The Influence of Pre-Milling on the Microstructural Evolution During Mechanical Alloying of a Fe$_{50}$Cu$_{50}$ Alloy.

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Keywords: Fe-Cu, Solid solution, Premilling, Mechanical alloying, Mössbauer spectroscopy.

Abstract. Fe$_{50}$Cu$_{50}$ solid solutions have been obtained by two different mechanical alloying routes, i.e. from Fe-Cu powders with and without premilling. These two alloying processes were studied and compared at two rotation speeds of a planetary ball-mill, namely 200 rpm and 300 rpm. The microstructural evolution of the ball-milled powders as a function of milling time was monitored using electron probe microanalysis, X-ray diffraction, scanning electron microscopy and Mössbauer spectroscopy. In both processing routes, a broad distribution of different local environments of the iron atoms was observed in the Mössbauer spectra of the fcc-FeCu phase. Mössbauer spectra show further that a fcc non magnetic Fe-rich phase is formed transiently besides the bcc-Fe phase. The Fe-Cu reactions are slowed down when premilled powders are used as starting powders but similar fcc solid solutions are formed during milling. As expected, the mechanical alloying process is faster at 300 rpm but the final results are similar using 200 rpm or 300 rpm.

Introduction

Far from equilibrium synthesis processes have been used to produce materials with non-equilibrium structures, e.g. amorphous alloys, supersaturated solid solutions, metastable crystalline phases and nanocrystalline materials, with good mechanical and physical properties suitable for high performance materials. For instance, a giant magnetoresistance effect has been measured in Fe-Cu alloys for different iron concentrations [1]. According to the Fe-Cu phase diagram, Fe and Cu are immiscible at room temperature. However, alloying of Fe and Cu is known to be feasible by high-energy ball-milling [see for instance [2-10]]. During mechanical milling, iron and copper powders quickly agglomerate and are repeatedly flattened and welded by the colliding balls. This gives rise to a lamellar structure which breaks down by further co-deformation into Fe needle-shaped particules via multiple necking to rupture [6]. This process induces the formation of nanosized grains in which Fe atoms are dissolved in the Cu matrix (fcc Cu(Fe) solid solution). Single-phase structures have been obtained by MA in a broad concentration range of the Fe-Cu system, e.g. fcc Cu(Fe) solid solution for iron contents lower than 45 at.% and bcc Fe(Cu) solid solution for iron contents higher than 55 at.% [3]. In between, the two phases, Cu(Fe) + Fe(Cu) co-exist [2, 8]. However, some results of Mössbauer and of atom probe investigations indicate that these solid solutions are chemically heterogeneous on a nanometer scale [2, 5, 6, 8, 9]. Further, supersaturated solid solutions formed deep inside a miscibility gap through rapid quenching or solid state routes may have decomposition features on the scale of a few nanometers as shown recently for sputtered fcc Ag-Cu solid solutions [10] or for mechanically alloyed Fe$_{30}$Cr$_{70}$ alloys (the same conclusion holds for milled as-cast Fe$_{30}$Cr$_{70}$ alloys) [11]. Similar results on heterogeneous alloys are reported.
by other authors on mechanically alloyed Fe-Cu mixtures with and without subsequent annealing [2, 5-8]. In a recent work on nanocrystalline Fe\textsubscript{y}Cu\textsubscript{1-y} (0.141\leq y\leq 0.450) alloys produced by ball milling [6], Fernandez et al. used small-angle x-ray scattering technique to conclude that the synthesized alloys are heterogeneous on a nanometer scale since composition fluctuations were detected inside nanocrystalline grains. Wanderka et al. [5] investigated the microstructure and composition on the nanometer scale of mechanically alloyed Cu\textsubscript{50}Fe\textsubscript{50} solid solution with a tomographic atom probe. They concluded that samples contain nanometer-sized regions with strongly varying composition. By contrast, the existence of homogeneous fcc Cu(Fe) solid solution with a random distribution of iron and copper atoms in the lattice was reported by Huang [12] on Fe-Cu alloys milled at low energy. The bimodal nature of hyperfine magnetic field distributions found in Fe\textsubscript{50}Cu\textsubscript{50} prepared by high-energy ball-milling and further ball-milled at low energy [7] was explained either by a nonrandom distribution of Fe atoms in the alloy, by variations in the magnetic moments of Fe atoms depending on their local environments, or by composition fluctuations of the order of 1 at.%. It was not possible to distinguish between these causes.

