSD

		Mean Difference			95% Confide	ence Interval
(I) STRESS	(J) STRESS	(l-J)	Std. Error	Sig.	Lower Bound	Upper Bound
100k	500k	-,171	.316	.949	984	.642
	postherm	.191	.316	.931	622	1.004
	preop	.511	.316	.371	302	1.324
500k	100k	.171	.316	.949	642	.984
	postherm	.363	.316	.661	451	1.176
	preop	.682	.316	.136	131	1.495
postherm	100k	191	.316	.931	-1.004	.622
	500k	363	.316	.661	-1.176	.451
	preop	.320	.316	.744	493	1.133
preop	100k	511	.316	.371	-1.324	.302
	500k	682	.316	.136	-1.495	.131
	postherm	320	.316	.744	-1.133	.493

Based on observed means.

RIDIT

Tukey HSD^{a,b}

		Subset
STRESS	N	1
preop	56	3.418
postherm	56	3.737
100k	56	3.929
500k	56	4.100
Sig.		.136

Means for groups in homogeneous subsets are displayed.
Based on Type III Sum of Squares
The error term is Mean Square(Error) = 2.804.

- a. Uses Harmonic Mean Sample Size = 56.000.
- b. Alpha = .05.

Microleakage 2-way ANOVA

Between-Subjects Factors

		N
ENERGY	12	16
	24	16
	4	16
ŀ	4w	16
	6	16
	8	16
	con	16
STRESS	post	56
	pre	56

Descriptive Statistics

Dependent Variable: PERCENT

ENERGY	STRESS	Mean	Std. Deviation	N
12	post	55.88	31.06	8
	pre	58.50	18.19	8
NEW YORK	Total	57.19	24.63	16
24	post	30.25	5.12	8
	pre	38.38	14.13	8
	Total	34.31	11.09	16
4	post	55.63	27.30	8
	pre	53.00	30.91	8
	Total	54.31	28.20	16
4w	post	52.25	21.50	8
	pre	53.00	30.91	8
	Total	52.63	25.72	16
6	post	54.88	30.27	8
	pre	57.63	28.41	8
	Total	56.25	28.39	16
8	post	47.13	23.85	8
	pre	56.88	25.33	8
	Total	52.00	24.30	16
con	post	40.88	37.34	8
	pre	67.75	25.48	8
	Total	54.31	33.86	16
Total	post	48.13	26.96	56
	pre	55.02	25.42	56
	Total	51.57	26.31	112

Tests of Between-Subjects Effects

Dependent Variable: PERCENT

Source	Type III Sum of Squares	df	Mean Square	F	Sig.
Corrected Model	9502.929 ^a	13	730.995	1.064	.399
Intercept	297876.571	1	297876.571	433.639	.000
ENERGY	5881.929	6	980.321	1.427	.212
STRESS	1330.321	1	1330.321	1.937	.167
ENERGY * STRESS	2290.679	6	381.780	.556	.764
Error	67318.500	98	686.923		
Total	374698.000	112			
Corrected Total	76821.429	111			

a. R Squared = .124 (Adjusted R Squared = .007)

Post Hoc Tests ENERGY

Multiple Comparisons

Dependent Variable: PERCENT

Tukey HSD

		Mean Difference			95% Confide	ence Interval
(I) ENERGY	(J) ENERGY	(l-J)	Std. Error	Sig.	Lower Bound	Upper Bound
12	24	22.88	9.27	.182	-5.02	50.7
	4	2.88	9.27	1.000	-25.02	30.7
	4w	4.56	9.27	.999	-23.33	32.4
	6	.94	9.27	1.000	-26.96	28.8
	8	5.19	9.27	.998	-22.71	33.0
	con	2.88	9.27	1.000	-25.02	30.7
24	12	-22.88	9.27	.182	-50.77	5.03
	4	-20.00	9.27	.328	-47.90	7.9
	4w	-18.31	9.27	.436	-46.21	9.5
	6	-21.94	9.27	.224	-49.83	5.90
	8	-17.69	9.27	.480	-45.58	10.2
	con	-20.00	9.27	.328	-47.90	7.90
4	12	-2.88	9.27	1.000	-30.77	25.02
	24	20.00	9.27	.328	-7.90	47.90
	4w	1.69	9.27	1.000	-26.21	29.5
	6	-1.94	9.27	1.000	-29.83	25.96
	8	2.31	9.27	1.000	-25.58	30.2
	con	.00	9.27	1.000	-27.90	27.90
4w	12	-4.56	9.27	.999	-32,46	23.3
	24	18.31	9.27	.436	-9.58	46.2
	4	-1.69	9.27	1.000	-29.58	26.2°
	6	-3.63	9.27	1.000	-31.52	24.2
	8	.63	9.27	1.000	-27.27	28.52
	con	-1.69	9.27	1.000	-29.58	26.2
6	12	94	9.27	1.000	-28.83	26.96
	24	21.94	9.27	.224	-5.96	49.83
	4	1.94	9.27	1.000	-25.96	29.83
	4w	3.63	9.27	1.000	-24.27	31.52
	8	4.25	9.27	.999	-23.65	32.15
	con	1.94	9.27	1.000	-25.96	29.83
3	12	-5.19	9.27	.998	-33.08	22.71
	24	17.69	9.27	.480	-10.21	45.58
	4	-2.31	9.27	1.000	-30.21	25.58
	4w	63	9.27	1.000	-28.52	27.27
	6	-4.25	9.27	.999	-32.15	23.65
	con	-2.31	9.27	1.000	-30.21	25.58
con	12	-2.88	9.27	1.000	-30.77	25.02
	24	20.00	9.27	.328	-7.90	47.90
	4	.00	9.27	1.000	-27.90	27.90
	4w	1.69	9.27	1.000	-26.21	29.58
	6	-1.94	9.27	1.000	-29.83	25.96
	8	2.31	9.27	1.000	-25.58	30.2

Based on observed means.

