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atadorno@iq.unesp.br

Universidade Estadual Paulista Júlio de
Mesquita Filho
Brasil

Gárlipp, Waldir
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Thermomechanical treatment of a Fe-Mn-Al alloy

Waldir Gárlipp¹

Resumo: Análises da dureza e da estrutura metalográfica mostraram em uma liga de Fe-Mn-Al a existência de fase γ e de precipitado β pela ativação do Si, durante o rápido esfriamento do material fundido. Durante o tratamento térmico nas temperaturas de recuperação e de recristalização são formadas franjas, grãos maiores do precipitado β e da fase recristalizada γ . O tratamento termomecânico dá ao material excelentes propriedades de resistência mecânica e de ductilidade, sem a colaboração do precipitado β .

Palavras-chave: liga, tratamento termomecânico, metalografia, microscopia óptica

Abstract: Analysis of the hardness and the metallographic structure, show in a FeMnAl alloy the existence of a γ phase and a β precipitate, activated by Si during the casting processes. During the recovery and recrystallisation temperatures there are the formation of fringes, coarsened grains of γ phase and β precipitate. The thermomechanical treatments give to the material excellent properties of mechanical resistance and good ductility, without the β collaboration.

Keywords: alloy, thermomechanical treatment, metallography and optical microscopy

¹ Prof. Titular Aposentado do Departamento de Materiais – USP - São Carlos.

INTRODUCTION

Studies of ordered alloys Fe₃Al and Fe₃Si [6,12] has enabled the development of the Fe-(5w%Al)-(1.5w%Si) magnetic alloy itself, with excellent resistance to corrosion and high resistance to oxidation at high temperatures, by forming a thin surface layer of alumina, Al₂O₃.

The composition limit for the alloy is 1.5w%Si and 5w%Al [1,17], which allows their use as well as resistors (80 $\mu\Omega$ cm), combined with moderate mechanical strength properties and resistance to oxidation.

Their mechanical fragility is attributed to the structure BCC and to the interstitial defects of the alloy.

Because their FCC structure the (55w%Fe-35w%Mn-10w%Al) alloy [1,16], presents at high temperatures a better mechanical and a good plasticity properties.

A combination of (35w%-40w%Mn) and 15w%Al of that alloy produces an improper structure

β -Mn, and a lamellar ferritic structure B2 that do not maintain their austenitic structure at high temperature [2,13,17].

The addition of Carbon to the alloy [8] stabilizes the austenite, gives inoxidizable property up to 650°C, similar to the inox steel Fe-Ni-Cr, and superior mechanical conditions up to 815°C [7,10] to the carbon steels 304 AISI and 347 AISI.

During cold rolling operation that alloy hardens quickly wanting intermediate thermal anneals.

Today a nominal composition proposal to agree mechanical properties and corrosion resistance for the alloy [1,2,3], is 53w%Fe, 35w%Mn, (8 - 9w%)Al, 2w%Si, (0.7-1.0w%)C.

Addition of 3w% of elements as B, Cr, Mo, Ni, Nb, Ti, do not improve the best properties of this alloy [3,14,15,18]. *corresponding

The table 1 shows for example some mechanical properties of different Fe cast-alloys,

Table 1 (weight%)

Fe-Cast-Alloy	σ_R (MPa)	σ_E (MPa)	Elong. (%)	Hv	Refer.
30Mn-10Al-1.3Si	605	-----	10	381Vickers	9
30Mn-7Al-1.25Si	550	-----	11	166Vickers	9
30Mn-8Al-1C*	890	495	54	-----	9
31.22Mn -7.54Al-1.34Si-0.93C	738	401	50	234Vickers	10
30Mn-8Al-1.5Si-1C	800	574	72	-----	3

