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Phase transformations and thermodynamic properties of nanocrystalline FePt powders


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Abstract: The solid state reactions and structural evolution in nanocrystalline FePt powders during mechanical ball milling at 77 K and subsequent annealing have been investigated. Above 310 °C the formation of L10 FePt is observed. As the milling time increases, the enthalpy evolved during the transformations is reduced, whereas the corresponding activation energy increases. For accelerating the ordering process a fine lamellar structure of Fe and Pt is favourable.

Keywords: Mechanical milling, X-ray diffraction (XRD), Rietveld method, Differential scanning calorimetry (DSC), Ordering, L10 phase

1. Introduction

The chemically ordered tetragonal L10 FePt phase has attracted much attention as a material for special permanent magnet applications and, more recently, for ultra-high density magnetic recording [1]. The interest in L10 FePt is a consequence of its excellent intrinsic magnetic properties ($K_1 = 6.6 \text{ MJ m}^{-3}$, $J_s = 1.43 \text{ T}$, $T_c = 750 \text{ K}$ [2]), as well as high ductility and good corrosion resistance. Several approaches are used to obtain ordered L10 phase [3]. It can be formed by annealing of the disordered face-centered cubic (fcc) (A1) phase [2], [4] and [5]. In the thin film-form, the L10 FePt phase can be directly prepared by deposition onto heated substrates [6]. Ordered FePt thin films have been prepared as well by the subsequent annealing of multilayers produced by sequential deposition of separate Fe and Pt layers having thickness in the order of several nanometers [7]. Nanocrystalline FePt powders with a coercivity up to 1 T can also be prepared by mechanical alloying of elemental powders at 77 K followed by annealing at moderate temperatures [8] and [9]. The mechanisms of phase formation and ordering (i.e. continues or discontinues) differ depending on the approach used to obtain the L10 phase [5], [6], [9], [10] and [11]. In the present work, we investigate structural transformations occurring upon milling and subsequent annealing of the FePt powders. An improved understanding of thermodynamics and phase formation in nanocrystalline FePt should be equally valuable for other nanoparticulate FePt systems.

2. Experimental

For the preparation of equiatomic FePt, mixtures of elemental Fe and Pt powders with a purity of 99.9% and a particle size less than 80 μm were mechanically ball-milled at liquid nitrogen and room temperature for 1–7 h in a Misuni vibration mill. Heat treatment of the as-milled powders was carried out in the temperature range of 350–550 °C under argon atmosphere. The structure of the as-milled and annealed powders was investigated by X-ray diffraction (Philips X’Pert diffractometer equipped with a diffracted beam monochromator) using Co-Kα radiation. Rietveld refinement (X’Pert Plus software) was used to determine the phase composition, lattice constants and order parameter. Average crystallite size ($D$) and root mean square (rms) strain ($e$) were determined by a modified Williamson–Hall analysis [12]. The line profile parameters were extracted from the Rietveld refinement, in the assumption of a pseudo-Voigt profile, and corrected for the instrumental broadening. As a standard reference material, a LaB6 powder specimen has been used. The microstructure of the FePt powders was investigated by means of field emission gun scanning electron microscopy (FEG-SEM LEO1530) equipped with energy dispersive X-ray (EDX) analysis. The thermal behaviour was examined by differential scanning calorimetry (DSC 404C Netzsch) over the temperature range of 50–1000 °C with heating rates of 5–40 °C/min under flowing argon.
3. Results