Homogeneous Subsets

PERCENT

Tukey HSD^{a,b}

		Subset
ENERGY	N	1
24	16	34.31
8	16	52.00
4w	16	52.63
4	16	54.31
con	16	54.31
6	16	56.25
12	16	57.19
Sig.		.182

Means for groups in homogeneous subsets are displayed.
Based on Type III Sum of Squares
The error term is Mean Square(Error) = 686.923.

- a. Uses Harmonic Mean Sample Size = 16.000.
- b. Alpha = .05.

1-Way ANOVA Microleakage Pre and Post Stressing

Between-Subjects Factors

		N
ENERGY	12	8
	24	8
	4 4w	8
	4w	8
	6	8
	8	8
	con	8

Descriptive Statistics

Dependent Variable: PERCENPR

ENERGY	Mean	Std. Deviation	N
12	58.50	18.19	8
24	38.38	14.13	8
4	53.00	30.91	8
4w	53.00	30.91	8
6	57.63	28.41	8
8	56.88	25.33	8
con	67.75	25.48	8
Total	55.02	25.42	56

Tests of Between-Subjects Effects

Dependent Variable: PERCENPR

Source	Type III Sum of Squares	df	Mean Square	F	Sig.
Corrected Model	3756.857 ^a	6	626.143	.966	.458
Intercept	169510.018	1	169510.018	261.440	.000
ENERGY	3756.857	6	626.143	.966	.458
Error	31770.125	49	648.370		
Total	205037.000	56			
Corrected Total	35526.982	55			

a. R Squared = .106 (Adjusted R Squared = -.004)

Post Hoc Tests ENERGY

Multiple Comparisons

Dependent Variable: PERCENPR Tukey HSD

		Mean			95% Confide	ance Intonvol
(I) ENERGY	(J) ENERGY	Difference (I-J)	Std. Error	Sig.	Lower Bound	Upper Bound
12	24	20.13	12.73	.695	-19.01	59.20
	4	5.50	12.73	.999	-33.64	44.6
	4w	5.50	12.73	.999	-33.64	44.6
	6	.88	12.73	1.000	-38.26	40.0
	8	1.63	12.73	1.000	-37.51	40.7
	con	-9.25	12.73	.990	-48.39	29.8
24	12	-20.13	12.73	.695	-59.26	19.0
	4	-14.63	12.73	.909	-53.76	24.5
	4w	-14.63	12.73	.909	-53.76	24.5
	6	-19.25	12.73	.736	-58.39	19.89
	8	-18.50	12.73	.770	-57.64	20.6
	con	-29.38	12.73	.261	-68.51	9.70
4	12	-5.50	12.73	.999	-44.64	33.6
	24	14.63	12.73	.909	-24.51	53.7
	4w	.00	12.73	1.000	-39.14	39.14
	6	-4.62	12.73	1.000	-43.76	34.5
	8	-3.87	12.73	1.000	-43.01	35.2
	con	-14.75	12.73	.906	-53.89	24.39
4w	12	-5.50	12.73	.999	-44.64	33.64
	24	14.63	12.73	.909	-24.51	53.76
	4	.00	12.73	1.000	-39.14	39.14
	6	-4.62	12.73	1.000	-43.76	34.5
	8	-3.87	12.73	1.000	-43.01	35.26
	con	-14.75	12.73	.906	-53.89	24.39
6	12	88	12.73	1.000	-40.01	38.26
	24	19.25	12.73	.736	-19.89	58.39
	4	4.62	12.73	1.000	-34.51	43.76
	4w	4.62	12.73	1.000	-34.51	43.76
	8	.75	12.73	1.000	-38.39	39.89
	con	-10.13	12.73	.984	-49.26	29.01
8	12	-1.63	12.73	1.000	-40.76	37.51
	24	18.50	12.73	.770	-20.64	57.64
	4	3.87	12.73	1.000	-35.26	43.01
	4w	3.87	12.73	1.000	-35.26	43.01
	6	75	12.73	1.000	-39.89	38.39
	con	-10.88	12.73	.978	-50.01	28.26
con	12	9.25	12.73	.990	-29.89	48.39
	24	29.38	12.73	.261	-9.76	68.51
	4	14.75	12.73	.906	-24.39	53.89
	4w	14.75	12.73	.906	-24.39	53.89
	6	10.13	12.73	.984	-29.01	49.26
	8	10.88	12.73	.978	-28.26	50.01

Based on observed means.

Homogeneous Subsets

PERCENPR

Tukey HSD^{a,b}

		Subset
ENERGY	N	1
24	8	38.38
4	8	53.00
4w	8	53.00
8	8	56.88
6	8	57.63
12	8	58.50
con	8	67.75
Sig.		.261

Means for groups in homogeneous subsets are displayed.
Based on Type III Sum of Squares
The error term is Mean Square(Error) = 648.370.

- a. Uses Harmonic Mean Sample Size = 8.000.
- b. Alpha = .05.

