Effect of Heat Treatments and SiC Content in the Mechanical Properties and Microstructure of Self-Lubricating Steels

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The objective of the present work was to study the effect of heat treatments in the microstructure, graphite nodules and mechanical properties of self-lubricating steels, to achieve this, self-lubricating steels (Fe + 0.45C + 4Ni + 1Mo % wt ) and additions of 2 and 3 % wt SiC were fabricated. They were consequently heat treated in 3 different conditions: martempering at 180 °C and tempering at 530 °C and 300 °C respectively and austempering at 300 °C. Hardness, yield strength, tensile strength and work hardening behavior were studied in as-sintered and heat-treated samples. The microstructure was analyzed by optical microscopy, scanning electron microscopy, Raman spectroscopy and micro-hardness. The transformation temperatures were determined using dilatometric tests. Results show that the presence of dissolved Si in the matrix due to SiC dissociation notably affects the morphology of the microstructure and transformation temperatures also affecting post heat treatment mechanical properties. The structure of graphite nodules produced by SiC dissociation is not significantly affected by the heat treatments.

Keywords: heat treatment, metal injection molding, plasma sintering, self-lubricating steel.

1. Introduction

In engineering a recurrent topic for research is the development of materials with a good combination of mechanical properties, heat resistance, high wear resistance, high corrosion resistance and low specific weight. When such materials are put into contact and relative motion, a tribological component adds up as minimizing energetic and economical losses due to deficient lubrication becomes an important issue for both companies and researchers. For applications were fluid lubrication is unsuitable (extreme temperatures, near vacuum pressures, contamination-free environments, etc) solid lubricants and self-lubricating materials appears to be an appropriate alternative or even the only alternative to operate under such conditions.

Currently several different methods have been developed for the application of solid lubrication, for example, it can be provided by deposited DLC layers 2,3 , BCN 4 , polymer based composite layers 5 , etc. Solid lubrication can also be provided by incorporating solid lubricant into the bulk material by powder pressing 6,7 or Powder Injection Molding (PIM) 8. In this context the authors of this paper have developed self-lubricating sintered steels produced by Metal Injection Molding (MIM) 9 and studied its mechanical properties and tribological behavior 10,11. Through MIM it is possible to obtain a fine microstructure consisting of a steel matrix with homogeneously distributed graphite nodules. The graphite nodules come from the addition of silicon carbide (SiC) which dissociates into the ferrous matrix leaving rings of stabilized ferrite surrounding a graphite nodule. This SiC dissociation and graphite formation provides a stock of solid lubricant which forms a fine graphite rich tribo-layer when the material is put into contact and relative motion with another surface, thus providing lubrication 12. This graphite has been found to be turbostratic 13: turbostratic graphite consist of disordered graphene layers with random parallel stacking without three dimensional order 14. Turbostratic graphite provides lower friction coefficients than crystalline graphite as found by Kumar et al. 15 this explains the low friction coefficients and wear rates found by previous studies 11,16 therefore, studying the structural changes in these nodules became relevant to evaluate the feasibility to perform heat treatments and retain or improve the tribological behavior of these materials.

Besides contributing to the formation and degradation of the tribolayer, the microstructure of the metallic surface has a large influence on the tribological behavior of any material as wear resistance depends on the appropriate combination of strength, ductility and fracture toughness 17.
However, the effect of the ferritic Si enriched zones in the hardenability, transformation temperatures and mechanical properties have not been studied yet, thus the objective of this work is to study the influence of the heat treatments on the microstructural evolution and mechanical properties of these materials and to determine if such heat treatments have an impact in the structure of the graphite nodules that are part of the microstructure.

