Study of chemical bond strength of methyl methacrylate (MMA) based bonding agent on type I dentin collagen at various humidity

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ABSTRACT

One of the basic agents used in dentin bonding solution is methyl methacrylate (MMA). This bonding agent is widely used in dentistry. It have been proved that the adhesion between dentin bonding agent and collagen fibril is chemically bond; though the chemical bonding contribution is smaller than physical mechanical bond. The purpose of the research was to examined the chemical bonding strength of MMA based dentin bonding on type I dentin collagen at various humidity. Samples of treatment group were put into desiccator with 60%, 70%, 80%, and 90% humidity, while for control groups at room humidity (65%). Chemical bond of pure MMA and MMA mixed with collagen were measured by FTIR. The lower the value of MMA carbonyl, the higher absorbance band speak of chemical bond strength between MMA and collagen. Data was statistically analyzed with One-Way ANOVA at 95% confidence level continued with Tukey-HSD test. The result showed that the highest chemical bond strength was at 65% humidity (p ≤ 0.05). In conclusion, many ester carbonyl MMA molecules reacted with amino collagen at 65% humidity. This can be shown by the lowest peak’s value of the MMA carbonyl absorbance at FTIR.

Key words: FTIR, humidity, MMA, type I dentin collagen, carbonyl group

INTRODUCTION

Science development of operative dentistry made bonding agents have been used excessively to restore dental caries. These agents are ideal to restore class V erosion which involving dentin, especially for the elderly patients. Since the bonding agents have hydrophilic characteristic, it can adhere well to dentin.1,2,3 Recently there are many bonding agents with hydrophilic and hydrophobic characteristic have been produced; such as bisphenyl dimethacrylate (BPDM), 4-methacryloxyethyl trimellitic anhydride (4-META) and others.

Bonding agents with hydrophilic characteristic will adhere to dentin collagen, while the one hydrophobic characteristic adhere on composite resin.3,4 Good wetting characteristics of bonding agents due to their low viscosity which increase surface energy. Surface energy is the potency of an agent surface to pull another substance surface. The adhesion between two agents occur if there is adhesive force at the interface area. The increase energy of each unit area is related with surface energy and surface tension.5 There are several monomer bonding agents available such as hydroxyethyl methacrylate (HEMA), BPDM, glutaraldehyde, 4-META, methyl methacrylate (MMA), etc. In order to make clinical application easier, usually the bonding agent is added with photo-initiator (camphoroquinone) which absorb the blue region of visible light spectrum at the wave-length from 400 to 500 nm.3,6 Accelerators from amine groups, which well interact with camphoroquinone such as dimethylaminoethyl methacrylate 0.15%, are often added to bonding agents. For long storage, the bonding agent is added with inhibitor agent, butylated hydroxytoluene 0.01%.

The adhesion of bonding agent on dentin surface occurs physical-mechanically or chemically. The physical-mechanical bonding is caused by penetration of bonding agent into nano space of collagen fibrils; resin penetration in dentin tubules; microscopically retention on dentin surface (undercut, crack, micro space) and Vander-Waals force based on dipolar attraction.2,3,4 Chemical bonding is caused by interaction of ester carbonyl groups of dentin bonding with amine groups at collagen which produce amide groups [C (O)NH]. This bonding is strong due to of its covalent characteristics called as inter-atomic primer bond.7 The bonding process of restoration materials on dental structure are a complex matter including MMA based resin bonding. The resin bonding failure are caused by some factors, such as smear layer on dental surface, topical application of fluoride, unhomogeneous tooth composition (organic and inorganic agents are very different in enamel and dentin), and saliva or blood contamination.