3.1. Milling of FePt

Fig. 1 shows the X-ray diffraction patterns of the Fe–Pt powders prior to and after milling at liquid nitrogen temperature for different time intervals. Considerable changes in the phase composition and microstructure are observed after short milling times. Initially sharp X-ray diffraction lines are appreciably broadened after only 1 h of milling. In order to separate the contributions to the overall breadth from crystallite size and lattice strain the modified Williamson–Hall analysis, the so-called “average size-strain” method of Langford [12] was applied. The effect of stacking faults to the peak broadening was not considered in this study. It has to be noted, however, that no peak displacements produced by deformation faulting were detected for the FePt powders studied. The “average size-strain” plot for the 1 h milled Fe/ Pt powder yields \( Y_D \) of about 29 nm for Pt and 43 nm for \( \alpha \)-Fe, i.e. substantial reduction in the grain sizes compared to the starting microcrystalline powders is observed. The root mean square (rms) strain \( \epsilon \) for Pt and \( \alpha \)-Fe was estimated to be about 0.16% and 0.21%, respectively. Further milling results in a reduction of the grain sizes of Pt \( (D \approx 23 \text{ nm}) \) and \( \alpha \)-Fe \( (D \approx 17 \text{ nm}) \) and an increase of the rms strain to 0.27% and 0.55%, respectively. High resolution scanning electron microscopy (SEM) investigation reveals that the microstructure of the powder milled for 2 h is a lamellar structure of the constituent elements having a thickness of about 20–300 nm (Fig. 2). The fine contrast within the Pt lamellae confirms the nanocrystalline nature of the powders. Moreover, Rietveld refinement reveals the presence of the fcc (A1) FePt phase having dimensions of ~5 nm in the powders milled for 2 h. Thus, alloying starts to occur at this milling stage. The following phase contents were determined for this powder: \( \alpha \)-Fe (~31 vol.%), Pt (~36 vol.%) and A1 FePt (~33 vol.%).

Increase of the milling time to 5 h yields a further refined lamellar structure of the constituent elements (not shown). The \( \alpha \)-Fe and Pt fractions are reduced, whereas the amount of A1 FePt increases (Fig. 3). The grain sizes of \( \alpha \)-Fe and Pt reduce to ~7 and 9 nm, respectively. However, this minimum grain size does not correspond to the maximum strain: rms strain of \( \alpha \)-Fe and Pt reduces to about 0.05%. That is, the rms strain increases rapidly at the first stage of the milling, but for longer milling times it decreases. A similar behaviour was observed for low energy ball-milled pure metals [13]. The grain size and lattice strain of the A1 FePt phase after milling for 5 h were determined to be about 15 nm and 0.65%, respectively.

Fig. 1. X-ray diffraction patterns of the FePt powders prior to and after milling for various times at liquid nitrogen temperature.

Fig. 2. High resolution SEM image (backscattered electron mode) of the FePt powder milled at liquid nitrogen temperature for 2 h.
After 7 h of milling at liquid nitrogen temperature A1 FePt fraction reaches 84.4(7) vol.%, the rest being α-Fe. The grain size of the A1 phase is almost unchanged, although the lattice strain decreases to 0.49%. The lattice constant of the A1 FePt phase formed by 7 h of milling was determined to be \( a = 0.38196(2) \) nm. SEM investigation revealed that the observed α-Fe is due to the iron from the balls and vial walls, which is usually observed in powders milled with steel as a grinding medium [14].

For comparison, milling at room temperature was also performed and X-ray analysis indicates that the formation of the A1 FePt phase occurs faster. However, the crystallite sizes are increased and the yield of the powder after room temperature milling is extremely low due to the cold welding of the material onto the milling media.

3.2. Thermal behaviour of the FePt powders milled at 77 K

As an example, Fig. 4 shows DSC traces for the Fe_{50}Pt_{50} powders milled for 2, 5 and 7 h. Rietveld refinement of X-ray data of the samples annealed to temperatures before and after the transformation confirms that the exothermic reaction is due to the formation and ordering of new phases. With increasing milling time, the minimum of the transformation peak \( T_m \) shifts towards lower temperatures: from about 440 °C for 2 h milled powder to 367 °C for 7 h milled powder. It has to be noted, that the onset of the transformation appears essentially at the same temperature of about 310(4) °C independent of milling time. The transformation peak becomes narrower with increase of milling time: its full width at half maximum decreases from about 120 °C (2 h milled) to 56 °C (7 h milled). The amount of heat \( \Delta H \) evolved during the transformations, determined from the area under the peaks, decreases correspondingly from about 110(20) J/g to 70(9) J/g and 55(2) J/g as the milling time increases from 2 to 5 and 7 h. Isothermal DSC measurements at 290 °C, i.e. below the onset temperature of the transformations, have shown a signal that only decayed with time.