To conduct the study sintered self-lubricating steels with a Fe + 0.45C + 4Ni + 1Mo steel matrix and additions of 2 and 3% wt. SiC were heat treated and characterized. The objective of the study was to analyze the influence of the heat treatments and SiC content in the phase transformations, microstructure, graphite nodules, mechanical properties and work hardening of these materials. Work hardening was considered of interest as a high work hardening coefficient have been reported to positively influence the wear resistance under dry sliding conditions in general\textsuperscript{17} and, in particular, of ferrous alloys containing graphite nodules\textsuperscript{18} while decreasing friction itself\textsuperscript{19}. Graphite nodules structure is also of special interest in regards of the wear resistance and friction coefficient of these materials\textsuperscript{20} so Raman spectra is used as it has proven to be an excellent tool to analyze carbon based materials in terms of crystallinity, bonding and defects\textsuperscript{21-23} being particularly useful to analyze the presence of defects in carbide derived carbons\textsuperscript{24}.

### 2. Materials and Methods

Three compositions of sintered steels were used for this study; they consisted of a metallic matrix of Fe + 0.45C + 4Ni + 1Mo (referred as base alloy) and 2 and 3% wt. of SiC. Table 1 details the powders used and their specifications.

<table>
<thead>
<tr>
<th>Element</th>
<th>Commercial name</th>
<th>Particle mean size (µm)</th>
<th>Purity (% wt)</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prealloyed Fe + 0.9C</td>
<td>CL-OM</td>
<td>7.84</td>
<td>98.3</td>
<td>Basf</td>
</tr>
<tr>
<td>(0.45 after sintering)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ni</td>
<td>PF-10F</td>
<td>6.06</td>
<td>99.9</td>
<td>ATMIX</td>
</tr>
<tr>
<td>Mo</td>
<td>OMP</td>
<td>5.50</td>
<td>99.8</td>
<td>HC</td>
</tr>
<tr>
<td>SiC</td>
<td></td>
<td>800</td>
<td>99.0</td>
<td>Cobral</td>
</tr>
</tbody>
</table>

Table 2. Binder composition.

<table>
<thead>
<tr>
<th>Element</th>
<th>Mass % of the mixture</th>
<th>% of the constituent in the binder</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polypropylene</td>
<td>3.40</td>
<td>42.56</td>
</tr>
<tr>
<td>EVA</td>
<td>1.32</td>
<td>16.51</td>
</tr>
<tr>
<td>Paraffin</td>
<td>2.90</td>
<td>36.19</td>
</tr>
<tr>
<td>Cocamide</td>
<td>0.36</td>
<td>0.45</td>
</tr>
<tr>
<td>DEA</td>
<td>0.02</td>
<td>0.20</td>
</tr>
<tr>
<td>Total</td>
<td>8.00</td>
<td>100</td>
</tr>
</tbody>
</table>

Table 3. Powder Injection Molding processing parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Injection Pressure (bar)</td>
<td>1000</td>
</tr>
<tr>
<td>Settlement Pressure (bar)</td>
<td>800</td>
</tr>
<tr>
<td>Injection Temperature (°C)</td>
<td>180</td>
</tr>
</tbody>
</table>
better identification of the carbides in the microstructure. For further microstructural characterization micro-hardness measurements using Vickers scale according to ASTM E386 standard with a 10g load were performed. A differential dilatometer (NETZSCH DIL 402C) was used to analyze phase changes at heating and cooling. For these tests samples taken from 5 different sintered tensile test specimens were used per alloy, with dimensions of 4.8 x 10 mm (diameter x height) the samples were heated at 950 °C for 20 min with a heating rate of 10 °C/min and then cooled at a rate of 10 °C/min up to 100 °C. To obtain the Raman spectra of the graphite nodules the samples 3 test specimens per condition were fractured and 5 spectra were obtained using a Renishaw InVia spectrometer with an Ar laser (λ = 514.5 nm) coupled to a Olympus microscope (BX41 TM). To test mechanical properties 5 samples per condition were tested by uniaxial tension at a strain rate of 0.0067 s⁻¹ according to MPIF 42 standard and also 10 Brinell hardness measurements per sample were done using a 2.5 diameter tungsten sphere according to ISO 6506.

