Effect of Copper Content on Compressive Strength and Microstructure of Dental Amalgams

Akram Hasheminezhad, Seyed Mojtaba Zebarjad*, Seyed Abdolkarim Sajjadi, Laleh Rahanjam
Department of Metallurgy and Materials Engineering, Engineering Faculty, Ferdowsi University of Mashhad, Mashhad, Iran
Email: *Zebarjad@um.ac.ir
Received November 18, 2011; revised January 4, 2012; accepted January 10, 2012

ABSTRACT
The main goal of this research is to investigate the effect of copper on compressive strength of dental amalgam. For this purpose amalgam capsules with two different content of copper were used. Cylindrical samples with diameter to height ratio 1 to 2, were prepared via molding method. To evaluate the role of copper element on compressive strength, compression test was done at different strain rates i.e. 0.02, 0.2, 0.4 and 2 min⁻¹. The results and microscopic evaluations showed that an increase in copper content in amalgam caused to eliminate gamma2 phase and leaded to promote compressive strength.

Keywords: Amalgam; Copper; Compressive Strength; Microstructure; Gamma2

1. Introduction
Dental amalgams constitute almost 75% of all restorative materials used in dental prak tice. They are obtained from the combination of mercury and a silver-copper-tin powder [1]. Early dental amalgams (1900-1965) composed of 66% - 67% Ag, 25% - 28% Sn, 0% - 6% Cu and 1% Zn so called as low copper amalgam. Reaction that occurs is observed below [2]:

\[ \text{Ag}_2\text{Sn}(\gamma) + \text{Hg} \rightarrow \text{Ag}_2\text{Hg}_3(\gamma_1) + \text{Sn}_{1.8}\text{Hg}(\gamma_2) + \text{Ag}_3\text{Sn(reacted)} \]

The reaction products are about 54% - 56% Ag2Hg3 (γ1) phase, 32% - 35% Ag3Sn (γ) phase and 11% - 13% Sn7-8Hg (γ2) phase. The γ2 phase is the most corrosion prone phase and its hardness is almost 10% of that of γ1 phase [3]. In order to eliminate γ2 phase, in 1965, the high copper amalgam with 12% - 30% copper were produced. They are much more corrosion resistant. In this amalgams copper reacts with tin and Cu6Sn5 phase is formed instead of Sn7-8Hg [4,5]. Thus it eliminates γ2 phase. This reaction is shown below [2]:

\[ \text{Ag}_2\text{Sn}(\gamma) + \text{Ag} + \text{Cu}(	ext{eutectic}) + \text{Hg} \rightarrow \text{Ag}_2\text{Hg}_3(\gamma_1) + \text{Cu}_6\text{Sn}_5(\eta) + \text{Ag}_3\text{Sn(reacted)} + \text{Ag} + \text{Cu}(	ext{reacted}) \]

Although there are a lot of papers in the field of physical and chemical properties of dental amalgam [6-14], only a few consider the mechanical behavior of this material.

2. Experimental Procedure
2.1. Materials
Two series of alloy powders and mercury were prepared. According to manufacturer claim the type of both alloy powders was lathe-cut and spherical. SEM micrograph and analysis of this powder are shown in Figures 1 and 2. The details of used materials are summarized in Table 1. Just for simplicity the materials were coded. The main goal of this research is to find out the effect of copper on compressive strength of dental amalgams.

2.2. Sample Preparation
The alloy powders and mercury were mixed mechanically at different ratio. Table 2 illustrates the mixing condition.

Figure 1. SEM micrograph of admixed high copper amalgam.
Figure 2. SEM analysis of admixed high copper amalgam.

Table 1. Specification of used alloy powders.

<table>
<thead>
<tr>
<th>Material</th>
<th>Commercial name</th>
<th>Percent of Ag (wt%)</th>
<th>Percent of Sn (wt%)</th>
<th>Percent of Cu (wt%)</th>
<th>Percent of Zn (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>SDI, permit</td>
<td>56</td>
<td>27.95</td>
<td>15.4</td>
<td>2</td>
</tr>
<tr>
<td>S2</td>
<td>SDI, gs-80</td>
<td>40</td>
<td>31.35</td>
<td>28.7</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 2. Mixing condition.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Per capsule</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>400mgl Alloy, 364mgl Hg</td>
</tr>
<tr>
<td>A2</td>
<td>600mgl Alloy, 552mgl Hg</td>
</tr>
</tbody>
</table>

Figure 3. Mold and prepared specimens.