Although MA has long been used for the preparation of Fe-Cu alloys, little attention has so far been paid to the effect of the characteristics of the starting powders on the microstructural evolution during milling. In fact, premilling of the initial powders, which modifies some of their mechanical characteristics and, consequently, the transfer of energy from colliding balls to entrapped powder particles [13], might change the way Fe and Cu are mixed at a nanometric scale. The highest solubility of magnesium in titanium was for instance achieved when both elemental powders were made brittle by premilling before mechanical alloying [14]. Mechanical mixing by shearing during plastic deformation of powder particles is expected to occur when mixtures of as-received powders of ductile elements are ground while mixing due to welding of the particles should occur predominantly when the powders are premilled separately and then mechanically alloyed together. We have investigated the phase transformations of a Fe\textsubscript{50}Cu\textsubscript{50} alloy during mechanical alloying of powder mixtures, with and without pre-milling, in order to characterize the influence of the mixing mode on the final stationary state. These two alloying processes were studied and compared at different milling speeds of a Fritsch P6 planetary ball-mill (200 and 300 rpm). Firstly, the classical method of mechanically alloying of as-received elemental Fe and Cu powder blends was used. Later on, as-received powders of Fe and Cu were milled separately and then mixed in the required proportions and milled again.

**Experimental details**

Fe\textsubscript{50}Cu\textsubscript{50} alloys were synthesised by mechanical alloying of Fe and Cu powders with a nominal purity of 99% and an average particle size of 60 µm and 150 µm, respectively. Milling was performed in a planetary ball mill (Pulverisette 6 from Fritsch) using hardened steel vial (250 ml) and balls (15 balls with 20 mm diameter each). A ball-to-powder weight ratio of 20:1 was used. The disc rotation speeds were either 200 or 300 rpm. Milling was performed in an argon atmosphere to avoid air contamination. The powders were mechanically alloyed either in the as-received condition (also named hereafter “straight condition”) or after premilling the elements separately at 550 rpm and 150 rpm in the case of Fe and Cu respectively. The tendency of Cu powder particles to agglomerate increases when the rotation speed increases. Thereby, two different batches were obtained. The Vickers hardness of the Cu and Fe powders, as-received and after milling, was measured with a 100g load, in a Shimadzu equipment, on cold compacted samples. Average values of 193 and 250 HV were obtained for milled Cu and Fe powders respectively, against 49 and 80 HV for the as-received powders. The as-mechanically alloyed powders were characterized by means of electron probe microanalysis (EPMA), X-ray diffraction (XRD), scanning electron microscopy (SEM) and Mössbauer spectroscopy (MS). The X-ray diffraction patterns were obtained using Co K\textsubscript{α} radiation (λ=0.17889 nm). The mean crystallite sizes and microstrains were
obtained from the widths of X-ray diffraction peaks using the Williamson-Hall method [15]. Mössbauer spectra were recorded at room temperature in transmission geometry with a spectrometer operating in conventional constant acceleration mode. A $^{57}$Co source in Rh matrix with a strength of $\approx$ 10 mCi was used. The spectra were analysed by the method of Le Caër [16], which extracts a hyperfine magnetic field distribution, $P(B)$, from an experimental spectrum. Lorentzian line-shapes were employed in this procedure.