3. Results and discussion

Figure 1 presents the microstructure of the samples of base alloy and base alloy + 3%wt SiC without heat treatment. The base alloy in the as-sintered condition shows zones of fine pearlite and ferritic zones with fine precipitated carbides it can be noticed from the figures that when SiC is added, proeutectoid ferrite surrounding graphite nodules appears. This graphite nodules are formed by carbon coming from SiC dissociation during sintering, the ferrite rings surrounding them is formed because dissolved Si (also coming from SiC dissociation) stabilize α-iron in the zones where SiC was present, this dissolved Si also inhibits the precipitation of carbidies in the microstructure as discussed in references also due to Si presence and the reactor cooling conditions regions of bainite can be found in the microstructure.

All the three alloys had white unetched areas. For the base alloy the etched regions (ferrite + carbides) had a hardness of 2800 ± 264 MPa and the unetched regions had a hardness of 4400 ± 500 MPa, which corresponds to regions of untempered martensite and austenite due to an incomplete dissolution of nickel.

Fig 2 shows a SEM image of a graphite nodule from the as-sintered sample with 3%wt SiC, the figure shows the chemical composition of both the graphite nodule and the area around it which consist in a region with a high amount of carbon surrounded by a metallic phase with a high content of silicon due to SiC dissociation as it is explained by Binder et al.13.

![Figure 1. Optical microscopy of the sintered samples corresponding to: (A) and (B) Base alloy, (C) and (D) Base alloy + 3%wt SiC.](image-url)
Figs 3 (a) and (b) shows dilatometric curves of heating and cooling for the Fe + 0.45C + 4Ni + 1Mo base alloys with three different SiC contents. The analysis shows that the samples with SiC additions undergo a previous transformation prior to ferrite transformation into austenite, this can be attributed to the tempering of martensite and bainite present in the samples in the sintered condition, being this transformation absent in the samples of base alloy without SiC. There is a slight rise of the tempering the tempering temperature from samples with 2% wt SiC to 3% wt SiC, this rise can be attributed to the amount of dissolved Si in the samples as has been previously discussed by Kozeschnik and Bhadeshia. Regarding the austenitic transformation, the SiC content decreases the severity of the volumetric change from ferrite to austenite: As part of the microstructure of the SiC containing alloys is stabilized ferrite hence, it doesn't transform into austenite in the A1-A3 range. This can be noticed by analyzing the slope of the curves in the A1-A3 range, while samples of base alloy exhibit a considerable slope in the austenitic transformation range, samples with SiC additions shows a narrow range with a minor slope, hence, a less severe volumetric change. The same applies for the transformation from γ to α + Fe3C during cooling. Also A1-A3 temperatures are influenced by the Si dissolved in the matrix as a product of the SiC dissociation during sintering: As Si difficult the transformation of ferrite into austenite, a rise in the transformation temperatures A1 and A3 is expected. These temperatures are shown in table 4.

Table 4. Transformation temperatures.

<table>
<thead>
<tr>
<th></th>
<th>Base alloy</th>
<th>Base alloy + 2%_wt SiC</th>
<th>Base alloy + 3%_wt SiC</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1(°C)</td>
<td>660 +/- 6</td>
<td>690 +/- 6</td>
<td>710 +/- 5</td>
</tr>
<tr>
<td>A3(°C)</td>
<td>710 +/- 9</td>
<td>730 +/- 2</td>
<td>750 +/- 8</td>
</tr>
</tbody>
</table>

Silicon dissolved into the matrix due to SiC dissociation also had an impact in the microstructure of the austempered samples. Figure 4 shows the microstructure of austempered samples containing 0, 2 and 3% wt SiC. Samples without SiC shows a regular arrangement of bainite + martensite, however SiC containing samples exhibit larger and coarser bainite grains growing from the ferritic zones surrounding the graphite nodules, this resembles the microstructure found by Malla, Grech and Smallman for high silicon ADI. Figure 4B and 4D shows a FE-SEM image of bainite from the samples without SiC and of the acicular ferrite surrounding a graphite nodule of a sample with 3% wt SiC. As silicon is a ferrite stabilizer it can be expected that acicular ferrite nucleation during austempering starts from the Si rich zones (which in this case is the area surrounding a graphite nodule) and then grow into the retained austenite, this results into a coarser and more unevenly distributed mixture of ferrite-austenite which explains the drop in the mechanical properties of the austempered samples with SiC additions that is shown in figure 5.

