Characteristics of hardness and microstructure of extraction forceps for dental and oral care made of stainless-steel

Nur Kholis¹, Nuryanto Nuryanto², Arif Mustofa³

¹-²Department of Mechanical Engineering, Faculty of Engineering, Universitas Wahid Hasyim, Semarang. 50236. Indonesia.

Corresponding author: nurkholis@unwahas.ac.id

Permalink (DOI): https://doi.org/10.23917/arstech.v1i2.184

ARTICLE INFO

Article history:
Received 20 Sep 2020
Revised 10 Oct 2020
Accepted 23 Nov 2020
Available online 29 Nov 2020
Published regularly 01 Dec 2020

Keywords:
Hardness
Microstructure
Stainless-steel
Scanning electron microscopy (SEM)
Medical material

ABSTRACT

The reliability of medical devices such as extraction forceps is vital for dental and oral care. Apart from having hygienic properties, the extraction forceps must be strong and resistant to corrosion. This study evaluates the effects of tempering temperature on the hardness and microstructure of a medical device's material made from stainless-steel DIN 4021. In the experiments, a heat treatment process was carried out previously with a temperature of 1,050°C and a holding time of 20 minutes. A quenching process was conducted using a cooling channel that flowed with water at 10-20°C. After the heat treatment, the material was subjected to a tempering process with temperature variations of 200, 400, and 600°C. The research results indicated that the heat treatment process could increase the material's hardness—the hardness of the raw material changed from 20 to 48.67 HRC with the heat treatment. The tempering parameters resulted in the highest hardness of 46.67 HRC at 200°C and the lowest value of 42.33 HRC at 600°C. Microstructure testing using optical microscopy showed that it produced ferrite, pearlite, and martensite structures. In contrast, the result of a microstructure testing using Scanning Electron Microscopy on the surface of the material is that the higher the tempering temperature, the greater the microstructures' dimension.

1. INTRODUCTION

Stainless steel or corrosion-resistant steel is known as magic metal has been widely used for human life. This material has reliable mechanical properties and good corrosion resistance [1]. Its applications range from everyday household goods to applications in boilers, automotive, biomedical, and chemical industries [2-4]. In an earlier time, stainless-steel has been applied to produce the best and most durable products. However, the materials or products made of stainless-steel could only be purchased or accessed by the upper class. As time goes by,
the application and production of products with stainless-steel has increased rapidly and become more common and accessible. As a result, the rapid development of technology has significantly impacted the industry, particularly medical devices, for oral and dental care. Most of these tools take advantage of stainless-steel as the raw material due to its corrosion resistance [5][6].

It is necessary to give a heat treatment of material, such as tempering appropriately in the production process. Heat treatment is commonly applied to the raw material to alter its softness property and increase its hardness as needed [7]. Heat treatment is a combination of heating and cooling a solid material for a certain period to obtain certain properties [8]. As recommended by the ISO 13485 standard, medical devices’ permissible hardness is 43 ± 3 HRC [9].

As commonly known, the negative impact of heat treatment is that the material examined hardness is above the desired standard and becomes brittle. However, tempering can overcome the effects after completing the heat-treatment process. The tempering process may reduce the hardness and the brittle property of a material [10]. The tempering is carried out by reheating a hardened material at a temperature below the critical point to obtain certain ductility and then quenching it. In principle, the process is intended to reduce the brittleness and hardness to meet its use requirements. In addition, tempering serves to increase its toughness and ductility without drastically reducing its hardness [11].

Besides the hardness, the quench hardening process also affects a material's microstructure [12]. The hardening process above the austenite temperature and the quenching process carried out on the raw material produce a martensite phase. The property of martensite microstructures tends to be hard and brittle because of their high hardness [13]. The increased austenitising temperature raises the amount of residual austenite content [10]. The retained austenite subsequently erodes Chromium (Cr) and improve the anti-rust film. The corrosion resistance of martensitic stainless-steel to the carbide volume fraction is closely related to the carbide precipitation during tempering [14].

