EVALUATION OF PRECIPITATION HARDENING HEAT TREATMENT OF PH 17-7 STAINLESS STEEL SPRING

A. A. Babakoohi Ashrafii, H. Mohammadii and A. Habibolla Zade1

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1 Metallurgical - Material Engineering Department of Semnan university, Semnan, Iran.
2 Mining and Metallurgical Engineering Department of Amirkabir University, Tehran, Iran.

Abstract: In this paper, the influence of heat treatment on PH17-7 stainless steel spring was evaluated. Precipitation hardening phenomenon of PH 17-7 steel was evaluated in three stages. First, the spring constant changes with time and temperature was evaluated. Second, the spring constant changes with respect to its original length at constant temperature and time with blocking (spring length after compression, 18 and 21 mm) were investigated. And finally, the spring heat treatment at 480°C for 80 min and then holding at 230°C in oil bath for 60 min without blocking were investigated. The results showed that the use of 18 mm block have large spring constant than 21 mm block. The optimal conditions (480°C for 80 min) for this spring to reaching maximum spring constant were determined.

Keywords: “PH 17-7 stainless steel, precipitation hardening, spring constant, blocking”.

1. INTRODUCTION

Steel 17-7 PH, WNo: 1.4568, is a chromium-nickel-aluminum precipitation hardening stainless steel that classified as high-strength austenite-martensitic stainless steel [1, 2]. Precipitation-hardening (PH) stainless steels are a family of stainless steels that combine high strength, ductility, and good corrosion resistance. This material is widely used as a structural material in a variety of applications in aircraft and spacecraft, and surgical instruments, beds, springs [3,4]. Depending on the thermal and mechanical history, alloys can contain variable amounts of austenite, ferrite, and martensite. The alloy 17-7 PH has approximately the same chromium and nickel contents as austenitic type 301 stainless but also contains 1.2 wt% Al for precipitation hardening. Precipitation hardening in these steels is achieved by homogeneously dispersed fine precipitates formed during aging [5].

The main reason for the development of precipitation-hardening stainless 17-7 PH steel, is the fact that Maraging steels, despite of very high degree of alloying, have not satisfactory corrosion resistance. Therefore, PH steels have been developed. These steels have high values of strength and fracture toughness, very close to Maraging steel, but with greater resistance to corrosion[6].

Strengthening of the steel is achieved by the transformation of austenite into martensite and subsequent precipitation of intermetallic compounds.

The Al-Ni compositions and carbides will precipitate in grain boundary from solid solution phase during hardening heat treatment [7].

After partial or complete transformation austenite into matrensate these steels are additionally strengthened with precipitation aging. This strengthening with precipitation of intermetallic phase NiAl is very simple and effective [8].

Precipitation hardening in these steels is achieved by homogeneously dispersed fine precipitates formed during aging.

PH stainless steels containing Ni and Al can be age-hardened by the formation of NiAl particles. In the marensitic condition, the aluminum in these steels is in supersaturated solid solution. Upon heating for precipitation hardening, the aluminum in the martensite is precipitated as Ni-Al (NiAl and Ni2Al3) intermetallic compounds, as had been identified by electron diffraction studies. The amount of precipitation depends on the time and temperature at which the reaction takes place [9, 10].

The carbon in this steel forms some carbide
such as $\text{M}_2\text{C}_6$, $\text{M}_3\text{C}_6$, $\text{M}_7\text{C}_3$. The important carbide is $\text{M}_2\text{C}_6$. When this steel is heated to 1100-1150 °C, the carbides dissolve and will obtain the austenitic phase free of carbide precipitation after quenching. Heating the steel in 480°C–500°C temperature range, $\text{M}_2\text{C}_6$ carbide will precipitate in the grain boundary [11].

Typically the 17-7 PH alloy is furnished in the annealed condition. In this condition, the material possesses an austenitic structure. As an austenitic material, the 17-7 precipitation hardening alloy possesses a relatively low strength. This is the condition in which formability is easiest.