Dentin is life tissues which containing approximately 60% inorganic components (hydroxyapatite) Ca_{10} (PO_4)_6 (OH)_2, 30% organic components and 10% water. Those
organic components are 90% collagen, and 10% non-
collagenous. Most of those collagens are type I and few of
them are type V. Type I collagen which frequently used as
research agent are a sequence of amino acid: prolin, prolin,
glisin \( (H_2N-(pro-pro-gly)_5-COOH) \). Dentin bonding
can bond well to collagen fibril if the collagen is in active or
permeable condition. The permeability of collagen is highly
influenced by humidity of dentin surface. If the humidity
is too high or too low, the bonding agent will hardly bonds
on collagen fibril. Therefore, it requires optimum humidity
condition to obtained a maximum bond of resin bonding
agent to collagen.

The humidity used in the research were in the range
of 60 to 90%. The other factors that influence the bond
doing bonding to collagen are viscosity, type and
centrination of dentin bonding agents, application of
acid agents as conditioner and temperature of surrounding
collagen. This research used resin monomer MMA
\( C_5H_8O_2 \). The MMA solution is a clear and transparent
liquid at room temperature. The physical characteristics
as follows: melting point at \(-48^\circ C\), boilling point
at 100.8\(^\circ\)C, density = 0.945 g/ml (20\(^\circ\)C), heat of
polymerization = 12.9 kcal/mol.

The aim of the research was to examined the chemical
bond strength of MMA based bonding on type I dentin
collagen at various humidity.

**MATERIAL AND METHODS**

Material which were used: bovine type I collagen
(Sigma Chemical, St. Louis, USA; batch # 031K7054),
methyl methacrylate liquid (Vertex; Dentimex, Holland).
Instruments which were used: desiccators with vacuum
faucet (DSC, China), hygrometer (Haar. Synth-Hygro,
Germany), air suction (Schuco, USA), micro injection
(Hamilton, USA), Fourier Transformed Infra Red
instrument (FTIR, Jasco FT/IR 5300, Japan).

The research was conducted in Laboratorium Dasar
Bersama (LDB) Airlangga University with room humidity
of 65% and room temperature at 25 \(\pm\) 2\(^\circ\)C. The method
of this research has been presented in detail in a previous
research. To arrange 90% of humidity, 150 cc water
were put in the bottom part of desiccators, then calibrated
hygrometer was set. In this condition, the humidity seen
from hygrometer was 94–95%. Then from faucet located
above desiccators, the air is pumped out with air suction
until humidity reached to 90% and the pump directly closed.
For 80% humidity, the air in desiccators is pumped out until
humidity reached to 80%. In order to fasten the work, we
put the silica gel which is activated first in to desiccators.
Afterwards the air was pumped out until the humidity in
desiccators reached to 60%.

To prepared kalium bromide (KBr) pellets: 300 mg
KBr powder were dropped with 10 µl MMA (9.45 mg of
weight) using micro-injection. Molecular weight of MMA
was 0.945 g/ml. Total weight of KBr powder and MMA
were 309.45 mg, then all of the materials were crushed
with mortar and pestle made from agate stone. After mixed,
50 mg mixed-powder were put in to KBr die and compressed
until 10 ton while it’s was vacuumed. The final result of
the process was a clear pellet. The pellets were observed
into FTIR.

To make samples at various humidity: 2 mg collagen
fibers were packed into desiccators with hygrometer that
has been calibrated. Then the humidity of desiccators
were arranged to 60%, 70%, 80%, and 90%. As soon as release
from the desiccators, the collagen was dropped with MMA
and added with KBr powder until reached 309, 45 mg of
weight. The mixed powders were compressed, and then 50 mg of the mixtures were put it in to KBr die. Next, the samples are compressed until 10 ton while it has been vacuumed. That pellets were finally observed in to FTIR.

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weight. The mixed powders were compressed, and then 50 mg of the mixtures were put it in to KBr die. Next, the samples are compressed until 10 ton while it has been vacuumed. The peak of carbonyl absorbance band (C = 0) of treatment groups and control groups was measured. The method to calculate the peak of carbonyl (P) absorbance band as below.

\[
P = \left( \frac{BC}{AB} \right) \times 100;
\]

where \(AB\) and \(BC\) measured in
transmittance, \(T = 30\%\) to 60%.

