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Synthesis and characterization of the Fe-Zn intermetallic phases using the Rietveld Method

Síntese e caracterização de fases intermetálicas Fe-Zn por intermédio do Método Rietveld

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Abstract

The Rietveld Method was used to investigate the structure of three Fe-Zn intermetallic phases and to quantify their proportion. The zeta, delta and gamma phases were successfully obtained by the sintering process. A simple mixture of the three intermetallic phases was made with a known mass percentage and the XRD pattern of this mixture was collected and analyzed by the Rietveld Method. A difference of 14% between the simulated and the experimental XRD patterns was found, indicating a satisfactory fitting. Slightly different lattice parameters from those given in the ICSD cards were observed. Because of the good fitting, the calculated phase percentages of the mixture agreed closely with the real ones. Thus, it was concluded that the Rietveld Method is a reliable method to quantify the phases in galvanized steel sheets and the same methodology applied in this laboratorial study can be applied to industrial sheets.

Keywords: Fe-Zn intermetallic phases, galvanized steel sheets, Rietveld Method, X-ray diffraction.

Resumo

O Método Rietveld foi utilizado para investigar a estrutura de três fases intermetálicas Fe-Zn e quantificar sua proporção. As fases zeta, delta e gama foram obtidas com sucesso pelo processo de sinterização. Uma mistura simples das três fases intermetálicas foi realizada com uma porcentagem de massas conhecida e seu padrão DRX foi coletado e analisado pelo Método Rietveld. Uma diferença de 14% entre os padrões DRX simulado e experimental foi encontrada, indicando um ajuste satisfatório. Uma pequena diferença entre os valores dos parâmetros de rede calculados e aqueles contidos nos cartões ICSD foi observada. Por conta do bom ajuste, as porcentagens de fase da mistura calculadas concordaram de maneira próxima com as porcentagens experimentais. Logo, concluiu-se que o Método Rietveld é um método confiável para quantificar as fases presentes em aços galvanizados e que a mesma metodologia aplicada nesse estudo laboratorial pode ser aplicada em chapas industriais.

Palavras chave: Fases intermetálicas Fe-Zn, chapas de aço galvanizado, difração de raios X, Método Rietveld.

1. Introduction

The combination of corrosion resistance, surface quality and workability is one of the biggest challenges in the

automotive industry. In the last decades, several improvements led to the development of techniques that increased the

surface properties of steels applied in this field. Among these advances is the usage of galvanized steel sheets, such

as dual phase or interstitial free steels, in the production of vehicle components. In the galvannealing process, steel sheets are subjected to a zinc bath followed by a heat treatment that converts the entire coating to a zinc-iron alloy (Kao *et al.*, 2009). Thus, the main difference of this process when compared with a typical galvanized coating is the formation of intermetallic compounds between the deposited zinc layer and the iron from the substrate (McDevitt *et al.*, 1997; Morimoto *et al.*, 1997; Kato *et al.*, 2000). The excellent weldability observed in galvannealed steels is one of the main advantages of this process, once the resultant coating presents a melting point similar to the melting point of the substrate, providing a better usage of the electrodes. Furthermore, the cohesion between coating and sheet allows the production of a thinner layer with higher mechanical strength, resulting in a steel sheet with superior workability and a surface that can be readily painted. Additionally, the galvannealed coating is less reactive when exposed to the atmosphere and the dissolution that occurs during corrosion process is more slowly than for a galvanized coating

(McDevitt *et al.*, 1997; Morimoto *et al.*, 1997; Kato *et al.*, 2000).

The galvannealed coating is usually composed of three Fe-Zn intermetallic phases denominated as zeta (ζ), delta (δ) and gamma (Γ) (Hansen, 1958). In galvannealed steel sheets, the zeta phase is the top layer of the multi-layer system. The gamma phase is the layer next to the steel substrate and usually is the compound with the lowest mass percentage. The delta phase is the middle layer with the highest mass percentage (Wienströer *et al.*, 2003). Since the crystallographic properties of these phases are different from each other, the mechanical and metallurgical properties presented by a galvannealed steel sheet can be dependent on the phase fraction found in the coating (Nakamori *et al.*, 1995). For example, the presence of a zeta phase may jeopardize the former process due to its high fragility, which makes a zeta phase amount of over ~8% not desirable for steel sheets applied in the automotive industry. Thus, it is very important to apply a method to quantify the phases present in the galvannealed coating to control the properties of this layer. Hong

et al. (2003) investigated and quantified Fe-Zn intermetallic phases using Mössbauer analysis, but this is too complex to be used as an industrial routine. On the other hand, X-ray diffraction (XRD) is an analytical method which already has been used in the characterization of crystalline steel sheet coatings for presenting reduced measuring time (Wienströer *et al.*, 2003). The phases present in a crystalline alloy can be easily quantified through its X-ray diffraction (XRD) pattern by means of the Rietveld Method. This method uses a least square approach to obtain a theoretical line profile that must match the experimental profile. Parameters as atomic positions and lattice parameters are refined to obtain this theoretical line (Young, 1993).