The 17-7 PH Alloy is covered by the AMS 5529, ASTM A313 and EN 10088-3 specification and standards. When 17-7 PH alloy with an austenitic structure is cold worked by substantial deformation, a transformation to martensitic structure results from the deformation. In this condition, the material may be precipitation hardened directly by heat treatment at 482°C. The aim of this paper is to study the influence of effective factors such as temperature and time, blocking and without blocking precipitation hardening on 17-7 PH steel.

2. MATERIALS AND METHOD

The chemical composition and dimension of the springs used in the present work is shown are Table 1 and 2 [12, 13].

Precipitation hardening of 1.4568 (17-7 PH) steel for obtaining the spring with K (Spring Constant) = 0.19 -0.21 in four stages was performed. The spring diameter in this study is low, so precipitation hardening heat treatment at high temperature (955°C) is prevented, because the high temperature cause to spring deformation and distortion. The melting point of this material is about 1430 °C.

<table>
<thead>
<tr>
<th>Description</th>
<th>Attribute (mm)</th>
<th>Description</th>
<th>Attribute (mm)</th>
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</thead>
<tbody>
<tr>
<td>Outside diameter of the spring</td>
<td>4.65</td>
<td>Primary length</td>
<td>36.5</td>
</tr>
<tr>
<td>The external diameter of the wire spring</td>
<td>0.65</td>
<td>Block Length</td>
<td>18-21</td>
</tr>
<tr>
<td>The number of ring diameter coil springs</td>
<td>18</td>
<td>Diameter Average spring</td>
<td>4</td>
</tr>
<tr>
<td>The number of useful ring diameter coil springs</td>
<td>16</td>
<td>Inside diameter coil springs</td>
<td>3.35</td>
</tr>
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</table>

Therefore, firstly should obtain an appropriate operating temperature range for precipitation hardening and strengthening. So the precipitation-hardening treatment consisted of heating at 400 °C to 540 °C in a resistance furnace for a period of 35 to 150 min was selected with regarding the condition CH 900 (AMS 5529). The low temperature allows minimum distortion compared to conventional hardening processes.

The samples were arranged inside the reticulated metal basket and then were placed into the furnace with 10°C accuracy and were cooled in air after heat treatment. The final step of this heat treatment cycle is age-hardening, which the specimens were held at 230°C in oil bath for one hour and then cooling, normally very slowly, as shown in Fig.1. The flow chart of the process is illustrated in Fig. 4-6.

In the second stage, the changes of spring constant (kg/mm) were studied by blocking and pressing the spring from initial length to

![Fig. 1. Heat Treatment Cycle for Spring](image-url)
secondary length or block length (18 and 21 mm). The springs were placed into the fixture to block the springs (as shown in Fig. 2) after after precipitation-hardening heat treatment (480°C for 45, 60, 70, 80 min), then were placed into oil bath at 230 °C for 1hr. The main reason of using this type of fixture is to prevent from deformation of the springs during blocking. The spring constant were measured with measurement hardness device in accordance to Fig(3). The spring constant is obtained by using this device by measuring the force changes (ΔF) in terms of distance.

In the third stage, the spring constant changes in terms of length were evaluated at constant time and temperature without blocking. And finally the samples that were heat treated at 480°C for 80 min and were aged at the 230°C temperature for 1h in oil bath without blocking (this state is the optimum condition for spring production) were prepared for additional analysis.

The general microstructure of the material was observed by an optical microscope and SEM (Scanning Electron microscopy). Metallographic samples from selected heat treatment conditions were prepared and etched in chemical etching (the etchant consists of nitric acid, acetic acid and glycerin) to reveal the microstructures [4, 11]. The analysis of the base metal characteristic were obtained with the EDX and XRD.

3. RESULTS AND DISCUSSION

3. 1. Evaluation of Spring Constant Changes With Time and Temperature

As shown in Fig (5), the spring constant changes at 400 °C- 540 °C temperatures from 35 to 170 min shows a peak at each of temperature. With prolonging holding time at these temperature, the precipitation of inter-metallic compounds of Ni-Al (Ni₃Al and NiAl) and carbides (Cr₃C₂) are increased, which were precipitated in grain boundaries from
supersaturated solid solution phase, so the hardness and strength of spring with the conventional precipitation strengthening mechanism are increased[12]. The amount of precipitation depend on the time and temperature.