Some properties of the austenitic alloy (56w%Fe. 30w%Mn, 10w%Al, 3w%Si, 1.0w%C) [4] are presented here:

remarkable corrosion resistance to boiling sea water during 118 hours [9], what is adequate to use as helix for vessels long journey;

imperceptible superficial corrosion after to be into sea water during 32 days at room temperature, as show similar works [1,5,9,10];

their properties σ_R (MPa), σ_E (MPa), between 24° C and 815° C, and of cold reduction increment, are higher than that austenitic Fe-Ni-Cr steels [10];

their ductility is similar to traditional inox steels, which permits various percentages of reduction;

the σ_E of the alloy can reach to high values of 1660(MPa) for 80% of cold reduction [10];

studies of that Fe-Mn-Al-Si-C austenitic steel, at three dynamic strain aging rates in the range of 172-345°C, show the occurrence of peaks in strength, in ductility and work hardening parameters, the three well-defined peaks for the work hardening were displaced to higher temperatures with increase in strain rate, and the activation energy involved was determined and the possible mechanism of strain ageing was proposed [4];

the oxidization resistance of the alloy at 700° C for 70 hours is almost the same for the inox AISI 304 at 800° C for 40 hours with an stable oxidized superficial layer [11];

their electrical resistivity varies from 168 $\mu\Omega\text{cm}$ to 180 $\mu\Omega\text{cm}$ between 80°C and 800° C, values that are superior on the Ni-Cr and Kanthal alloys [11].

MATERIAL & METHOD

The following alloy was cast using purest components,

**Fe(58.8w%)-Mn(30.2w%)-Al(8.6w%)-
Si(1.30w%)-C(0.63w%)-Cr(0.40w%)-Ni(0.05w%)**

Chemical analysis of the cast-alloy indicated a composition close to the nominal one, with Cr, Ni as main impurities.

The existence of two metallographic structures β and γ , was revealed by the optical metallography and the Scanning Electronic Microscope (SEM-Instituto de Quimica-UNESP), having the β phase a little concentration of Cr and a smallest concentration of Fe and Mn than in the γ phase.

The existence of β phase it is attributed to the Si influence on the casting procedure.

These two phases stay inside the $(\gamma+\beta)$ band of the Horizontal Section at 7600C of the Fe-Mn-Al Ternary Equilibrium Diagram [2].

ANALYTICAL PROGRAMS

Two programs were settled, trough micro hardness and micrographic analysis, to determine the temperatures of stress relief, and of re-crystallization, as well as to evaluate the mechanical properties for technological applications.

1°) Program – Sequence of thermomechanical treatments;

a) Cold rolling to 40% of reduction (0.2 cm thickness), b) Anneal at 1473K, 1 hour and quench to normal temperature, c) Anneal from 300K to 1200K, in successive steps of 100K for 1 hour to measure the respective micro-hardness, figure 1, and metallographic structure observation, Table 2.

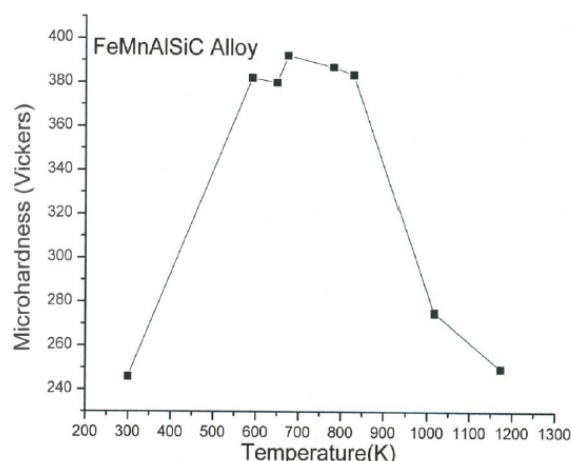


Figure 1 - (1st Program)

Table 2 - Micrographic observation of annealed samples

Temperature (K)	GamaPhase(γ)	Beta Phase (β)	Observation	Micrography (Figure)
300	Coarse grains with L of 44 μm to 800 μm	Few grains bordering γ phase Lm=30 μm	Fine fringes drawn on all the optical field Lengths L and medium Lm	III
640	Larger granulation than at the 300K Lm=44 μm	More grains bordering γ phase	Idem	IV
660	Some sub-grains Lm=150 μm	Spreaded and bordering γ Lm=60 μm	Long size due on the pre-cold rolling	V (3)
800	Decomposition of grains and creation of sub-grains	Idem	Idem	VI (4)

1000	Decomposition of more grains Lm \approx 150 μ m	Long and cracked in various parts	Decomposition of fringes into spheres and small batons	VII
1180	Scattered decomposition Lm \approx 15 μ m	Scattered decomposition Lm \approx 15 μ m	_____	VIII (5)

Figures 3, 4, 5 respectively micrographics of V,VI,VIII from table 2

2º) Program – Sequence of thermomechanical treatments;

a) Cold rolling to 40% of reduction (0.2 cm thickness), b) Anneal at 1473K, 1 hour and quench to normal temperature, c) Another cold rolling to 40% of reduction (0,2 cm of thickness), d) Anneal from 300K to 1200K, in successive steps of 100K for 1 hour to measure the respective micro hardness, figure 2, and metallographic structure observation, Table 3.