**Figure 1.** Peak of the MMA carbonyl absorbance band.

**Description:**

- \(P = \left( \frac{BC}{AB} \right) \times 100;\)
- \(AB\) and \(BC\) measured in
- centimeter. The calculation is accurately enough and trustworthy if the intensity of absorbance band at transmittance, \(T = 30\%\) to 60%.
RESULTS

Figure 2. The IR ray spectrum of MMA and collagen (KBr pellets).
Value of the peak carbonyl absorbance band is 33.3; point 7 (arrow).
Humidity: 80%.

Figure 3. The IR ray spectrum of MMA + collagen (KBr pellets).
Value of the peak carbonyl absorbance band is 29.2; point 11 (arrow).
Humidity: 90%.

Chemical bond measurement is done by using KBr pellets in which then irradiated by the IR ray. After that the peak value of the MMA ester carbonyl absorbance band was recorded. When MMA carbonyl groups excessively bond toward collagen amino groups, the peak value of the MMA carbonyl absorbance band decreased. The more MMA carbonyl groups bond with amino collagen groups the bond between those agents will increase. Mean value of MMA carbonyl absorbance band and standard deviation were shown at Table 1.

Mean value of the MMA carbonyl absorbance was 48.7 ± 3.7 then decreased at 60%, 65%, 70%, 80%, and 90% humidity. To know whether the chemical bond data was normal, one sample of Kolmogorov Smirnov test was done. The p value at the experimental groups at 60–90% humidity showed the higher number than 0.05 (p > 0.05). It means that the experiment of chemical bonds groups at 60–90% humidity was homogen.

Table 1. Mean of the peak of MMA carbonyl absorbance and standard deviation at various humidity

<table>
<thead>
<tr>
<th>Humidity</th>
<th>N</th>
<th>(\bar{X}) (mean)</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMA</td>
<td>5</td>
<td>48.7200</td>
<td>3.73992</td>
</tr>
<tr>
<td>60%</td>
<td>5</td>
<td>27.3600</td>
<td>2.21314</td>
</tr>
<tr>
<td>65% (control)</td>
<td>5</td>
<td>19.1800</td>
<td>1.74557</td>
</tr>
<tr>
<td>70%</td>
<td>5</td>
<td>20.5000</td>
<td>1.33791</td>
</tr>
<tr>
<td>80%</td>
<td>5</td>
<td>30.9400</td>
<td>1.56301</td>
</tr>
<tr>
<td>90%</td>
<td>5</td>
<td>34.5000</td>
<td>3.09031</td>
</tr>
</tbody>
</table>

Description:
N = Amount of sample; \(\bar{X}\) = Mean of the peak of the MMA carbonyl absorbance band; SD = Standard Deviation.

Table 2. The Tukey-HSD test in carbonyl absorbance band of MMA at various humidity