The Ni₃Al phase known as γ’ phase, has ordered cubic, and phase NiAl has a cubic crystal lattice. Intermetallic γ’ phase precipitate as coherent precipitate [13]. Coherence shows high stability of precipitates, that means very low surface energy per unit area of the precipitate/matrix. Precipitation hardening is achieved by the precipitation of coherent second phase from a supersaturated solution created by an aging heat treatment. The intermetallic phases formed impede the dislocation movement and therefore strengthening the alloy [1,15].

The mechanism of strengthening by precipitation hardening involves the formation of coherent clusters of solute atoms (that is, the solute atoms have collected into a cluster but still have the same crystal structure as the solvent phase). This causes high amount of strain because of mismatch in size between the solvent and solute atoms. The cluster stabilizes dislocations, because dislocations tend to reduce the strain. When dislocations are trapped by coherent solute clusters, the alloy is considerably strengthened and hardened. However, if the precipitates are semi-coherent, incoherent or are incapable of reducing strain behavior, a dislocation can circumvent the particles only by bowing into a roughly semi circular shape between them under the action of an applied shear stress. Consequently, the presence of the precipitate particles, provide higher strength by obstructing and retarding the movement of dislocations. The characteristic that determines whether a precipitate phase is coherent or non-coherent is the closeness of match or degree of disregistry between the atomic spacing’s on the lattice of the matrix and on that of the precipitate[16].

Figure (5) shows that spring constant is reached to a maximum value by heating for 80 min at 480°C temperature and the maximum constant spring is low in below and above 480°C temperature.

In the temperature range between 400°C-480°C, the amount of Ni₃Al, NiAl and M₂₃C₆ precipitation in grain boundaries increases gradually, because the above mentioned precipitation particles are stable in this range[17,18,19].

With increasing temperature from 400°C to 480°C, the amount of precipitation particles at grain boundaries and matrix phases as barriers against of dislocations movement increases and thus the hardness and strength of springs increases.

With increasing temperature from 480°C to 540°C is facilitated the conditions for diffusion and aggregation of the precipitation particles in matrix phase and result in non-uniform distribution of the precipitation particles.

This non-uniform distribution of precipitation particles in matrix phase and also their aggregation in special places, is caused to removal obstacles against dislocations movement and therefore some dislocations are pile-uped in back of the obstacle, will release and continue to their movement. So the hardness and strength of spring reduced and therefore the spring constant decreased (Fig. 4). Therefore, a gradually increase of spring strength and constant with increasing temperature to 480°C, while a gradually decrease of spring strength and constant with increasing temperature to 520°C is showed in Fig. 5.

According to Fig (5) the maximum spring constant for temperatures 460, 470 and 490 °C are seen in time 75, 90, 100 min, respectively. But the height of these peaks are lower than the 480°C temperature and 80 min. The time to achieve the maximum spring constant is decreased by increasing the temperature of heat treatment from 460 to 490°C.

The results show that the strength and spring constant distribution occurs at range of time and temperature, but any peak is not seen in the temperature ranges lower than 460°C or higher than 490°C.

The changes of spring constant with primary length are shown in Figs (6) and (7). The spring are heated at 480°C temperature and are blocked to amount of 18-21 mm and then heated at 230°C temperature for 1 h in oil bath. Three major factors are determined the strength and spring constant are stress (applied force), temperature and
heating time.

Figures (6) and (7) shows that the spring constant changes with heat treatment at 480°C at times of 45 to 80 min for 18 and 21 mm spring. As shown, the spring constant for 18 mm spring length after blocking more than 21 mm, so, the amount of applied stress is the effective factor in strengthening of the spring. For austenitic stainless steels, as the cold work increases, the optimum aging temperature decreases [16].