For the analysis of the DSC curves, the Kissinger method [15] was employed using heating rates \( \beta \) between 5 and 40 °C/min. By using the linear relationship

\[
\ln\left(\frac{T_m^2}{\beta}\right) = \frac{E_a}{k_B T_m}
\]

it is possible to obtain values for the activation energy \( E_a \) of the processes involved in the transformations, where \( T_m \) is a temperature of the transformation peak and \( k_B \) is the Boltzmann’s constant. The corresponding Kissinger plot is shown in Fig. 5. The obtained activation energies were found to increase with milling time from 1.03(8) eV for 2 h milled powder to 1.24(11) and 1.45(10) eV for the powders milled for 5 and 7 h, respectively.
3.3. Structural analysis of the milled and annealed FePt powders

In order to obtain information on structural changes induced by heat treatment the isothermally annealed and subsequently quenched FePt powders were investigated using quantitative X-ray diffraction Rietveld analysis and high resolution SEM. Annealing was carried out at temperatures in the vicinity of the DSC transformation peaks and above (350–550 °C) for a series of times between 10 min and 48 h.

Heat treatment of the 2 h milled FePt powder at 450 °C induces an immediate formation and ordering of L10 FePt and two additional Fe3Pt (L12) and FePt3 (L12) phases. After a short heat treatment (e.g. after 10 min), unreacted α-Fe (7 vol.%) and Pt (5 vol.%), as well as A1 FePt (13 vol.%) are present. L10 FePt, Fe3Pt and FePt3 fractions are equal to 53, 6 and 15 vol.%, respectively. With increasing annealing time the amount of residual Fe and Pt is reduced and no elemental Fe and Pt are observed upon annealing at 450 °C for 16 h and longer. At the same time, L10 FePt fraction increases and the Fe3Pt and FePt3 fractions are reduced. The Rietveld refinement of X-ray diffraction patterns (not shown here) of the FePt powders annealed at 450 °C/48 h yields the following phase contents: about 84 vol.% of L10 FePt, 8 vol.%-A1 FePt, 5 vol.%-FePt3 and 3 vol.%-Fe3Pt.

Increased annealing temperature promotes faster reaction rates. For instance, after annealing for 10 min at 500 and 550 °C the amounts of the L10 FePt phase are 62(1) vol.% and 71.1(7) vol.%, respectively. The higher the annealing temperature, the smaller is the amount of the Fe3Pt and FePt3 phases. However, these phases are still present in the samples annealed for 48 h. For the annealing conditions studied, the maximum FePt (L10) content was about 96(1) vol.% after heat treatment at 550 °C/48 h, the rest being A1 FePt (~2 vol.%) and L12 FePt3 (~2 vol.%).

The following lattice parameters of the L10 FePt phase in the heat treated FePt powders were determined from the Rietveld analysis: 

$$a = 0.3858(2) \text{ nm}$$

$$c = 0.3716(2) \text{ nm}$$

$$c/a = 0.964.$$ Lattice constants of the A1 phase were equal to 

$$a = 0.3820(3) \text{ nm}.$$ No significant changes in the lattice parameter of the L10 and A1 FePt phases were observed in dependence on the annealing conditions studied.

Rietveld analysis was also employed for the investigation of site occupations in the L10 FePt phase and the determination of the L10 FePt order parameter: 

$$S = r_d + r_e - 1,$$

where 

$$r_d$$

is the fraction of Pt(Fe) sites occupied by the correct atom [16]. When the order is perfect, the order parameter S reaches unity, while for a completely random atom arrangement S is equal to zero. The order parameter of the L10 FePt phase in the heat-treated FePt powders was found to be approximately 0.9, independent of annealing conditions. Thus, heat treatment of the powders milled for 2 h at liquid nitrogen temperature results in the formation of the chemically highly ordered L10 FePt phase.

The high resolution SEM micrograph of the FePt powder annealed at 450 °C for 48 h (Fig. 6) shows that the microstructure preserves a lamellar character after annealing. Three types of compositional contrast can be distinguished. Bright layers have the stoichiometry of Fe25Pt75. They are surrounded by Fe50Pt50 layers, followed by a layer of Fe75Pt25 (darkest contrast). The interfaces between the layers are sharp. The high resolution SEM image reveals as well that the size of the crystallites within the lamellae is substantially smaller than the lamellae thickness.