Thus, in this work, the Rietveld Method was used to quantify the phases contained in the Fe-Zn system, proving that it is an adequate method to be applied to the control of galvannealed coating phases. Additionally, since the crystalline structure of the contained phases are related to their properties, some structural information, such as lattice parameter, was also investigated.

2. Materials and methods

In order to synthesize the pure Fe-Zn intermetallic phases, the powders of iron (purity: 99.99%) and zinc (purity: 99.98%) were mixed in the nominal compositions of $\text{Fe}_{11}\text{Zn}_{40}$, FeZn_{13} and FeZn_{10} . The powder mixtures were pressed in tablet shape and then sealed into a quartz tube under an argon atmosphere. The tablets were sintered at 420°C for 11 hours, following the conditions previously reported by Kubaschewski (1982). After the sintering process, the tablets were milled and returned to the powder shape to avoid possible crystallographic

texture due the pressing step. The sample phases were identified by the qualitative analysis of XRD patterns using a powder diffractometer (Rigaku, model Miniflex, Japan) with Cu K α radiation ($\lambda = 1.5418$ Å). The XRD patterns were recorded from 30° to 60°, which is a sufficient range to obtain the main peaks of the Fe-Zn intermetallic phases, to prove that the zeta, delta and gamma phases were successfully obtained.

A specific quantity of each synthesized Fe-Zn intermetallic phase was selected and mixed. The selected mass

percentage of the mixture contained 50% of delta phase, 30% of zeta phase and 20% of gamma phase. This proportion was arbitrarily chosen, only as a matter of theoretical study. The XRD pattern of the mixture was also collected under the same conditions described previously. Finally, the XRD pattern of the mixture was analyzed by the Rietveld Method, using the DBWS 9708 software package. The theoretical quantities calculated by the Rietveld Method were then compared with the already known phase quantities to prove the accuracy of the method.

3. Results and discussion

Three Fe-Zn intermetallic phases, such as zeta, delta and gamma, which are

often formed in the galvannealed coating layer, were investigated in this work. Their

crystallographic properties are presented in Table 1.

Phases	Formula	Crystal structure	Space group	Lattice Parameter	ICSD
zeta	FeZn_{13}	monoclinic	$C12/m1$	$a = 13.3940 \text{ \AA}$ $b = 7.5980 \text{ \AA}$ $c = 5.0660 \text{ \AA}$ $\beta = 100.53^\circ$	240050
delta	$\text{Fe}_{0.64}\text{Zn}_{10.48}$	hexagonal	$P6_3mc$	$a = 12.7870 \text{ \AA}$ $c = 57.2220 \text{ \AA}$	150199
gamma	$\text{Fe}_{11.68}\text{Zn}_{39.32}$	cubic	$F43m$	$a = 17.9630 \text{ \AA}$	103711

Table 1
Summarization of crystal structures, lattice parameters and chemical formulas of the Fe-Zn intermetallic phases (ICSD, 2005).

The XRD patterns recorded for the Fe-Zn intermetallic samples are shown in Figures 1-3. By comparing the peak patterns of figures 1, 2 and 3 with the reference ones (ICSD, 2005), it is possible to identify the Fe-Zn intermetallic samples as zeta, delta and gamma phases, respectively. Figure 2 shows the presence of pure Zn and ZnO phases because of the non-completion of the sintering process and oxidization during this process. However, the low intensity of these peaks represents a low quantity of these compounds, which

is not relevant to this study. The same is shown in Figure 3 where the presence of a low quantity of ZnO was detected.

Once the XRD results showed that the zeta, delta and gamma phases were successfully synthesized, the mixture of the three intermetallic phases (50% of delta, 30% of zeta and 20% of gamma) was performed and analyzed by XRD and the resultant pattern was used as input data to the Rietveld refinement. The experimental XRD pattern, the theoretical line profile calculated by Rietveld and

their difference are shown in Figure 4. The total difference between experimental and Rietveld calculated patterns was 14%, which is considered as a good fitting, especially in the case of a pattern containing many peaks. Through the difference line, it is possible to observe that the most relevant differences were observed in the 42° to 44° range. It was already expected because a considerable peak overlapping occurs in this range, resulting in eventual mismatches between experimental and calculated patterns.

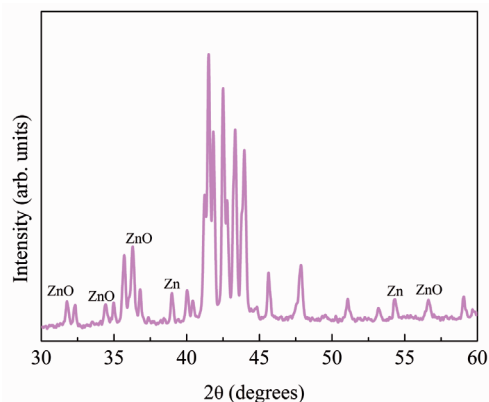


Figure 1
Experimental XRD pattern for the zeta phase.