With increasing heating time to 80 min at 480°C, the spring constant increases for both of blocking springs, but the amount of this effect is high in the 18 mm length of spring after blocking. As expected, increasing the amount of cold work will cause the final spring constant to increase, as seen in Fig (8). For 17-7 PH, as the cold work increases, the optimum aging temperature decreases. According to Fig (8), the spring with 18 mm block shows more spring constant at low temperatures and reached the optimum at 480°C temperature for 80 min. As the heating time is increased, the spring constant increase to a maximum, and then drop off.

The decrease in spring constant signifies a condition that is called “over-aging”. Over-aging occurs as the particles that caused the increase in strength continue to grow in size. As these particles grow, they begin to coarsen and cause a decrease in the spring constant.

3. 2. Evaluation of the Spring Constant Changes With Time at Constant Temperature

The results of spring constant changes wit time at constant temperature without blocking are shown in Figs (9) and (10). In Fig (9) is observed that spring constant will gradually rise with increasing heat treatment time at 480°C temperature from 45 to 90 min and increasing heat treatment time from 90 to 150 min is decreased again. In Fig (10), at 520°C temperature, the spring constant was increased from 45 to 80 min and then was faced with severe downfall from 80 to 150 min. The diagrams of spring constant changes with time at 480°C and 520°C temperatures, indicate that the maximum spring constant is attainable about 80 min at 480°C temperature and is about 70 min at 520°C
temperature, respectively (Fig.11). With increasing precipitation hardening time and temperature the amount of precipitations in grain boundaries and also inside grains increases. With increasing heat treatment temperature, the required time to reaching maximum spring constant is decreased.

3.3. Evaluation Microstructure PH17-7 Spring

The microstructure images were obtained on transverse and longitudinal direction after heat treatment of 17-7PH steel at 480°C for 80 min and aging at 230°C for 60 min in oil bath without blocking (Fig. 12). In these microstructures, the elongated grains in wire forming direction is observed. Due to the cold forming process on this spring, the amount of internal stresses are high. Whatever, the elongated intermetallics were participated inside matrix phase as shown in the cross-section.

The analysis and characteristic of the base metal and components were obtained with EDX and XRD (Fig.13). SEM images of microstructure in Figs (14a,b) show typical compounds in matrix phase such as A and B Areas. The presence of metallic elements such as Ni, Al, Cr and Fe was confirmed using EDX and XRD in the A and B Areas (Figs. 15 and 16).

These Areas are created by precipitation of Ni-Al inter-metallics (Ni₃Al and NiAl) and carbides such as Cr₂₃C₆ in matrix phase. The precipitated particle was elongated in forming direction in local Areas so the crystal lattice was more

Fig. 9. Diagram depending on changes of spring constant in terms of primary length without blocking at 480°C and times (a) 70 (b) 80 min.

Fig. 10. Diagram depending on changes of spring constant in terms of primary length without blocking at 520°C and times (a) 70 (b) 80 min.

Fig. 11. Diagram depending on changes of spring constant in terms of heat treatment time without blocking (a) 480°C (b) 520°C, respectively.

Fig. 12. Image of drawn austenite texture in directions (a) cross section (b) longitudinal section of spring steel 1.4568
irregular and these factors create a lot of big and large obstacles in front of dislocations movement. The results showed that with prolonging time of heat treatment at 480°C to 80 min, the number of these Areas with inter-metallic compounds and carbids (Ni₃Al, NiAl and C₂₃C₆) increases [20]. From 80 to 150 min, these Areas and mentioned compounds decreases which adversely effects on strength and spring constant of samples.

4. CONCLUSIONS

1. The results showed that heating temperature and time and stress factors (applied force) playing a major role in the spring constant.
2. The maximum of spring constant is obtained at 480°C temperature for 80 min for PH17-7 springs.
3. The use of 18mm block shows more spring
constant than 21mm block.
4. The XRD and EDX analysis confirm the presence of metallic elements such as Ni, Al, Cr and Fe that these elements precipitate in matrix phase as Ni3Al and NiAl and Cr23C6 inter-metallic combinations.
5. The results showed that with prolonging time of heat treatment at 480°C temperature to 80 min, the number of Areas with intermetallic combinations increases and with prolonging time from 80 to 150 min these Areas decreases.

REFERENCES

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