Univariate Analysis of Variance

Between-Subjects Factors

		N
ENERGY	12	8
	24	8
	4	8
	4w	8
	6	8
	8	8
	con	8

Descriptive Statistics

Dependent Variable: PERCENPO

ENERGY	Mean	Std. Deviation	N
12	55.88	31.06	8
24	30.25	5.12	8
4	55.63	27.30	8
4w	52.25	21.50	8
6	54.88	30.27	8
8	47.13	23.85	8
con	40.88	37.34	8
Total	48.13	26.96	56

Tests of Between-Subjects Effects

Dependent Variable: PERCENPO

Source	Type III Sum of Squares	df	Mean Square	F	Sig.
Corrected Model	4415.750 ^a	6	735.958	1.014	.427
Intercept	129696.875	1	129696.875	178.775	.000
ENERGY	4415.750	6	735.958	1.014	.427
Error	35548.375	49	725.477		
Total	169661.000	56	- 6		
Corrected Total	39964.125	55			

a. R Squared = .110 (Adjusted R Squared = .002)

Post Hoc Tests
ENERGY

Multiple Comparisons

Dependent Variable: PERCENPO

Tukey HSD

		Mean Difference	10.0		95% Confide	ence Interval
(I) ENERGY	(J) ENERGY	(I-J)	Std. Error	Sig.	Lower Bound	Upper Bound
12	24	25.63	13.47	.488	-15.77	67.02
	4	.25	13.47	1.000	-41.15	41.65
	4w	3.63	13.47	1.000	-37.77	45.02
	6	1.00	13.47	1.000	-40.40	42.40
	8	8.75	13.47	.995	-32.65	50.1
	con	15.00	13.47	.921	-26.40	56.40
24	12	-25.63	13.47	.488	-67.02	15.7
	4	-25.38	13.47	.500	-66.77	16.0
	4w	-22.00	13.47	.662	-63.40	19.40
	6	-24.63	13.47	.536	-66.02	16.77
	8	-16.88	13.47	.869	-58.27	24.52
	con	-10.63	13.47	.985	-52.02	30.77
4	12	25	13.47	1.000	-41.65	41.15
	24	25.38	13.47	.500	-16.02	66.77
	4w	3.38	13.47	1.000	-38.02	44.77
	6	.75	13.47	1.000	-40.65	42.1
	8	8.50	13.47	.995	-32.90	49.90
	con	14.75	13.47	.927	-26.65	56.15
4w	12	-3.63	13.47	1.000	-45.02	37.77
	24	22.00	13.47	.662	-19.40	63.40
	4	-3.38	13.47	1.000	-44.77	38.02
	6	-2.63	13.47	1.000	-44.02	38.77
	8	5.13	13.47	1.000	-36.27	46.52
	con	11.38	13.47	.979	-30.02	52.77
6	12	-1.00	13.47	1.000	-42.40	40.40
	24	24.63	13.47	.536	-16.77	66.02
	4	75	13.47	1.000	-42.15	40.65
	4w	2.63	13.47	1.000	-38.77	44.02
	8	7.75	13.47	.997	-33.65	49.15
	con	14.00	13.47	.942	-27.40	55.40
8	12	-8.75	13.47	.995	-50.15	32.65
	24	16.88	13.47	.869	-24.52	58.27
	4	-8.50	13.47	.995	-49.90	32.90
	4w	-5.13	13.47	1.000	-46.52	36.27
	6	-7.75	13.47	.997	-49.15	33.65
	con	6.25	13.47	.999	-35.15	47.65
con	12	-15.00	13.47	.921	-56.40	26.40
	24	10.63	13.47	.985	-30.77	52.02
	4	-14.75	13.47	.927	-56.15	26.65
	4w	-11.38	13,47	.979	-52.77	30.02
	6	-14.00	13.47	.942	-55.40	27.40
	8	-6.25	13.47	.999	-47.65	35.15

Based on observed means.

Homogeneous Subsets

PERCENPO

Tukey HSD^{a,b}

	Į.	Subset
ENERGY	N	1
24	8	30.25
con	8	40.88
8	8	47.13
4w	8	52.25
6	8	54.88
4	8	55.63
12	8	55.88
Sig.		.488

Means for groups in homogeneous subsets are displayed.
Based on Type III Sum of Squares
The error term is Mean Square(Error) = 725.477.

- a. Uses Harmonic Mean Sample Size = 8.000.
- b. Alpha = .05.

PART THREE

KHN		n = 3	FI	exure B	ars		
	4 9	ecs		avg	st dev		
surface	20.7	27.0	19.9	22.5	3.9	31.4	% of max
inside	16.6	14.8	16.0	15.8	0.9	22	% of max
	8 8	ecs		avg	st dev		
surface	37.8	37.1	32.8	35.9	2.7	50	% of max
inside	39.5	33.4	36.4	36.4	3.1	50.7	% of max
	16:	secs		avg	st dev		
surface	50.3	51.6	49.0	50.3	1.3	70.1	% of max
inside	50.7	51.0	47.2	49.6	2.1	69.1	% of max
	80 :	secs		avg	st dev		
surface	66.1	64.4	65.7	65.4	0.9	91.1	% of max
inside	62.4	71.8	62.3	65.5	5.5	91.2	% of max