To determine the average crystallite size of the L10 FePt phase, the “average size-strain” method was applied. Fig. 7 shows the dependence of the L10 FePt grain size on annealing time at different temperatures. With increasing annealing time a comparatively slow grain growth is observed. The grain growth, however, is more pronounced at 550 °C. Heat treatment at this temperature for 48 h results in a grain size
of about 39 nm, which is noticeably higher than that of the powder annealed at 450 °C (\(\langle D \rangle \approx 30 \text{ nm}\)). The rms strain is in the range of 0.11–0.15%. In all the studied FePt powders, no peculiar broadening of the superstructure reflections relative to the fundamental reflections was detected, i.e. no indication of the presence of the antiphase domains in the L10 phase was observed.

Heat treatment at 350 °C of the FePt powders milled for 7 h induces the A1 FePt → L10 FePt transformation. A substantial amount of the A1 FePt phase (67%) is transformed after only 10 min of annealing at this temperature. The transformed L10 fraction increases with annealing time and reaches about 89% after heat treatment at 350 °C for 16 h. The order parameter of the L10 FePt phase was found to increase from \(S = 0.72(2)\) to \(S = 0.87(3)\) for the powders annealed for 10 min and 16 h, correspondingly. Along with the disordered and ordered FePt phases, \(\alpha\)-Fe was detected in the annealed powders. Its fraction (~15 vol.%) is constant with annealing time, i.e. it is not consumed during the transformation. Considering that 15 vol.% of \(\alpha\)-Fe was already present in the as-milled powders, it can be concluded that the observed \(\alpha\)-Fe is due to the additional iron from the milling media.

At 450 °C, the transformation of the A1 phase into L10 FePt phase in the 7 h milled FePt powders is essentially finished after 10 min of annealing. The amount of L10 FePt reaches about 78 vol.%, A1 FePt and \(\alpha\)-Fe fractions are equal to 7 and 15 vol.%, respectively. Again, assuming that \(\alpha\)-Fe is essentially milling media debris, one obtains that about 91% of A1 FePt is consumed during the A1 → L10 transformation. Further increase of annealing time does not lead to substantial changes in the phase composition. However, order parameter of the L10 FePt phase slightly increases from 0.92(1) to 0.97(2) in the powders annealed for 10 min and 48 h, respectively. Average crystallite size of L10 FePt was found to slowly increase from 23 to 30 nm upon increasing the annealing time from 10 min to 48 h.

In order to compare L10 fractions transformed during annealing, the 5 h milled FePt powders were heat treated at 450 °C for 48 h. The following phase composition was determined: 86.7(7) vol.% of L10 FePt, 3.4(5) vol.% of A1 FePt and 9.9(7) vol.% of \(\alpha\)-Fe. Assuming that about 10 vol.% of \(\alpha\)-Fe is from milling media, the transformed L10 FePt fraction is estimated to be 97%. The order parameter of the formed L10 phase is about 0.98(2).

4. Discussion

The evolution of structure observed during ball milling of elemental Fe and Pt powders at liquid nitrogen temperature is consistent with the mechanism of alloying in a system with two different ductile components [14]. The yield of the powder is much higher after milling at 77 K compared to that at room temperature due to the minimised cold welding of the powder onto the milling media. When milling is performed at 77 K, the formation of the disordered A1 FePt phase starts later compared to room temperature milling. This allows to adjust the microstructure of the FePt powders and, depending on the aim, to finish with either a lamellar structure of the constituent elements (multilayer-type), disordered A1 FePt phase or an intermediate structure, consisting of fine lamellae of Fe and Pt, with the A1 FePt phase formed at their interfaces [17].

Heat treatment of the as-milled powders induces combined solid state reactions that take place in solids in which thermodynamic equilibrium is approached by more than one elementary reaction [18]. It leads simultaneously to the decrease of defect density, nucleation and growth of the new phases concurrent with the ordering process. Comparing the formation of the L10 phase starting from A1 or lamellar precursors, one should not overlook a principal difference in the mechanisms of the phase formation. In a recent study [9], we have shown that on annealing Fe\(_{100−x}\)Pt\(_x\) (\(x \neq 50\)) powders with a lamellar structure it is energetically more favourable to form highly ordered FePt (L1\(_0\)) and additional Fe\(_3\)Pt (L12) or FePt\(_3\) (L12) phases, rather than nonequilibrium FePt (L1\(_0\)), as it could have been expected according to the equilibrium Fe–Pt phase diagram. Here it is observed that neither the lattice constants nor the order parameter of the L10 FePt phase are changed substantially once the phase is formed. However, if the initial powders predominantly have a disordered A1 structure, heat treatment induces ordering of the A1 FePt phase into L10 FePt. In the 7 h milled FePt powders, the order parameter \(S\) of the L10 phase formed as a result of A1 FePt → L10 FePt transformation increases gradually with increasing annealing time and temperature.