The heat treatment process involving hardening, quenching, and tempering turns the microstructures of the steel’s initial carbon into martensite, pearlite, or bainite microstructures. This transformation of microstructure also alters its physical and mechanical properties. Therefore, a more in-depth study regarding this kind of change is needed, especially a stainless-steel material for medical devices [15].

Several kinds of research on the material properties of stainless-steel DIN 4021 have been conducted to analyse the effect of a holding time in the material properties' tempering process. A holding time significantly impacts the material properties in terms of hardness and microstructure form [3][16].

This research examined the relationship between the tempering temperature and the material’s hardness concerning the problem discussed above. The study analyses the relationships between the tempering temperature and the microstructural change using optical microscopy (OM) and scanning electron microscopy (SEM). This study considered some treatment of holding time carried out by Jiang et al. [17].

2. RESEARCH METHODOLOGY

This research was realised experimentally in the Material Laboratory of the Faculty of Engineering, Universitas Wahid Hasyim Semarang Indonesia. Figure 1 shows a product of extraction forceps and the samples of the cut specimen. This product is a medical device for oral and dental care, considering that stainless-steel has a relatively high corrosion resistance [18]. The material of stainless-steel DIN 4021 contains C=0.16 - 0.25%, Si ≤ 1, Mn ≤ 1.5, Cr=12-14, Mo=0, and Ni=0.25.

Figure 1. (a) Extraction forceps, (b). Specimen.

Several steps were taken in this experiment, including preparing for the raw material, cutting it into samples, heating them in a furnace room, quenching, and re-entering them into the furnace room for tempering according to the parameters. Hardness testing and microstructure testing were then carried out to evaluate changes in mechanical and physical characteristics.

In general, the heat treatment process's first step was to heat a metal or another alloy at a specific temperature, hold it for a particular period, and then cool it down at a specific cooling rate. Microstructural changes would appear in terms of the phase, shape, or size of the crystal grains in the heating and cooling process. This transformation would alter the properties of the metal or alloy as well.

Heat treatments were carried out on a material with a temperature of 1,050°C in a furnace and a holding time of 20 minutes, followed by quenching. The quenching was mainly intended to increase the hardness of steel, and it was typically realised to obtain better wear resistance or strength as suggested by some researchers [3][19][20]. Coupled with a rapid cooling rate, it would result in hard
martensite. Concerning the study, the quenching process was carried out in a cooling canal with a temperature range of 10 - 20°C, as seen in Figure 2.

Figure 2. Heating furnace and quenching canal.

After heating and quenching, tempering was performed with temperature variations of 200, 400, and 600°C with a holding time of 20 minutes. As known in the open literature, the tempering process was oriented to reduces steel's residual stress and increases its toughness and ductility [21]. During the tempering process, the steel would decrease hardness and strength, including an increase in ductility. A decrease would follow the brittleness. The tempering process was carried out in a heating furnace, as shown in Figure 3.

Figure 3. Tempering furnace.

Once the hardening tempering process was completed, a series of hardness and microstructural testing was employed for all specimens referring to the ASTM E384 [22]. The hardness of the materials was tested to examine their mechanical properties. Even though the measurement was carried out only at specific points or areas, the hardness obtained were valid to describe their strength. By this kind of testing, materials would be much easier to be classified as ductile or brittle. Microstructural testing was done to determine the phase arrangement of a test object or specimen. The shapes and sizes of metal crystals, deformation-driven metal damage, and beyond could be identified through the microstructural examination. Before all specimens entered the testing phase, some stages had to be done, such as mounting, grinding, polishing, and etching as other researchers [23][24]. All the preparation process are presented in Figure 4.

Figure 4. Specimen preparation. (a) Mounting process. (b) Polishing. (c) Etching.

Figure 4(a) is the process of mounting the materials to manage the polishing process. Figure 4(b) is the polishing process after the grinding process with a grit of sandpaper P120, P220, P400, P800, P1000, P1500, P2000. The results have to be smooth like a mirror, with no scratches. Figure 4(c) is the etching process using chemical liquids, namely HNO₃, HCl, and alcohol, as done by Kisason et al. [24].