<table>
<thead>
<tr>
<th>Dentin bonding agent based on MMA</th>
<th>Significance level</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMA 60% humidity</td>
<td>21.360*</td>
</tr>
<tr>
<td>MMA 65%</td>
<td>29.540*</td>
</tr>
<tr>
<td>MMA 70%</td>
<td>28.220*</td>
</tr>
<tr>
<td>MMA 80%</td>
<td>17.780*</td>
</tr>
<tr>
<td>MMA 90%</td>
<td>14.220*</td>
</tr>
<tr>
<td>60% humidity MMA</td>
<td>– 21.360*</td>
</tr>
<tr>
<td>65%</td>
<td>8.180*</td>
</tr>
<tr>
<td>70%</td>
<td>6.860*</td>
</tr>
<tr>
<td>80%</td>
<td>– 3.580</td>
</tr>
<tr>
<td>90%</td>
<td>– 7.140*</td>
</tr>
<tr>
<td>65% humidity MMA</td>
<td>– 29.540*</td>
</tr>
<tr>
<td>60%</td>
<td>– 8.180*</td>
</tr>
<tr>
<td>70%</td>
<td>– 1.320</td>
</tr>
<tr>
<td>80%</td>
<td>– 11.760*</td>
</tr>
<tr>
<td>90%</td>
<td>– 15.320*</td>
</tr>
<tr>
<td>70% humidity MMA</td>
<td>– 28.220*</td>
</tr>
<tr>
<td>60%</td>
<td>– 6.860*</td>
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<tr>
<td>65%</td>
<td>1.320</td>
</tr>
<tr>
<td>80%</td>
<td>– 10.440*</td>
</tr>
<tr>
<td>90%</td>
<td>– 14.000*</td>
</tr>
<tr>
<td>80% humidity MMA</td>
<td>– 17.780*</td>
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<td>70%</td>
<td>10.440*</td>
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<tr>
<td>90%</td>
<td>– 3.560</td>
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<tr>
<td>90% humidity MMA</td>
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Note: *: significant difference at \(\alpha = 0.05\)
ANOVA test was used to know the influence of humidity towards chemical bonds between MMA and dentin collagen. There was a significant difference among the experimental groups (p < 0.05). Turkey-HSD test was done to give evidence of distinctions for every experimental group. Peak value of the MMA carbonyl absorbance band among the experimental groups at 60%, 65%, and 70% humidity showed significant difference (p < 0.05). There was no significant difference (p > 0.05) between 80% and 90% humidity in experimental groups. Experimental groups with 70% humidity compared with 60%, 80% and 90% humidity showed significant difference (p < 0.05) and if compared with 65% humidity showed no significant difference (p > 0.05). Furthermore, in experimental group with 90% compared with 60%, 65%, and 70% humidity showed significant difference (p < 0.05), but not for 80% humidity (p > 0.05).

**DISCUSSION**

Infra red spectrum also can be used to count the quantitative value of the of mixing matter components. However the number of each factor definitely must be calculated accurately in order to get accurate and trustworthy datas.

If collagen fiber mixes with MMA, chemical reaction between amino group of collagen and ester carbonyl group of MMA will occur. In fact, not all of the ester carbonyl group of MMA fully reacted with amino collagen groups. This matter can be seen at peak of the mixture of MMA carbonyl and amino collagen absorbance band which is decrease but not disappear. Esther carbonyl wave number ($v_{C=O}$) absorbance ranging around 1700 to 1750 cm$^{-1}$.

In this research, the peak of ester carbonyl at MMA absorbance band is used as the control or comparison with the peak of ester carbonyl absorbance band at experimental groups by various kind of humidity treatments. If many carbonyl groups at MMA react with amino collagen groups, the IR ray spectrum will occur the reduction of peak of the mixture of ester carbonyl at MMA and collagen absorbance band. So it can be assumed that if there is great reduction of peak of the MMA carbonyl absorbance band, it means that will be there are many ester carbonyl groups bond with amino groups. This means that chemical bond between MMA and collagen can be stronger than ever.

Chemical bond between bonding agents MMA based and collagen fibril should be considered even though from the previous research chemical bond only 30% compared with physical-mechanical bond strength. The chemical bond strength depends on the condition of surrounding collagen, such as humidity and temperature.$^{5,19}$ The study used constant temperature (25 ± 2 °C). At an optimum humidity, the condition of collagen fibril is very permeable, not only made a strong bonding between MMA carbonyl group and collagen amino group, but also a maximum hydrogen bond inside molecule chain, or among collagen molecules. Moreover a good penetration of MMA into nano interfibriler space will occur and polymerization of MMA will form a mechanical retention.