**Results and discussion**

The X-ray diffraction patterns of the two Fe$_{50}$Cu$_{50}$ sample batches milled at 200 and 300 rpm as a function of milling time are shown in figures 1 and 2, respectively. In all cases, a broadening of the diffraction peaks of both Cu and Fe occurs gradually with increasing milling time. The intensities of the diffraction peaks of Fe decrease more rapidly than those of Cu do. The diffraction peaks of Cu are no more observed after different periods of time depending on the milling process.

![Figure 1 - XRD patterns of two Fe$_{50}$Cu$_{50}$ samples as a function of milling time (200 rpm) of: (a) starting mixtures of as-received powders, (b) starting mixtures of premilled powders.](image)

![Figure 2 - XRD patterns of two Fe$_{50}$Cu$_{50}$ samples as a function of milling time (300 rpm) of: (a) starting mixtures of as-received powders, (b) starting mixtures of premilled powders.](image)
Starting from as-received powder mixtures, Figs. 1(a) and 2(a), the X-ray peaks of Fe are no longer visible for milling times longer than 40h and 8h, respectively. When initial blends are prepared from premilled powders, Figs. 1(b) and 2(b), vanishing of the Fe diffraction lines occurs for longer milling times, i.e. longer than 50 h and 16h, respectively. In both processes, iron is incorporated in the fcc-Cu structure, giving rise to a fcc-Cu(Fe) solid solution whose diffraction peaks shift to higher angles with milling time as a consequence of a steadily increasing solubility of iron in copper. In equilibrium, the elements Fe and Cu are immiscible and the equilibrium Fe-Cu phase diagram does not exhibit intermediate phases. The lattice parameters of the fcc-Cu(Fe) solid solutions obtained after the two milling processes (straight and premilled powders) are equal to 0.3638 nm for 200 rpm and 0.3636 and 0.3635 nm, for 300 rpm and for straight and premilled processes respectively, which correspond to expanded fcc-Cu lattices (a=0.3615 nm for pure Cu). In spite of the larger atomic size of copper when compared to iron, the fcc Cu lattice is expanded by the incorporation of iron atoms in agreement with previous works [2,4,12]. Harris et al [17] propose a model in which the strong interaction between like atom pairs (Fe-Fe, Cu-Cu) provides Coulombic forces which establish geometrical limitations which prohibit the unlike Fe-Cu pairs from coming any closer than the measured distances. In fact, the dilated bonds measured between Cu and Fe indicate that either a very weak interaction or a repulsion exists between unlike neighbours as a result of a positive heat of mixing in this system [16]. The crystallite sizes and microstrains in the stationary state of the milled powders, as determined by X-ray analysis are 9.5 nm and 0.4% (straight process) and 6.3 nm and 0.5% (premilled powders), respectively, for 200 rpm and 23 nm and 1.5% (straight process) and 13 nm and 1% (premilled powders), respectively, for 300 rpm. The crystallite size and microstrain measured in premilled Fe are 42 nm and 0.3%, respectively, while the premilling of Cu reduces the crystallite size down to 52 nm with a microstrain of 0.2%.

Figure 3 shows the standard-deviation $\sigma$ of the distribution of chemical composition, as estimated from EPMA measurements performed at 10 different points in each sample, as a function of milling time.

![Figure 3 - Standard deviation of the distribution of chemical composition in particles of milled powders as a function of milling time and of milling conditions. A.R. = as-received powders; P-M. = premilled powders.](image-url)
As expected, $\sigma$ decreases with milling time because the powder particles become gradually chemically homogeneous during mechanical alloying. The premilling process slows down the chemical homogenization for both 200 and 300 rpm speeds, in agreement with the XRD results. Moreover, when the rotation speed is increased, the temperature and the injected specific power are increased. These phenomena speed up diffusion and lead, consequently, to an acceleration of the chemical homogenisation process and formation of a Cu(Fe) solid solution.

The final overall chemical composition of the samples is also presented in fig. 3. As can be seen, samples prepared from premilled powders are enriched in iron while the opposite is found for samples prepared from as-received powders. The sample obtained from premilled powders with a milling speed of 200 rpm for 150h is the one presenting the higher iron contamination. This means that the initial hardness of powders play an important role on their contamination during milling. In fact, the higher is the initial hardness, the more abrasive is the mixture and consequently the higher is the iron contamination. This phenomenon is increased by long-term milling processes.