3. RESULTS AND DISCUSSION

3.1. The characteristics of microstructure using optical microscopy

Figure 5 gives the microstructure of raw material with 200X and 400X magnification based on the optical microscopy testing.

Figure 5. The microstructure of raw material. (a) 200X magnification. (b) 400X magnification.
From the microstructural testing, the raw material contains both ferrite and pearlite structures. The ferrite structure is white in colour, rough in shape, and typically with a maximum carbon content of 0.025% at 723°C. Besides, the structure is soft, ductile, and has a body centre cubic (BCC) crystal structure, as found by another researcher [25]. On the other hand, the pearlite structure is grey and large (coarse pearlite). The pearlite structure was formed from two phases, namely the ferrite phase and the cementite phase.

Figure 6 shows the microstructural testing results on stainless steel due to hardening treatment at a temperature of 1,050°C and a holding time of 20 minutes. The resulting microstructure consists of soft and dense ferrite structure (light colour), smooth and thick pearlite structure (grey colour), and martensite structure. The blade martensite microstructure is related to a mixture of ferrite and cementite phases, not in lamellar but cementite needles. The martensite phase's existence is due to a rapid cooling rate, preventing the transformation process's atomic particles. This structure has typically hard and brittle properties. It is found that the pearlite and martensite microstructures are more dominant and denser. Consequently, as the austenite temperature in stainless steel increases, the austenite grains grow significantly, resulting in coarse martensite grains rather than fine ones. The coarse grains' sizes provide less nucleation for the pearlite transformation, causing the martensite matrix to be more challenging.

Figure 6. The microstructure of 1,050°C hardened material. (a) 200X magnification. (b) 400X magnification.

The materials used are hardened at 1,050°C, held for 20 minutes, and then quenched in a cooling canal. Subsequently, the material is tempered for microstructural testing. The microstructural testing is carried out at varied tempering temperatures of 200, 400, and 600°C held for 20 minutes. These temperatures are further classified as low, medium, and high, respectively. The tempering treatment is principally meant to reduce the risk of cracks and internal stress resulting from a rapid cooling rate. The primary purpose is to compare the particle structures' resulting microstructures and shapes after various tempering temperatures. The results of the materials' microstructures with varying temperatures are given in Figure 7.

Figure 7. Results of microstructures of tempered materials. (a) The temperature of 200°C. (b) The temperature of 400°C. (c) The temperature of 600°C.

Ferrite, pearlite, and tempered martensite microstructures are formed at a tempering temperature of 200°C, referring to Isfahany et al. [18]. The microstructure of tempered martensite comes from the decomposed hardened martensite phase. Therefore, it appears to be more ductile but remains challenging. At 200°C, the microstructure is dominated by tempered martensite structure and pearlite structure, and the atomic arrangement was smoother and tighter. This result is not much different from that of the hardened material. Accordingly, the tempering temperature below 200°C has no significant effect on the microstructure and mechanical properties, as found by Sayed and Kheirandish [26].

At a tempering temperature of 400°C, ferrite, pearlite, and slightly tempered martensite microstructures have been observed. At that temperature, ferrite and pearlite microstructures are quite dominant.

At 600°C, ferrite and pearlite microstructures are clearly seen. The ferrite and pearlite phases are formed when the austenite phase is altered due to a slow cooling rate. As the tempering temperature increases, the martensite phase shifts into the pearlite and ferrite phases. This transformation eventually changes the physical and mechanical properties.
3.2. The characteristics of microstructure using scanning electron microscopy

Microstructural testing using scanning electron microscopy aims to determine the particles’ morphology and size distribution on the specimen materials’ surfaces [27-29]. The particle analysis was carried out to obtain the mean areas and diameters of the particles for all tested specimens, namely the raw material, 1,050°C hardened specimens, tempered specimens at 200, 400, and 600°C with 2000X magnification. The analysis was performed using ImageJ software and the Threshold method, as suggested by Banerjee et al. [30][31]. The image analysis results using scanning electron microscopy are provided in Figure 8 for the raw material.