Flexural Strength n = 5

	Force	(lbs.)				Flexural Stre	ength (MPa)		
	Control	16 sec	8 sec	4 sec	1	Control	16 sec	8 sec	4 sec
1	6.96	4.32	3.98	1.84	1 [116.13	72.08	66.41	30.70
2	7.29	5.22	4.46	2.33	1 [121.63	87.10	74.42	38.88
3	7.78	6.37	3.84	2.54	1 [129.81	106.28	64.07	42.38
4	7.38	5.60	4.52	1.75	1 [123.14	93.44	75.42	29.20
5	8.10	5.51	4.60	2.09	1 [135.15	91.93	76.75	34.87
avg	7.50	5.40	4.28	2.11	1 [125.17	90.17	71.41	35.21
st dev	0.44	0.74	0.34	0.33	1 [7.41	12.35	5.76	5.51
5. 50					% of max	92.62	66.72	52.84	26.05

Flexural Modulus n = 5

	Force	e(lbs.)				Flexural Mo	dulus (GPa)		
	Control	16 sec	8 sec	4 sec		Control	16 sec	8 sec	4sec
1	5.50	3.60	2.50	1.25		12.04	7.88	5.47	2.74
2	5.50	5.00	3.00	1.00	9	12.04	10.95	6.57	2.19
3	5.75	4.25	2.75	1.50		12.59	9.31	6.02	3.28
4	5.50	4.25	3.00	1.00		12.04	9.31	6.57	2.19
5	5.75	3.75	2.40	1.00		12.59	8.21	5.26	2.19
avg	5.60	4.17	2.73	1.15		12.26	9.13	5.98	2.52
t dev	0.14	0.55	0.28	0.22	Ÿ	0.30	1.20	0.61	0.49
					%max	97.4	72.5	47.5	20.0

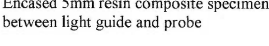
Curing Light Transmission Study

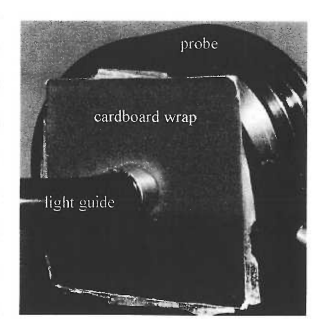
Purpose: To determine what percentage of the power density delivered to the occlusal surface actually reaches the gingival margin.

Materials and Methods: Three 5 millimeter Z250 resin composite specimens were created in a flattened template (Figure 7). The composite was incrementally placed and represented the control group as described Encased 5mm resin composite specimen All three specimens were

previously. mounted simultaneously in a square acrylic ring (Figure 8) with epoxy resin as before. The acrylic ring was encased in a black opaque cardboard wrap with a window cut out to allow light transmission only through the occlusal and gingival portion of a specimen. The encased acrylic ring was placed between the center

of the tip of the standard VIP light guide





and the center of the external orifice of the probe of the Power Max industrial radiometer. See figure above. The VIP curing light was calibrated to deliver 600 mW/cm². The specimen was exposed to the light and three readings were taken and an average calculated. The black cardboard wrap was moved to reveal the next resin composite specimen and three more readings were taken until all three specimens were tested. Next the acrylic ring with mounted specimens was replaced with only the outer sheet of the black cardboard wrap containing the entrance orifice to determine the total power exposure to the occlusal surface. Three power readings were recorded with the Power

Max and an average calculated.

In order to create the most reproducible data using the Power Max the following technique was developed.

- 1) specimen in black wrap was placed between probe and light tip
- 2) Power Max was auto-zeroed
- 3) specimen was removed
- 4) VIP curing light was activated
- 5) specimen replaced
- 6) lowest, stabilized reading recorded

The same procedure was repeated with only the outer sheet of the black cardboard wrap containing the entrance orifice representing the occlusal surface.

The area of the occlusal and gingival surfaces of the black cardboard window was determined corresponding to the respective surfaces of the resin composite specimens.

Results:

Surface areas: occlusal surface of resin composite (entrance orifice): 0.0925 cm² gingival surface of resin composite (exit orifice): 0.064 cm²

Power at gingival surface (n = 3)

Power (mW)			
1.00	0.97	1.10	
1.10	0.98	1.16	
0.98	1.05	0.91	
	1.10	1.00 0.97 1.10 0.98	

Average Power: 1.03 mW standard deviation: 0.08

Power density at gingival surface: $1.03 \text{ mW} / 0.064 \text{ cm}^2 = 16.1 \text{ mW/cm}^2$

Power at occlusal surface: (n = 1) Average: 92.4 mW

Power density at occlusal surface: $92.4 \text{ mW} / 0.0925 \text{ cm}^2 = 998.9 \text{ mW/cm}^2$

Percent change: $16.1 \text{ mW/cm}^2 / 998.9 \text{ mW/cm}^2 = 1.6\%$

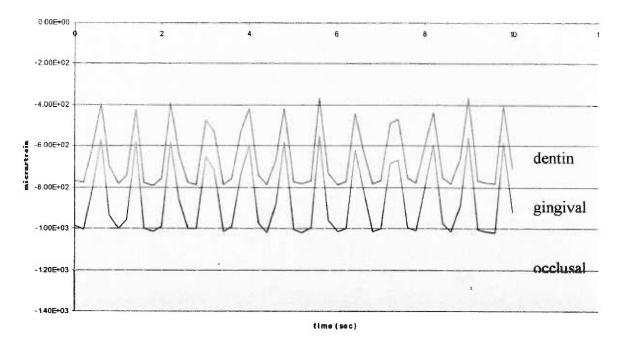
Strain Gauge Study

Purpose: To determine the magnitude of strains developed at various levels in the resin composite restoration during cyclic loading and to estimate the resultant stresses.