The peak of the transformations observed in the constant heating rate DSC curves is shifted towards lower temperatures with increase of milling time. This can be attributed first, to the finer lamellae and smaller grain size of the phases, favouring diffusion along grain boundaries and second, to the alloying of the elements on an atomic level. In the isothermal DSC measurements at 290 °C no distinct peak was detected. Rietveld analysis of the samples after the measurements reveals the presence of the L10 FePt and, in the case of the 2 h milled powder, additional Fe\(_3\)Pt and FePt\(_3\) phases. The apparent absence of a peak in the isothermal DSC measurements has been ascribed to transformations starting with fast nucleation from random positions followed by a comparatively slow diffusion-controlled growth [19] and [20]. Slow growth rates could be supported, for instance, by the data shown in Fig. 7. Interdiffusion between the constituent elements, which has already occurred during the milling, may provide the necessary mixing of atoms, thus favouring rapid nucleation of the new phases. This could explain the appearance of the transformation onset at almost
the same temperature of 310 °C in the powders milled for different times.

For the 7 h milled powder, consisting mainly of the A1 FePt phase, no long distance diffusion is required to form the L10 phase and consequently the reaction is completed at lower temperatures. The highest apparent activation energy of 1.45(10) eV was observed for this powder. The amount of heat ΔH ≈ 155 J/g evolved during the transformation was smallest, implying that less energy was stored compared to the powders milled for 2 and 5 h. The lowest activation energy Ea = 1.03(8) eV and highest enthalpy change of about 110 J/g was obtained for the powders milled for 2 h, i.e. for the lamellar structure of Fe and Pt with a ~5 nm thick A1 FePt at their interfaces. An additional driving force resulting from the concentration gradient presence could be expected for this powder. However, higher temperatures would be needed for the completion of the reactions, since to form the new phases atoms should diffuse over longer distances.

When comparing the L10 fractions transformed after annealing at 450 °C/48 h for the 7 h (Ea = 1.45(10) eV) and 5 h (Ea = 1.24(11) eV) milled powders, it turns out that the L10 amount is higher in the latter case and apparently no additional phases, like Fe3Pt or FePt3, are formed. Therefore, to achieve larger amounts of highly ordered L10 FePt, a fine lamellar structure is favourable.

The observed activation energies are substantially lower than the activation energy of 3.1 eV for interdiffusion in single-crystalline A1 FePt [21]. As shown in Ref. [20], activation energies for grain growth, phase formation and ordering in nanocrystalline materials are lower than for bulk diffusion and are of the same order of magnitude as those for fast diffusion along defects, such as dislocations or grain boundaries. Therefore, one could expect that in the studied FePt powders diffusion over the grain boundaries is a determining factor for the phase transformations, ordering and grain growth. The obtained activation energies are comparable to those found for the A1 → L10 transformation in sputter deposited FePt films [10] and [11].

5. Conclusions

Nanocrystalline FePt powders were prepared by mechanical ball milling at liquid nitrogen temperature. Depending on the milling time, either a lamellar (multilayer-type) structure of Fe and Pt, or A1 FePt or an intermediate structure, consisting of fine lamellae of Fe and Pt with A1 FePt formed at their interfaces was obtained. On annealing at temperatures just above 310 °C, combined solid state reactions become activated resulting in the formation of L10 FePt phase and, in the case of the 2 h milled powder, additional Fe3Pt and FePt3 phases. The amount of heat evolved during the transformations, decreases from about 110 to 70 J/g and 55 J/g as the milling time increases from 2 to 5 and 7 h. Diffusion over the grain boundaries is supposed to be the determining factor for the phase transformations, ordering and grain growth. The transformation activation energy of 1.03(8) eV obtained for the lamellar structure of the constituent elements was found to be lower than that for the disordered A1 → ordered L10 transformation (1.45(10) eV).

It is concluded that in order to accelerate the ordering process a fine lamellar structure should be favoured.

References