![Figure 8. The SEM image processing using ImageJ and Threshold. (a) initial image. (b) Threshold image.](image)

There is a difference between the photos in terms of brightness for the initial SEM image and the Threshold image. The measurement of the raw material using Threshold resulted in a rounded and broad form. This form is similar to that of the microstructural testing using Optical Microscopy as provided in Figure 5.

Figure 9 indicates the results of SEM images of the 1,050°C hardened material using ImageJ and Threshold. The threshold method is applied to measure the average area and dimension of the material's particles. The results indicated that the structure's shape is similar to that of the microstructural testing using Optical Microscopy, as shown in Figure 6.

![Figure 9. SEM image processing using ImageJ and Threshold.](image)

Table 1 provides the calculation results of the particles' average sizes in the raw material and the heat treatment specimen. The raw material particle is 465.689 μm², while that of the heat treatment specimen is 0.099 μm². It means that the raw material's structure is more extensive than that of the heat treatment specimen's construction, following the fact that the heat treatment specimen has smooth structures. The quenching process at a rapid speed prevents the atoms from transformation and further influenced its hardness. The smaller the dimensions, the higher the hardness, as displayed in Figure 9.

<table>
<thead>
<tr>
<th>No</th>
<th>Specimen</th>
<th>Area of Particle (μm²)</th>
<th>Particle Diameter (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Raw Material</td>
<td>465.689</td>
<td>2.596</td>
</tr>
<tr>
<td>2</td>
<td>Heat Treatment 1050°C</td>
<td>0.099</td>
<td>0.330</td>
</tr>
</tbody>
</table>

The tempered specimens have been analysed at 200, 400, and 600°C with 2000X magnification using SEM. The results of SEM image processing with ImageJ and Threshold are shown in Figure 10.

![Figure 10. SEM image processing with ImageJ and Threshold at various tempering temperature.](image)

Table 2. Calculation results of average particle sizes

<table>
<thead>
<tr>
<th>No</th>
<th>Tempering temperature</th>
<th>Particle Area Average (μm²)</th>
<th>Average Particle Diameter (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>200°C</td>
<td>0.149</td>
<td>0.392</td>
</tr>
<tr>
<td>2</td>
<td>400°C</td>
<td>0.255</td>
<td>0.492</td>
</tr>
<tr>
<td>3</td>
<td>600°C</td>
<td>0.527</td>
<td>0.587</td>
</tr>
</tbody>
</table>

Based on Table 2 and Figure 10, the specimens' analysis with varying temperatures of 200, 400, and 600°C results in the particle areas of 0.149, 0.255, and 0.527 μm², respectively. It also produces the average particle diameters of 0.392, 0.492, and 0.587 μm, respectively. Therefore, it can be inferred that the higher the tempering temperature is, the larger the size of the average particle area and
diameter will be. A high tempering temperature causes a slow cooling pace, allowing the atomic particles to transform with ease. With such alteration, the hardness of the material decreases.

3.3. The characteristics of hardness

In the study, Rockwell Hardness scale C has been applied to examine the hardness as suggested by ASTM [32]. As seen in Table 3, the hardening and quenching process increases the hardness of Stainless-steel DIN 4021. The hardness of the material changes from 20 to 49 HRC. At a hardening temperature of 1,050°C, the hardness reached 49 HRC as shown in Figure 5. The ferrite structure was white in colour, rough in shape, and typically with a maximum carbon content of 0.025% at 723°C. The structure is soft, ductile, and has a body centre cubic (BCC) crystal structure.