Materials and Methods: Three single strain gauges (CEA-06-032UW-120, Measurements Group, Raleigh, NC) were bonded on the resin composite (Control Group) near the occlusal marginal ridge, at the gingival margin, and the dentinal gingival margin (Figure 24) using cyanoacrylate. The tooth was subjected to the same cyclic forces as before (18 to 85 Newtons at 1.25 Hertz). The generated strain was conditioned by a strain gauge conditioner (2100 System, Measurements Group, Raleigh, NC).

Results: Strain was recorded at 5 per second and the results are shown graphically over a 10 second span below.

Cyclic microstrain recorded over 10 seconds simultaneously at the dentinal gingival margin, composite gingival margin and near occlusal marginal ridge of 5 mm resin composite restoration of control group.



The average of 10 maximum and minimum microstrain values was determined from the graph and the results are shown below.

Maximum and minimum microstrain over 10 seconds

occi	usal	gingi	val	den	tin
max	min	max	min	max	min
1250	754	1000	570	778	397
1260	753	1000	578	781	418
1260	763	1010	579	790	392
1260	830	1000	651	784	473
1240	788	1020	596	786	415
1270	753	1020	581	787	414
1260	742	1020	551	784	369
1270	799	1010	619	785	442
1240	866	1010	667	783	470
1250	776	1010	595	775	434
1256	782	1010	599	783	422
15.5	9.6	12.4	7.4	11.5	6.2

avg Stress (MPa)

Assuming the control group was uniformly cured and using the flexural modulus of 12.3 GPa for Z250 as determined previously (Table 12) in Part 3, the resulting stress could be estimated by the product of the microstrain and the flexural modulus as shown above.

Oneway ANOVA Time vs flexural strength or modulus

Descriptives

FLEXURE

					95% Confiden Me	
	N	Mean	Std. Deviation	Std. Error	Lower Bound	Upper Bound
4	5	35.220	5.520	2.468	28.366	42.074
8	5	71.420	5.754	2.573	64.275	78.565
16	5	90.160	12.345	5.521	74.832	105.488
80	5	125.140	7.405	3.312	115.946	134.334
Total	20	80.485	34.164	7.639	64.496	96.474

Descriptives

FLEXURE

	Minimum	Maximum
4	29.2	42.4
8	64.1	76.8
16	72.1	106.3
80	116.1	135.1
Total	29.2	135.1

ANOVA

FLEXURE

	Sum of Squares	df	Mean Square	F	Sig.
Between Groups	21093.846	3	7031.282	103.859	.000
Within Groups	1083.200	16	67.700		
Total	22177.046	19			

Post Hoc Tests

Multiple Comparisons

Dependent Variable: FLEXURE

Tukey HSD

		Mean Difference			95% Confide	ence Interval
(I) TIME2	(J) TIME2	(I-J)	Std. Error	Sig.	Lower Bound	Upper Bound
4	8	-36.200*	5.204	.000	-51.089	-21.311
	16	-54.940*	5.204	.000	-69.829	-40.051
	80	-89.920*	5.204	.000	-104.809	-75.031
8	4	36.200*	5.204	.000	21.311	51.089
	16	-18.740*	5.204	.012	-33.629	-3.851
	80	-53.720*	5.204	.000	-68.609	-38.831
16	4	54.940*	5.204	.000	40.051	69.829
	8	18.740*	5.204	.012	3.851	33.629
	80	-34.980*	5.204	.000	-49.869	-20.091
80	4	89.920*	5.204	.000	75.031	104.809
	8	53.720*	5.204	.000	38.831	68.609
	16	34.980*	5.204	.000	20.091	49.869

^{*.} The mean difference is significant at the .05 level.

Homogeneous Subsets

FLEXURE

Tukey HSD^a

			Subset for a	lpha = .05	
TIME2	N	1	2	3	4
4	5	35.220			
8	5		71.420		
16	5			90.160	
80	5				125.140
Sig.		1.000	1.000	1.000	1.000

Means for groups in homogeneous subsets are displayed.

Oneway

a. Uses Harmonic Mean Sample Size = 5.000.

Descriptives

ELASTIC

					95% Confiden Me	
	N	Mean	Std. Deviation	Std. Error	Lower Bound	Upper Bound
4	5	2.520	.487	.218	1.916	3.124
8	5	6.000	.604	.270	5.250	6.750
16	5	9.120	1.180	.528	7.655	10.585
80	5	12.240	.329	.147	11.832	12.648
Total	20	7.470	3.763	.842	5.709	9.231

Descriptives

ELASTIC

	Minimum	Maximum
4	2.2	3.3
8	5.3	6.6
16	7.9	10.9
80	12.0	12.6
Total	2.2	12.6

ANOVA

ELASTIC

	Sum of Squares	df	Mean Square	F	Sig.
Between Groups	260.694	3	86.898	165.363	.000
Within Groups	8.408	16	.526		
Total	269.102	19			

Post Hoc Tests

Multiple Comparisons

Dependent Variable: ELASTIC

Tukey HSD

		Mean Difference			95% Confidence Interval	
(I) TIME2	(J) TIME2	(I-J)	Std. Error	Sig.	Lower Bound	Upper Bound
4	8	-3.480*	.458	.000	-4.792	-2.168
	16	-6.600*	.458	.000	-7.912	-5.288
	80	-9.720*	.458	.000	-11.032	-8.408
8	4	3.480*	.458	.000	2.168	4.792
	16	-3.120*	.458	.000	-4.432	-1.808
	80	-6.240*	.458	.000	-7.552	-4.928
16	4	6.600*	.458	.000	5.288	7.912
	8	3.120*	.458	.000	1.808	4.432
	80	-3.120*	.458	.000	-4.432	-1.808
80	4	9.720*	.458	.000	8.408	11.032
	8	6.240*	.458	.000	4.928	7.552
	16	3.120*	.458	.000	1.808	4.432

^{*.} The mean difference is significant at the .05 level.