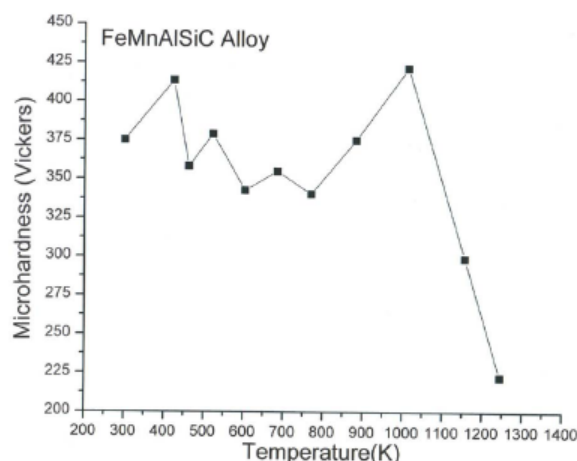


Figure 2 - (2nd Program)

Table 3 - Micrographic observation of annealed samples

Temperature(K)	Gama Phase (γ)	Beta Phase (β)	Observation	Micrography (Figure)
423	Grain boundary not observed	Lengthened by initial cold rolling Lm =70 μ m	Fringes at some regions. Lengths L and Medium Lm	_____
461	Idem	Idem	Fringes and initial cold rolling with directions of 90º	_____
521	Small numbers of grains with Lm=500 μ m Dm=150 μ m	Some grains bordering the γ phase	Existence of depressed γ phase with internal fringes Lüders bands?	XI
683	Large grains	Idem	Nitid fringes or Lüders bands!	XII
768	The number of grains is greater than that at 683K	Idem	In some places an high density of fringes	XIII-XIV

880	Poly crystals Lm=800µm	Interstitials grains on the γ phase borderline	Elongated black fringes filaments	XV
1010	More Poly- crystals Lm = 400 µm - 450 µm	Interstitials cracked grains on the γ phase borderline Small Lm	Black filaments inside the γ phase and Serrated traces	XVI
1155	Decomposition of grains with Lm<5 µm to 15 µm in distints regions	Cracked grains	Ausence of fringes	XVII (6)
1245	Polyhedral crystals Lm \approx 12 µm and twins bands	Decayed with Lm \approx 60 µm	Idem	XVIII

CONCLUSION

Under the metallographic point of view

The component Si during the cooling speed of the molten state alloy produces a blend of γ phase and β precipitates [2]. The calculated grain proportion of γ and β was $\gamma/\beta = 522$.

Micrographic photos, of (1°) and (2°) programs, show that the heat treatments produce, coarse grains of γ and β phases, fringes during the stress relief, figures 3, 4, 5 and re-crystallized γ , figures 6, 7.

By collaboration of the fringes the re-crystallization of γ phase grains starts at temperature of 800K (1°) program) and at 1000K (2°) program) with dimensions less than 12 µm.

The β phase starts to be multi-fractured at 1000K and no collaborates on the hardness of the material.

Under the technological point of view

1°) Program - In the range of 600K- 800K the heat treatment ensures to the alloy an excellent strength, hardness of 385Vickers, with a adequate grain sizes, and at 1180K, an hardness of 250 Vickers, a favourable mechanical conditions to produce by cold rolling process, plates blades, wires, nuts or bolts.

2°) Program - Internal stress relieves is proven in the heat treatment (range 600K - 700K), because the low hardness, 337Vickers.

The alloy hardens from 800K (337Vickers) to 1000K (422Vickers), and softens to 1250K (220Vickers), which recommends to be good for cold metal-forming.

For technical applications the higher hardness recommended is 422Vickers, despite of more energy consumption and less resistance to oxidation, figure 4

General hardness values, here marked, agree with the data contained in the works cited on the bibliography.

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