Figure 5 shows the mechanical properties obtained by heat treating samples of the base alloy and samples with additions of 2 and 3% wt SiC. For all the alloys, as expected, decreasing the tempering temperature increases the mechanical resistance and, in general, the addition of SiC to the base alloy improves its mechanical properties by increasing the hardness and mechanical strength without a considerable decrease of the ductility. It should be noticed that there are two effects that have to be considered, on one hand the SiC additions add dissolved Si to the matrix that
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Figure 4. Optical (OM) and electronic (SEM) micrographs of the austempered samples corresponding to: (A) OM of base alloy (B) OM of base alloy + 3 %wt SiC (C) SEM of base alloy (D) SEM of base alloy + 3 %wt SiC.

Figure 5. Mechanical properties of as-sintered and heat-treated samples of (A) Base alloy, (B) Base alloy + 2 %wt SiC, (C) Base alloy + 3 %wt SiC.
causes solid solution hardening and an increment in the hardenability\textsuperscript{42}, on the other hand the zones with a high amount of Si remains ferritic which is ductile\textsuperscript{33} this explaining the positive impact of SiC in the hardness and ductility. However, for the austempered samples the trend is reversed: without great differences in ductility the hardness and mechanical resistance decreases with SiC additions. Generally, Si additions into bainite produces an increase in hardness and the other mechanical properties of steels as stated in\textsuperscript{34} and also reported by\textsuperscript{35}. However, as the same austempering treatment were performed in the 3 alloys, disregarding the SiC content, Si dissolution into the matrix caused a deviation on the kinetics of the bainite formation, therefore resulting into a coarser and less homogenous microstructure which is detrimental to the mechanical properties.

To better understand the effect of SiC in the mechanical properties of these alloys the work hardening behavior was studied: several work hardening models have been developed to fit experimental stress - strain data into a mathematical model where one or more work hardening exponents can be extracted\textsuperscript{36}. The Hollomon equation is regarded as the most simple and practical way to obtain this parameter. The Hollomon work hardening exponent \((n)\) is calculated according to equations 1, 2 and 3:

\[
\sigma = K\varepsilon^n \quad (1)
\]

Where:

\[
\text{True Stress } \sigma = S(1 + \varepsilon) \quad (2)
\]

\[
\text{True Strain } \varepsilon = \ln(1 + \varepsilon) \quad (3)
\]

Being \(S\) the engineering stress and \(\varepsilon\) the engineering strain. However a deviation from this behavior had been reported at low and high strains\textsuperscript{37,38}. Nevertheless calculations of instantaneous \(n\) values over true strain has proven to be a good method to analyze the behavior of steels with mixed microstructures allowing to compare the work hardening exponents and plastic deformation obtained\textsuperscript{39}. An instantaneous \(n\) value means that the work hardening coefficient is calculated for each step of plastic deformation, this can be deduced from equation (1) as shown by Zhang et al.\textsuperscript{39}.

\[
n_i = \left(\frac{\varepsilon_i}{\sigma_i}\right) \left(\frac{d\sigma_i}{d\varepsilon_i}\right) \quad (4)
\]

Where \(n_i, \sigma_i\) and \(\varepsilon_i\) are the instantaneous work hardening exponent, the true stress and true strain respectively\textsuperscript{39}. Figure 6 a, b and c shows the instant work hardening exponent \((n)\) for Fe + 0.45C + 4Ni + 1Mo sintered steel with additions of 2 and 3\%\textsubscript{wt} SiC under as-sintered, martempered and tempered at 300°C, martempered and tempered at 530°C and austempered conditions.