Homogeneous Subsets

ELASTIC

Tukey HSD^a

		Subset for alpha = .05				
TIME2	N	1	2	3	4	
4	5	2.520				
8	5		6.000		11	
16	5			9.120		
80	5				12.240	
80 Sig.		1.000	1.000	1.000	1.000	

Means for groups in homogeneous subsets are displayed.

a. Uses Harmonic Mean Sample Size = 5.000.

Oneway ANOVA Time vs KHN surface or KHN center

Descriptives

KHNSURF

					95% Confidence Interval for Mean	
	N	Mean	Std. Deviation	Std. Error	Lower Bound	Upper Bound
4	3	22.533	3.889	2.245	12.873	32.194
8	3	35.900	2.707	1.563	29.174	42.626
16	3	50.300	1.300	.751	47.071	53.529
80	3	65.400	.889	.513	63.192	67.608
Total	12	43,533	16.840	4.861	32.834	54.233

Descriptives

KHNSURF

	Minimum	Maximum
4	19.9	27.0
8	32.8	37.8
16	49.0	51.6
80	64.4	66.1
Total	19.9	66,1

ANOVA

KHNSURF

	Sum of Squares	df	Mean Square	F	Sig.
Between Groups	3069.620	3	1023.207	164.151	.000
Within Groups	49.867	8	6.233		
Total	3119.487	11			

Post Hoc Tests

Multiple Comparisons

Dependent Variable: KHNSURF

Tukey HSD

		Mean				***
		Difference			95% Confidence Interval	
(I) TIME	(J) TIME	(I-J)	Std. Error	Sig.	Lower Bound	Upper Bound
4	8	-13.367*	2.039	.001	-19.895	-6.839
	16	-27.767*	2.039	.000	-34.295	-21.239
	80	-42.867*	2.039	.000	-49.395	-36.339
8	4	13.367*	2.039	.001	6.839	19.895
	16	-14.400*	2.039	.000	-20.928	-7.872
	80	-29.500*	2.039	.000	-36.028	-22.972
16	4	27.767*	2.039	.000	21.239	34.295
	8	14.400*	2.039	.000	7.872	20.928
	80	-15.100*	2.039	.000	-21.628	-8.572
80	4	42.867*	2.039	.000	36.339	49.395
	8	29.500*	2.039	.000	22.972	36.028
	16	15.100*	2.039	.000	8.572	21.628

^{*.} The mean difference is significant at the .05 level.

Homogeneous Subsets

KHNSURF

Tukey HSD^a

	1	Subset for alpha = .05					
TIME	N	1	2	3	4		
4	3	22.533					
8	3		35.900				
16	3			50.300			
80	3				65.400		
Sig.		1.000	1.000	1.000	1.000		

Means for groups in homogeneous subsets are displayed.

Oneway

a. Uses Harmonic Mean Sample Size = 3.000.

Descriptives

KHNCENT

					95% Confidence Interval for Mean	
	N	Mean	Std. Deviation	Std. Error	Lower Bound	Upper Bound
4	3	15.800	.917	.529	13.523	18.077
8	3	36.433	3.050	1.761	28.856	44.010
16	3	49.633	2.113	1.220	44.385	54.881
80	3	65.500	5.456	3.150	51.946	79.054
Total	12	41.842	19.241	5.554	29.617	54.067

Descriptives

KHNCENT

	Minimum	Maximum
4	14.8	16.6
8	33.4	39.5
16	47.2	51.0
80	62.3	71.8
Total	14.8	71.8

ANOVA

KHNCENT

	Sum of Squares	df	Mean Square	F	Sig.
Between Groups	3983.536	3	1327.845	119.689	.000
Within Groups	88.753	8	11.094		
Total	4072.289	11			

Post Hoc Tests

Multiple Comparisons

Dependent Variable: KHNCENT

Tukey HSD

		Mean Difference			95% Confidence Interval	
(I) TIME	(J) TIME	(I-J)	Std. Error	Sig.	Lower Bound	Upper Bound
4	8	-20.633*	2.720	.000	-29.343	-11.924
	16	-33.833*	2.720	.000	-42.543	-25.124
	80	-49.700*	2.720	.000	-58.409	-40.991
8	4	20.633*	2.720	.000	11.924	29.343
	16	-13.200*	2.720	.006	-21.909	-4.491
	80	-29.067*	2.720	.000	-37.776	-20.357
16	4	33.833*	2.720	.000	25.124	42.543
	8	13.200*	2.720	.006	4.491	21.909
	80	-15.867*	2.720	.002	-24.576	-7.157
80	4	49.700*	2.720	.000	40.991	58.409
	8	29.067*	2.720	.000	20.357	37.776
	16	15.867*	2.720	.002	7.157	24.576

^{*.} The mean difference is significant at the .05 level.