If the humidity of surrounding dentin is too high, MMA solution difficult to react with collagen fibril, because many water molecules in surrounding fibril will block the MMA penetration. An excessive hydrogen bond between water molecule and dentin collagen made bonding between collagen molecule and MMA can not chemically occur. Reis et al.$^{20}$ reported that the use of air stream is not effective to remove water from mixture of bonding agents and water. The location of the highest water concentration is predicted in the deepest collagen fibril (profunda site), so it is difficult to remove water using air stream and the bonding agents is difficult to diffuse. A study of Micro-Raman Spectroscopy showed that monomer concentration decrease from the upper side to hybrid layer base which formed by adhesive resin. For example, bonding agents concentration will decrease almost 55% at 2 μm under surface region and decrease almost 21% at hybrid layer base. But if the surrounding dentin is very dry (minimum humidity) the collagen will collapsed. It will make the hydrogen chain of inter and intra molecule collagen are broken and collagen amino group are covered with remain of collagen fibril. The condition not only makes amino collagen groups can not react with MMA carbonyl groups, but also interaction between functional groups of MMA and functional group (carboxylate, amino, amide) collagen do not occur and no complex reaction between Ca$^{++}$ dentin and MMA. It has been reported that in water free region (dry area), the polymer chain formation is weaker and unstable. The pH of monomer solution plays role in chemical bond strength between bonding agent and collagen.$^{52}$ The experiment, proved that high bonding strength between HEMA and collagen occur at pH 2. If pH rise up to 6.6, the bonding strength will decrease. However if pH is rise from pH 9.0 to 12.5, the bonding strength will increase again. Bonding strength is studied using method of collagen functional groups dissociation, such as, carboxylate and amino. If carboxylate acid dissociation or amino group is being inhibited, the hydrogen bond between resin and collagen will increase. If pH of collagen suspension is low, carboxylate acid will not be dissociated, so the interaction of resin and collagen will increase. The pH of MMA solution in the study was 5.0.

Nakabayashi and Pashly$^5$ reported that the process of collagen collapse known as passive theory. Passive theory is demineralized collagen network will float or suspended in water. Each fibril will be separated one another by water molecule in the space between fibril. Prior to water molecule the space is filled by apatite crystal. At drainage process, the water which supports collagen network will be eliminated, collagen will close and aggregate three dimensionly. This condition is called collapse or shrinkage. The fibril bonding becoming stiff and creating the interfibriler hygrogen bond which interact electrostatically or hydrophobically.$^5$ Water additional changed the condition into passive
re-expand. Water molecule can perform hydrogen bond with collagen peptide, breaking interfebriler hydrogen bond so the collagen network is re-expand. The re-expand network functions as hydrogel, where water osmotically entering interfibriller space.

In this research, pure MMA solution is used to examine role of MMA on chemical bond to pure collagen. Mean of band’s peak of the MMA carbonyl absorbance is 48.7 ± 3.7, this value decrease at 60% humidity (27.3 ± 2.2); 65% humidity (19.1 ± 1.7); 70% (20.5 ± 1.3); 80% (30.9 ± 1.5) and 90% (34.5 ± 3.1). The lower the band’s peak value of the MMA carbonyl absorbance the greater the bonding of MMA carbonyl groups to collagen amino groups. It makes the bonding strength between two agents increase. The highest strength of chemical bonds occur at 65% humidity, since the water molecule surrounding around fibril in optimum condition. Water molecule makes collagen re-expand, and collagen become more active and permeable to MMA. Comparing with previous research, the bond strength of bonding agents HEMA based and MMA based are different. The highest bonding strength of HEMA based, is at 70% humidity, while bonding agent MMA based is at 65% humidity. It may be caused by the difference of agent density. HEMA’s density is 1.07 g/ml while MMA density is 0.945 g/ml.

Table 1 showed that mean of the band peak’s value of MMA carbonyl absorbance from 60% to 90% humidity are significantly increase. It caused chemical bond strength decrease. The increase of humidity makes water molecules around fibril rise and the penetration of MMA solution to collagen fibril is blocked. If we compare the 65% and 60% humidity, chemical bond strength at 65% humidity is higher than 60%, since at 60% humidity, the condition of fibril is drier, so as the results fibril collapsed and low chemical interaction between MMA and collagen occur.

The research suggest that maximum chemical bond strength between MMA carbonyl groups and amino collagen groups occur at optimum humidity of 65%.

REFERENCES