Figures 4 and 5 show room-temperature Mössbauer spectra of two sample batches, i.e. (a) straight and (b) premilled, for different times of milling and for 200 rpm and 300 rpm, respectively.

![Figures 4 and 5](attachment:image.png)

Figure 4 – Room temperature $^{57}\text{Fe}$ Mössbauer spectra and the corresponding hyperfine magnetic field distributions for Fe50Cu50 powders after different milling times (200 rpm) of : (a) starting mixtures of as-received powders, (b) starting mixtures of powders of Fe and of Cu premilled separately.

Three different components can be distinguished. The first is a broadened sextet caused by the occurrence of a hyperfine magnetic field distribution which is attributed to the fcc-FeCu phase and is due to a broad distribution of environments of Fe atoms. The latter broad sextet demonstrates that Fe and Cu are alloyed at the atomic level but it does not necessarily imply that the corresponding
alloy is compositionally homogeneous. The second is a sextet with a hyperfine field of 33 T characteristic of the bcc-Fe phase. It should be noted that the bcc-Fe phase was not detected by X-ray diffraction for milling periods equal or longer than 40h (straight process) or than 50h (premilled powders) for 200 rpm and longer than 16h (straight process) or than 24h (premilled powders) for 300 rpm. The latter sextet can be explained by the very small amount and the small crystallite size of the residual bcc-Fe phase which may at least partly come from a contamination by milling tools. The third transient contribution is a single line with an isomer shift of –0.09 mm/s characteristic of the fcc-Fe phase [19]. The latter phase was not detected by X-ray diffraction, likely for similar reasons. These observations are in agreement with a model proposed by Gente et al [20] and with results of Jiang et al [21] for samples obtained from powder mixtures of as-received elements. Similar results are obtained when starting from premilled powders.

Figure 5 – Room temperature $^{57}$Fe Mössbauer spectra and the corresponding hyperfine magnetic field distributions for Fe50Cu50 powders after different milling times (300 rpm) of: (a) starting mixtures of as-received powders, (b) starting mixtures of powders of Fe and of Cu premilled separately.

Figures 6 and 7 show some SEM micrographs of particle morphologies as a function of milling time. As can be seen, in the final state the samples have a pellet shape of about 0.7 mm in diameter and about 0.3 mm in thick, independently of the milling process (Fig. 6 (a) and (b)). This reflects the fact that the fracture-welding equilibrium is shifted in the direction of welding, as a result of the high ductility of copper. This shape is obtained earlier in the straight milling process. Fig. 7 compares the morphologies of the particles milled for 1h using the two milling processes. In the case of inverted back-scattered (BE) images, black and white zones correspond to copper and iron respectively. In the straight milling process, mixing is faster than in the second process, in which plates of iron and copper can be easily differentiated. Quantitative electron microprobe analysis confirms these observations. No significant compositional variations were observed in particles obtained from the straight process after 1h of milling. By contrast, there are some compositional fluctuations in particles alloyed from premilled powders for 1h (Fe- and Cu-rich zones). Some iron particles were detected in the latter powders. The Cu-rich particles are actually Cu(Fe) solid
solutions with a significant iron amount. This means that, in contrast to the bcc-Fe structure, there is an incorporation of iron in the fcc-Cu structure from the beginning of the milling process.

Figure 6 – SEM particle morphologies (70x) as a function of milling time (300 rpm): (a) initial mixture of as-received powders, (b) initial mixture of powders of Fe and of Cu premilled separately.

Figure 7 – SEM images (1500x) of powder particles as a function of milling time. (a) initial mixture of as-received powders, (b) initial mixture of powders of Fe and of Cu premilled separately. SE = secondary electrons, BE = back-scattered electrons. (300rpm).