Homogeneous Subsets

KHNCENT

Tukey HSD^a

		Subset for alpha = .05					
TIME	N	1	2	3	4		
4	3	15.800					
8	3		36.433				
16	3			49.633			
80	3				65.500		
Sig.		1.000	1.000	1.000	1.000		

Means for groups in homogeneous subsets are displayed.

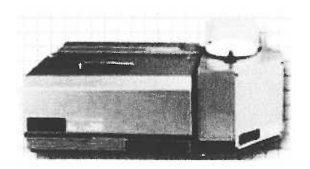
a. Uses Harmonic Mean Sample Size = 3.000.

EQUIPMENT

FOURIER TRANSFORM INFRARED SPECTROSCOPY (FTIR)

(DS20, Analect Instruments, Irvine, CA)

A Fourier transform is a complex mathematical computation that is performed by your computer. Infrared, an invisible part of the electromagnetic spectrum between visible light and radio waves refers



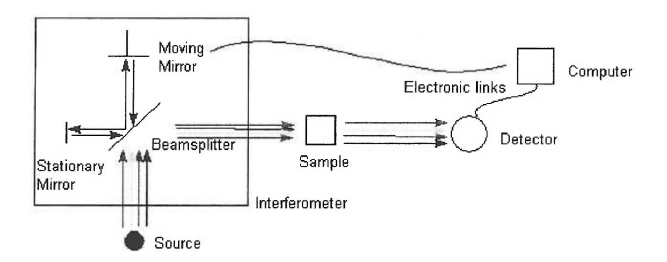
to the radiation used by the spectrometer to perform its measurement. An FTIR spectrometer is an instrument that provides information about the molecules present in a given specimen of matter as well as the quantities.

An infrared source emits light that travels through the optical system of the spectrometer (the interferometer) to the resin composite specimen on the KCl crystal. The light excites the molecules in the specimen and the molecules in turn absorb some of the wavelengths of light while the remainder are transmitted to the detector. The detector then measures the amount of transmitted light. This measurement is then converted by the analog-to-digital converter (ADC) to a form that can be used by the computer software. The software performs the Fourier transform and some other operations. A display called a spectrum then appears on your computer screen. The spectrum is actually a graphic representation of your specimen in terms of the amount of light absorbed or transmitted by your sample at various frequencies along the spectrum.

The heart of the spectrometer is the interferometer, the instrument's optical system. The interferometer is a device that encodes the light from the infrared source.

After the specimen absorbs some of the light, the Fourier transform decodes the signal into the infrared spectrum. Infrared energy is transmitted from the source via mirrors to

Interferometer Diagram



the beam-splitter where approximately half of the beam intensity is reflected toward the fixed-position mirror and the other half toward the moving mirror. The returning beam from each mirror travels back to the beam-splitter where the two beams are recombined, but the difference in path lengths creates constructive and destructive interference: an interferogram. The recombined beam passes through the sample. The sample absorbs all the different wavelengths characteristic of its spectrum, and then subtracts specific wavelengths from the interferogram. The detector now reports variation in energy versus time for all wavelengths simultaneously. A laser beam is superimposed to provide a reference for the instrument operation. A mathematical function called a Fourier tranform allows us to convert an intensity-vs-time spectrum into an intensity-vs-frequency spectrum.

The Fourier transform:

$$A(r) = \sum X(k) \exp(-2\pi \frac{irk}{N})$$

A(r) and X(k) are the frequency domain and time domain points, respectively, for a spectrum of N points.

The Fourier transform then converts the interferogram into a single-beam spectrum that is a representation of the same data in the frequency domain. That is, the single-beam spectrum shows the signal intensity for each frequency point in the range. After background data is taken (and then compensated for) this spectrum can be viewed in two modes: transmittance or absorbance.

In this project, tiny chips of resin composite, 10 to 30 microns thick, to be placed in the FTIR were removed from resin composite specimen using the tip of a scapel. The intensities of the double carbon bond (C=C) absorbance peak at 1637.3 cm⁻¹ and the aromatic (C...C) reference peak at 1608.3 cm⁻¹ were measured. The C...C peak originates from the aromatic rings in the Bis-GMA molecule and remains unchanged during the polymerization reaction. The ratio of the absorbance intensities of C=C/C...C were compared before and after polymerization using the following equation to determine the percent of reacted carbon double bonds or degree of conversion:

1 - [Abs (C=C) / Abs (C...C)] cured resin / [Abs (C=C) / Abs (C...C)] uncured resin X 100

Reference: Thermo Mattson Inc. "Chemist's Corner" website: www.mattsonir.com

KNOOP HARDNESS TESTER

(Kentron Microhardness, Torsion Balance Co., Clifton, NJ)

Hardness may be defined as the resistance to permanent surface indentation or penetration and is a force per unit area of indentation. The various hardness tests differ in the indenter material, geometry, and load. The indenter may be made of steel, tungsten carbide or diamond and be shaped as a sphere, cone or pyramid. The choice of hardness test depends on the material of interest, the expected hardness range and the desired degree of localization ¹⁸