Instantaneous work hardening exponent v/s true strain plots shows that steels with additions of SiC exhibits larger plastic strain and work hardening than the base alloy due to the presence of stabilized ferrite and dissolved Si originated from SiC dissociation being this effect previously reported by Cai et al. for ferrite-bainite dual phase steels\textsuperscript{35} and by Zhou et al. for ferrite-martensite dual phase steels\textsuperscript{40}. The as-sintered and austempered samples show larger strains and work hardening coefficients than the martempered samples, SiC additions increase these differences thanks to the effect of Si in the ductility of the alloys as previously discussed.

Raman spectroscopy was used to analyze the influence of the heat treatment in the turbostratic structure of the graphite nodules. Figure 7 shows the Raman spectra of graphite nodules from the base alloy + 3\%\textsubscript{wt} SiC in the as-sintered and heat-treated conditions. The spectra shows the G band at 1580 cm\textsuperscript{-1} which corresponds to a first order mode with \(E_{2g}\) symmetry being typical for graphite materials, the disorder induced bands D and D’ at 1360 and 1620 cm\textsuperscript{-1} respectively which corresponds to double resonance processes which are inhibited in defect-free graphite\textsuperscript{41} and finally the 2D band which is a second order resonance mode which varies with the number of graphene layers and the stacking order of such layers.

Figure 8 shows a 2D band of an as-sintered sample which is adjusted by two Lorentzian peaks, the presence of two peaks within the bands indicates a transition process between a turbostratic structure common to carbide derived carbons (CDCs) to a more crystalline structure which is known to present two well defined peaks within the 2D band\textsuperscript{42}. All the samples show the same peaks, the presence of D and D’ bands indicates the presence of defects which is expected in graphitic materials derived from carbides\textsuperscript{43-45}. The intensity ratio ID/IG has been largely used to measure crystallite size and to compare the quantity of defects in graphitic samples. Figure 9 shows a plot of ID/IG ratio for the samples in the as-sintered and heat-treated conditions.

Statistical analysis of the ID/IG ratios shows that there is no statistically significant difference between the samples as determined by one-way ANOVA (F(3,56) = 1.769, \(p = 0.05\)) which means that even if the means of the ID/IG ratio differs between conditions it cannot be concluded with a 95% confidence that the heat treatments had an impact on the structure of the graphite nodules measured by Raman spectroscopy, therefore, for these materials, heat treatments can be used to improve the mechanical properties of these materials without significantly affecting the turbostratic
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Figure 6. Instantaneous work hardening exponents (n) v/s true strain for: (A) Base alloy, (B) Base alloy + 2%wt SiC and (C) Base alloy + 3%wt SiC.

Figure 7. Raman spectra for samples of base alloy + 3%wt SiC in the as-sintered and heat-treated conditions.
4. Conclusions

Sintered self-lubricating steels were produced by powder injection molding and plasma assisted debinding and sintering. These were heat treated and its mechanical properties and microstructural evolution were studied. From these studies, it was found that:

- SiC addition to the base alloy decreases the severity in terms of volumetric changes of the transformation $\gamma \leftrightarrow \alpha + Fe_3C$.
- Si dissolution into the matrix rises the $A_1$ temperature due to ferrite stabilization, it also modifies the microstructure of the specimens, not only by generating highly stabilized ferrite zones around graphite nodules but also by modifying the morphology of the matrix’s microstructure after heat treatment. This is particularly noticeable in the austempered samples where dissolved Si generates coarser bainite grains which are detrimental for the mechanical properties.
- SiC addition improves mechanical resistance for all the samples with a maximum at 2% wt SiC. This trend is reversed for austempered samples as its microstructure is coarser than the one found in the samples without SiC due to Si dissolved into the matrix. Also SiC addition greatly influences the plastic strain and work hardening exponents due to ferrite stabilization and dissolved Si.
- The presence of the D and D’ band in the Raman spectra of the graphite nodules indicates the presence of defects and disorder in the structure of these nodules, this is further confirmed by the shape of the 2D band which does not corresponds with a perfectly crystalline graphite.
- Heat treatments don’t significantly affect the structure of the graphite nodules generated by SiC dissociation.

5. Acknowledgements

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6. References