Conclusions

The XRD and MS results show that it is possible to incorporate iron in the fcc-Cu matrix by the two milling processes used in this work. In both procedures, a broad distribution of different local environments of iron atoms was observed in the Mössbauer spectra of the fcc-FeCu phase. Mössbauer spectra show further that a fcc non magnetic Fe-rich phase is formed transiently. The reactions are slowed down when premilled powders are used as starting materials. In our milling conditions, similar fcc solid solutions are formed for 50h (straight condition) and 150h (premilled powders) when 200 rpm are used, and for 24 h (straight condition) and 32h (premilled powders) when 300 rpm are used. As expected, it was observed that at 300 rpm the alloying process occurs faster than at 200 rpm, both in the straight and premilled conditions.

Acknowledgements

This work was partly funded by the Portuguese-French ICCT-Ambassade de France à Lisbonne cooperation. The authors also thank Pierre Delcroix (LSG2M) for useful suggestions.

References

[1] C.L. Chien, J.Q. Xiao and J.S. Jiang, J. Appl. Phys. 73 (1993), p. 5309.
[2] L. B. Hong and B. Fultz, Acta mater. 46 (1998) 2937.
[3] J.Z. Jiang, C. Gente and R. Bormann, Mater. Sci. Eng. A 242 (1998), p. 268.
[4] P. Crespo and A. Hernando, Recent Res. Devel. Nanostructures, 1 (1999), p. 63.
[5] N. Wanderka, U. Czubayko, V. Naundorf, V.A. Ivchenko, A. Ye. Yermakov, M.A. Uimin and H. Wollenberger, Ultramicroscopy 89 (2001), p. 189.
[6] M.B. Fernandez van Raap, L.M. Socolovsky, F.H. Sanchez and I.L. Torriani, J. Phys. Condens. Matter 14 (2002), p. 857.
[7] J. Xu, G.S. Collins, L.S.J. Peng and M. Atzmon, Acta mater. 47 (1999), p. 1241.
[8] P.J. Schilling, J.H. He, R.C. Tittsworth and E. Ma, Acta mater. 47 (1999), p. 2525.
[9] S.D. Kaloshkin, I.A. Tomilin, G.A. Andrianov, U.V. Baldokhin, E.V. Shelekhov, Mater. Sci. Forum 235-238 (1997), p. 565.
[10] J.H. He, H.W. Sheng, J.S. Lin, P.J. Schilling, R.C. Tittsworth and E. Ma, Phys. Rev. Lett. 89 (2002), p. 125507.
[11] P. Delcroix, T. Ziller, C. Bellouard, G. Le Caër, Mater. Sci. Forum 360-362 (2001), p. 329.
[12] J.Y. Huang, J. Mater. Sci., 31 (1996), p. 4165.
[13] D. Choulier, R. Rahouadj and E. Gaffet, Ann. Chim. Sci. Mat. 22 (1997), p.351.
[14] E. Zhou, C. Suryanarayana and F.H. Froes, Mater. Lett. 23 (1995), p. 27.
[15] G. Williamson and W.H. Hall, Acta Metall. 1 (1952), p.22.
[16] G. Le Caër and J.M. Dubois, J. Phys. E 12 (1979), p. 1083.
[17] V.G. Harris, K.M. Kemner, B.N. Das, N.C. Koon, A.E. Ehrlich, J.P. Kirkland, J.C. Woicik, P. Crespo, H. Hernando, A. Garcia Escorial, Phys. Rev. B 54 (1996), p. 6929.
[18] E. Ma, H.W. Sheng, J.H. He and P.J. Schilling, Mater. Sci. Eng. A 286 (2000), p. 48.
[19] R. Halbauer and U. Gonser, J. Magn. Magn. Mater. 35 (1983), p. 55.
[20] C. Gente, M. Oehring and R. Borman, Phys Rev B 48 (1993), p.13244.
[21] J.Z. Jiang, U. Gonser, C. Gente and R. Borman, Appl. Phys. Lett. vol 63, no.20 (1993), p. 2768.