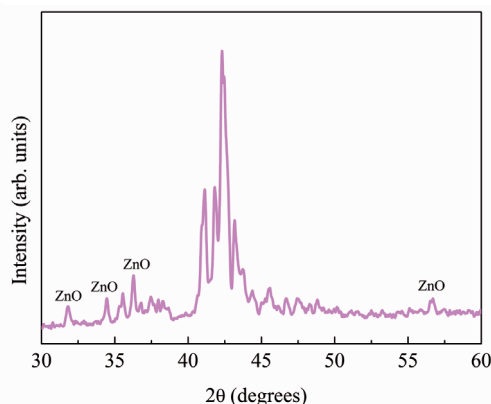


Figure 2
Experimental XRD pattern for the delta phase.

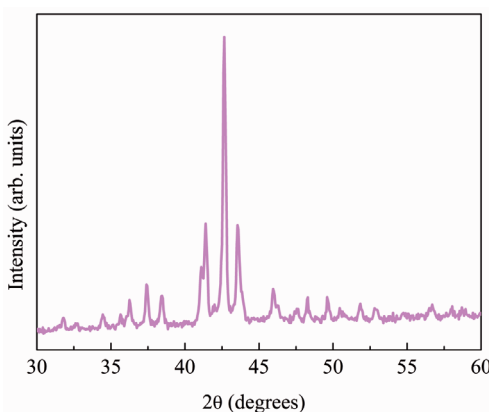


Figure 3
Experimental XRD pattern for the gamma phase.

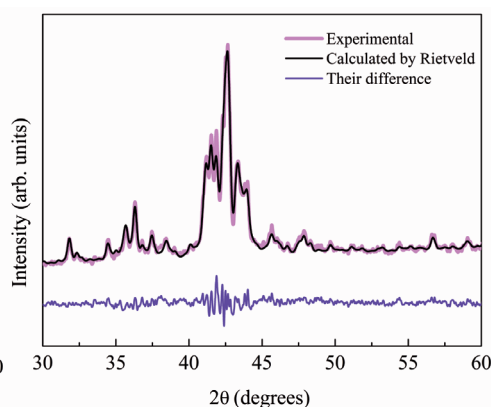


Figure 4
Rietveld refinement for the phase mixture.

The mass percentage of the phase mixture calculated by the Rietveld refinement is shown in Table 2. Since the mass percentage of the phases is related to the peak intensities, the eventual differences found between the calculated and experimental patterns could result in unreliable results. However, it is observed in Table 2 that the calculated

mass percentages are very close to the real ones. Thus, it is possible to affirm that the Rietveld Method is an accurate method to calculate the unknown mass percentage of Fe-Zn intermetallic phases. Since the Rietveld Method refines atomic parameters, such as lattice parameters, to obtain a line profile, this information was also calculated.

The best simulation was achieved with the lattice parameters $a = 13.3540 \text{ \AA}$, $b = 7.5891 \text{ \AA}$, $c = 5.0773 \text{ \AA}$, $\beta = 100.32^\circ$ for the monoclinic FeZn_{13} phase, $a = 12.7937 \text{ \AA}$, $c = 57.2871 \text{ \AA}$ for the hexagonal $\text{Fe}_{0.64}\text{Zn}_{10.48}$ phase and $a = 17.8893 \text{ \AA}$ for the cubic $\text{Fe}_{11.68}\text{Zn}_{39.32}$ phase. These values are slightly different from those given in the ICSD cards.

Phase	Real	Calculated
zeta	20%	19%
delta	30%	27%
gamma	50%	54%

Some limitations of this study should be pointed out. The Cu K α radiation used in this work is not the ideal one to perform XRD analysis on alloys containing iron. This radiation interacts with iron causing fluorescence, what may degrade the data quality. It was avoided in this study by means of a monochromator and selection of a low angle range. Furthermore, the Rietveld Method is

known for being a powder method, in order to randomize the crystals on the analyzed surface, avoiding eventual mistakes generated by crystallographic texture, a common effect found in cold-rolled steel sheets. However, the galvanized coating is resultant from a heat treatment performed after cold work, and then it is possible to affirm that the main effects of the crystallographic texture

Table 2

Comparison between the real and the Rietveld calculated mass percentage of the phase mixture.

are eliminated on the surface. Thus, performing this same analysis on industrial sheets would not represent considerable differences in the results. Additionally, once the reference patterns are known, the Rietveld refinement can be promptly performed in any XRD pattern from different galvanized steel sheets produced industrially by using the same methodology demonstrated in this study.

4. Conclusions

The Rietveld Method was used to investigate the structure of three Fe-Zn intermetallic phases and to quantify their proportion. The main conclusions of this study are the following:

- The zeta, delta and gamma phases were successfully obtained by a sintering process. This was proved through the XRD pattern analysis.

- A simple mixture of the three intermetallic phases was performed containing a known mass percentage. The XRD pattern of the mixture was collected and then analyzed by the Rietveld Method. A good agreement between experimental and calculated patterns was obtained with slightly different lattice parameters from those

given in the ICSD cards.

- Consequently, a good accordance among the mass percentage values (calculated by the Rietveld Method and the real values) was found. It shows that the Rietveld Method can be used to quantify phases in galvanized steel sheets produced industrially.

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