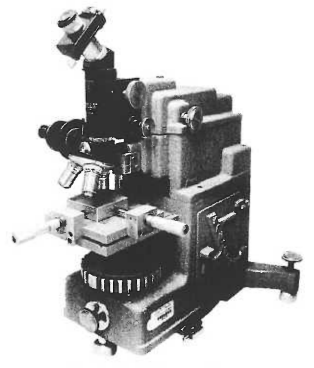


Photo by Patrick Pattamanuch

The Knoop indenter is cut in the shape of a diamond-based pyramid giving a diamond-shaped impression in which the long diagonal is very nearly 7 times the length of the short diagonal. Because of the difference in the lengths of the two diagonals, almost all of the elastic recovery of the indentations made with the Knoop indenter takes place in the transverse direction. Hence the measurement of the long diagonal together with the computed indenter constant gives a very close approximation of the unrecovered projected area of the indentation in square millimeters. The ratio of the applied load in kilograms to the approximate un-recovered projected area in square millimeters is called the Knoop Hardness Number for the specimen for that applied load.

The Knoop Hardness Number may be expressed by the formula:

$$KHN = L/Ap = L/l^2 x C_p$$
 where

KHN = Knoop Hardness Number

L = Load in kilograms applied to the indenter

indentation. For this indenter, $C_p = 0.07028$

Ap = Un-recovered projected area in square millimeters

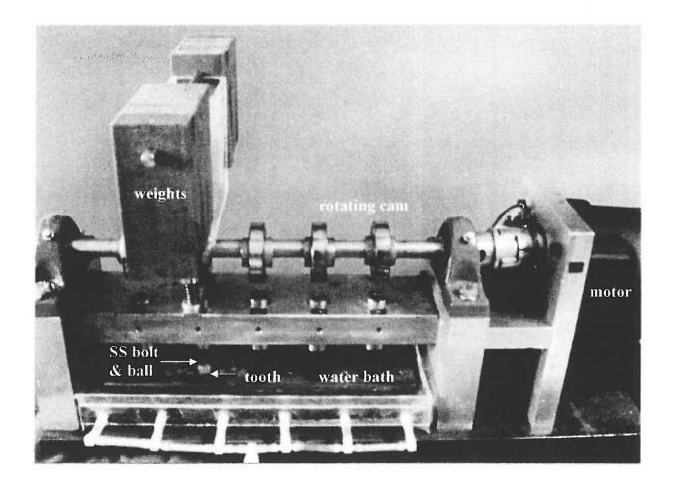
l = Measured length of the long diagonal of the indentation in millimeters

C_p = Constant relating length (l) to the un-recovered projected area of the

The Knoop indenter appears to be most useful in the investigation of the surface hardness of brittle materials such as glasses, hard carbides and oxides, thin layers of metals and small samples. The advantage of this method is that materials can be tested with a great range of hardness simply by varying the test load. The chief disadvantages are the need for a highly polished surface and a flat test sample.¹⁸

Reference: Kentron Micro Hardness Tester Operating Instructions

CUSTOM FATIGUE CYCLER



The OHSU custom-made fatigue cycler is capable of handling up to five specimens simultaneously. The number of dead weights and oscillations can be varied with a maximum of 85 Newtons and 1.25 Hertz. The unit operates with a DC motor rotating a cam system. The weights are supported by a spring assembly. The specimens may be bathed in re-circulating water at various temperatures with the use of a water bath. An electronic counter is available to terminate the oscillations at a preset number. The teeth were mounted in an acrylic ring. The ring and tooth assembly was locked into the cycler with adjustable plates. A 2 millimeter stainless steel sphere was cemented onto the occlusal surface of the resin composite restoration with chemically curing resin

cement. A stainless steel bolt, attached to the weight assembly system, was placed in contact with the surface of the steel sphere. The restorations were fatigued for up to 500,000 cycles from 18 to 85 Newtons at 1.25 Hertz while bathed in 37 degree centigrade water.

SCANNING ELECTRON MICROSCOPY

(JXA-6400, JEOL, Tokyo, Japan)

The Scanning Electron Microscope, or SEM, is an incredible tool for seeing the unseen worlds of micro-space. The SEM is capable of imaging samples of widely varying origin with a minimal preparation effort. A single SEM machine will allow magnifications from 10x up to 100,000x (the latter only under ideal conditions and with suitable samples). When an SEM is equipped with X-ray detectors, it becomes a microanalysis device capable of measuring the elemental composition of a sample with high spatial resolution. The composition of the sample can be studied on a micrometer scale and can be visualized as an X-ray map. Conventional light microscopes use a series of glass lenses to bend light waves and create a magnified image. The Scanning Electron Microscope creates the magnified images by using electrons instead of light waves. Samples have to be prepared carefully to withstand the vacuum inside the microscope. Biological specimens are dried in a special way that prevents them from contracting. Because the SEM illuminates them with electrons, they also have to be made to conduct electricity. Our SEM samples are coated with a very thin layer of gold by a machine called a sputter coater. The sample is placed inside the microscope's vacuum column through an air-tight door. After the air is pumped out of the column, an electron gun [at the top] emits a beam of high-energy electrons. This beam travels downward through a series of magnetic lenses designed to focus the electrons to a very fine spot. Near the bottom, a set of scanning coils moves the focused beam back and forth across the specimen, row by row. As the electron beam hits each spot on the sample, secondary

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electrons are knocked loose from its surface. A detector counts these electrons and sends

the signals to an amplifier. The final image is built up from the number of electrons

emitted from each spot on the sample.

Reference: Metalogic Products, Inc. www.metalogic.com