2.3. Microscopic Evaluation

To study the role of copper element on amalgam microstructure, the surface of some samples were polished to achieve a surface roughness of about 5μm. Then the polished surfaces of the samples were investigated using both optical and SEM microscopes.

2.4. Mechanical Test

Compressive test were carried out with a Zwick tensile machine at different strain rates 0.02, 0.2, 0.4 and 2 min⁻¹. Microhardness measurement carried out with a Vickers indenter under a load of 1 Kg and 500 g.

3. Results and Discussion

The microstructure of the A2 sample shows a continuous Ag-Hg matrix that simultaneously is formed with Cu₆Sn₅ phase and embeds η phase and Sn-Ag alloy particles. Also Ag-Cu eutectic particles are surrounded with η phase. In this sample in addition to η layers, a small amount of η crystals are found (Figures 4-6).

Figures 7-9 show microstructures of A1 samples. Comparison of both microstructure (i.e. A1 & A2) shows gamma₂ phase amount decreases and eta phase amount increases as copper content add.

Figures 10 and 11 illustrate the dependency of fracture stress of amalgam alloy on copper content. However residual Ag₃Sn particles in amalgam increase, amalgam strength is also increased. This is important because the matrix phase is the weaker and more corrosion prone than the residual alloy particles gamma phase is stable in mouth periphery.

Since gamma₁ phase plays like a barrier against crack propagate, thus fracture of amalgam alloy should be under control of copper content. Figure 10 and 11 illustrate the dependency of fracture stress of amalgam alloy on copper content. Figure 10 and 11 illustrate the dependency of fracture stress of amalgam alloy on copper content. Further residual Ag₃Sn particles in amalgam increase, amalgam strength is also increased.

This is important because the matrix phase is the weaker and more corrosion prone than the residual alloy particles gamma phase is stable in mouth periphery. Also pure gamma₁ phase is stable but existence gamma₁ in amalgam have a small quantity tin that is the most corrosion prone phase in mouth, so gamma₂ phase have low corrosion resistant and low compressive strength and can cause to crack propagation in amalgam. Furthermore interface between gamma phase and matrix is very magnitude. If interface strength is not coefficient can be caused crack nucleation and growth. Also Cu₆Sn₅ phase increases the strength of high copper amalgams.

for making amalgam. After mixing and before amalgamation, the standard samples were made using molding method. The heights of the samples were varied from 11 to 13 mm. Mold and prepared specimens are shown in Figure 3.

Figure 2. SEM analysis of admixed high copper amalgam.
Figure 4. The microstructure of the A2 sample.

Figure 5. SEM micrograph of A2 sample.

Figure 6. Optical micrographs of A2 sample.

Figure 7. SEM micrograph of A1 sample.

Figure 8. Gama1 and gama2 appear in A1 sample.

Figure 9. Optical micrograph taken from the polished surface of A1 sample.
copper content (A1), alloy powder consists of more lathe-cut particles than spherical particles. So particles total area and also required mercury content for diffusion in particles increase, on the other hand mercury diffusion causes to contraction, therefore contraction for this amalgam decreases. On the other hand $\gamma_1$ and $\gamma_2$ phases formation result in expansion. These changes neutralize one another in a good amalgam, but usually its expansion is more than contraction. It induces porosity that is a location for stress concentration and decreases compressive force, thus spherical particles not only harden rather stronger in compression than lathecut particles and Figure 12 Confirms this claim.

Whereas compressive strength is a function of strain rate, samples were tested in 0.02, 0.2, 0.4 and 2 min$^{-1}$ strain rates. The required force versus strain rate for A1 and A2 samples are shown in Figure 13. It can be observed a decrease in force with increasing strain rate for both samples. Also can be seen that A1 sample has less fracture force than A2 sample. Deviation of displacement and elasticity modulus versus strain rate as a function of copper content in turn are shown in Figures 14 and 15. Results of microhardness measurements (Table 3) are in agreement with results of compression test.
4. Conclusions

The results of study may be summarized as follows:

1) Addition of copper content to dental amalgam alloy causes an increase in compressive strength and hardness.

2) Required fracture stress for amalgam with specific Cu weight percent (wt% Cu) is raised when strain rate increases.

3) Increasing copper content in amalgam alloy powders cause to deform the powder particles from lathe-cut to spherical.

REFERENCES