<table>
<thead>
<tr>
<th>No.</th>
<th>Specimen</th>
<th>Hardness (HRC)</th>
<th>Average of Hardness</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Raw Material</td>
<td>20 19.5 20.5</td>
<td>20.00</td>
</tr>
<tr>
<td>2</td>
<td>Hardening 1050°C</td>
<td>49 49 49</td>
<td>49.00</td>
</tr>
<tr>
<td>3</td>
<td>Hardening 1050°C, Temp 200°C</td>
<td>47 48 45</td>
<td>46.67</td>
</tr>
<tr>
<td>4</td>
<td>Hardening 1050°C, Temp 400°C</td>
<td>44 43 44</td>
<td>43.67</td>
</tr>
<tr>
<td>5</td>
<td>Hardening 1050°C, Temp 600°C</td>
<td>41 44 42</td>
<td>42.33</td>
</tr>
</tbody>
</table>

In the process, the ferrite and pearlite phases are transformed into the martensite phase. In the hardening process above the austenitising temperature, the specimen’s carbide also increases, causing the physical and mechanical properties to shift; the martensite phase enlarged, and the hardness increased. In the tempering process at 200°C, the hardness is 46.67. There is a slight decrease in the hardness compared to that of the 1,050°C hardened material. It occurs due to the low temperature of tempering, allowing the phase transformation to start towards the martensite finish up to the tempered martensite phase. The most insufficient average hardness of all tempered materials is 42.33 HRC, which is possessed by the test material with a temperature of 600°C. It is caused by the fact that the austenite phase-shifted into the pearlite phase at this temperature, and the cooling reached the slowest, allowing for coarser and bigger metal grains. However, the hardness remains higher than that of the raw material. The higher the material’s tempering temperature, the lower the hardness, as stated by Hariningsih [3]. As illustrated in Figure 10, namely regarding the particle area and particle diameter, where the smaller the values, the higher the material’s hardness. The decreased hardness is linked to transforming the martensite phase into pearlite and ferrite phases, resulting from a slow cooling rate [26].

From the average hardness s of the tempered materials examines with varying variables, the permitted hardness for medical devices with the ISO 13485 standard stands at 43±3 HRC [2]. The upper limit is 46 HRC, and the lower limit is 40 HRC. Therefore, this research’s best value is 43.67 HRC, obtained at a variable temperature of 400°C. This figure is close to the acceptable standard of hardness.

4. CONCLUSION

An extraction forceps made of DIN 4021 stainless steel has been investigated to evaluate the effects of tempering temperature on microstructure and hardness. The hardness of that medical device materials increases from 20 to 49 HRC by the hardening process. The transformation of the pearlite and ferrite phases into the martensite phase triggers their hardness and ductility. In the tempering process, the permitted hardness for medical devices with the ISO 13485 standard stands at 43 ± 3 HRC, with the upper limit being 46 HRC, and the lower limit being 43 HRC. The microstructure of stainless-steel DIN 4021 is dominated by ferrite and pearlite structures. It adheres that the raw material is composed of a carbon composition of 0.16 to 0.25%. The martensite structure reduces by 30%, which cause a slight decrease of the material hardness. The largest average area and diameter of the tempered specimens examined are 0.527 μm² and the diameter of 0.587 μm respectively, obtained at a tempering temperature of 600°C. It means that the higher the tempering temperature, the greater the particle dimension and the lower the hardness.

5. ACKNOWLEDGEMENT

The authors would like to thank the Material Laboratory of the Faculty of Engineering, Universitas Wahid Hasyim Semarang Indonesia, for this research’s full support and facilities.

REFERENCES


https://doi.org/10.1007/s11665-017-2605-x.


https://doi.org/10.1016/jijpvp.2015.07.001.

https://doi.org/10.1016/j.jmrt.2019.06.028.


https://doi.org/10.1007/s11041-017-0081-5.

https://doi.org/10.1016/j.matdes.2013.08.099.

https://doi.org/10.1016/j.msea.2011.10.056.

https://doi.org/10.1016/j.matpr.2020.05.396.

https://doi.org/10.1016/j.compchemeng.2012.05.014.

https://doi.org/10.1111/j.1365-3091.2006.00786.x.

https://doi.org/10.1016/j.measurement.2019.03.046.

https://doi.org/10.1080/01913120701719189.