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PHASE TRANSFORMATIONS ON ANNEALING OF NON-CRYSTALLINE FERROUS HALIDES (*)

F. J. LITTERST, G. M. KALVIUS and A. J. F. BOYLE (**)  
Physik-Department der Technischen Universität München  
D 8046 Garching, James-Franck-Str., BRD

Résumé. — Des échantillons non cristallins de FeF₃, FeCl₂ et FeBr₂ obtenus par condensation de vapeurs à basse température, ont été convertis en phase cristalline par recuit à des températures allant jusqu'à 800 K. On trouve que FeCl₂ et FeBr₂ ne se convertissent jamais directement de l'état non cristallin en l'état cristallin habituel. Ils forment entre 60 et 350 K une phase intermédiaire inconnue. Par contre, FeF₃ se convertit directement. Ceci a lieu à des températures beaucoup plus élevées (~ 800 K). L'existence d'une phase intermédiaire pour FeCl₂ et FeBr₂ mais non pour FeF₃, est discutée en fonction des différents changements structuraux ayant lieu lors du processus de recristallisation des dihalogénures de fer non cristallins.

Abstract. — Non-crystalline samples of FeF₃, FeCl₂ and FeBr₂ obtained from vapour deposition at low temperatures were converted into the crystalline phase by annealing with temperatures up to 800 K. It is found that FeCl₂ and FeBr₂ never convert directly from the non-crystalline to the usual crystalline state. They form an unknown intermediate phase in the temperature range between 60 and 350 K. In contrast, FeF₃ shows direct conversion. This occurs at much higher temperatures (~ 800 K). The existence of an intermediate phase for FeCl₂ and FeBr₂, but not for FeF₃, is discussed in terms of the different structural changes taking place in the recrystallization processes of the non-crystalline iron di-halides.

1. Introduction. — We have previously reported Mössbauer studies of the magnetic properties of non-crystalline samples of the anhydrous iron dihalides FeF₃, FeCl₂, and FeBr₂ [1, 2]. In this paper we report on the observations made as these samples were converted, by annealing at increasing temperatures, into the normal crystalline state. It is shown that this conversion takes place via an intermediate phase of unknown structure.

2. Experimental results. — The samples with thicknesses of the order of ~ 5 000 Å were prepared by deposition from a molecular beam onto a cold (< 10 K) Al substrate. The vapour pressure in the molecular beam oven was usually 10⁻⁵-10⁻⁴ torr. The Mössbauer spectra observed for FeCl₂ and FeBr₂ at temperatures between 100 and 300 K are shown in figures 1 and 2. In our previous studies we have concentrated on the behaviour of the material as it was freshly deposited at low temperatures. It was shown [1, 2] that the Mössbauer spectra obtained differed considerably from those reported for the crystalline material [3, 4, 5, 6], and for one sample (FeF₃) direct evidence could be obtained by X-ray diffraction that the material was non-crystalline. In particular, all three iron di-halides investigated (FeF₃, FeCl₂, and FeBr₂) experience a magnetic transition at ~ 20.5 K and show about the same saturation hyperfine (hf) field of ~ 140 kOe for T → 0 K. Above the magnetic transition temperature a slightly broadened quadrupolar doublet is seen with a splitting about twice as large as those reported for the crystalline samples of the iron di-halides. Between 1.5 K and 30 K the samples could be thermally cycled without any noticeable irreversible effects. At higher temperatures, however, new hf pattern begin to appear which constitute the target of the present investigation.

The Mössbauer spectra for FeCl₂ and FeBr₂ in the temperature range between 100 and 300 K are shown in figures 1 and 2. The quadrupolar splitting labeled A corresponds to the spectrum observed slightly above the magnetic transition temperature (20.5 K) of the non-crystalline material. The quadrupolar doublet seen at high temperatures (> 300 K) is labeled C. It is seen in rather pure form in figure 1, while we have not shown the pure spectrum C for FeBr₂ in figure 2, although it can be obtained rather easily at temperatures around 350 K. From its hf parameters spectrum C can be easily identified as belonging to the normal crystalline phases of FeCl₂ and FeBr₂. Not only is the value of the quadrupole splitting identical to that previously observed for crystalline samples, but also...
spectra obtained when the samples were again cooled to low temperatures (4.2 K) were those characteristic of the crystalline material.

Figure 3 shows such spectra for FeBr₂. It should be kept in mind that crystalline FeBr₂ has a Néel temperature of 11 K. In the magnetic regime the hf field at

The most interesting feature of the spectra in figures 1 and 2 is the appearance of an intermediate phase which is characterized by the quadrupole splitting labeled B. It appears upon heating the non-crystalline samples to about 60 K. From figures 1 and 2 it can be seen that the conversion from phase A to phase B and from phase B to phase C (normal crystalline) takes place over a wide temperature range. The crystalline phase (C) begins to form at 120 K. All three phases can occur simultaneously. The relative intensities of the various phases in a Mössbauer spectrum depend primarily on the maximum annealing temperature and much less on the time spent at constant annealing temperature. This point is demonstrated in the top spectra of figure 2: Keeping the sample for long times at constant temperatures results in a steady, but rather slow growth of phase B, while only a short time heating to more elevated temperatures leads to a much more pronounced increase in phase B. Temperature cycling shows that both phase transitions (A to B, and B to C) are not reversible. In particular, recooling from above 100 K to 4.2 K indicates that the magnetic character of the original non-crystalline phase as described above is suppressed by the presence of phase B. The low temperature spectrum only shows the quadrupole splittings A and B but not the broad magnetic spectrum typical for
pure phase A. Samples of FeF\(_2\) show a slightly different behaviour. First, a much higher temperature (\(\sim 800\) K) is needed for the conversion to the crystalline form (spectrum C): secondly no evidence of an intermediate phase B could be detected. The direct transition from spectrum A to spectrum C is again rather sluggish and irreversible.

Figures 4, 5 and 6 show the temperature variation of the quadrupole interaction in the various phases for three iron di-halides.

**Fig. 4.** — Temperature dependence of the quadrupole splitting for non-crystalline (A), intermediate (B), and crystalline (C) phases of FeCl\(_2\).

**Fig. 5.** — Temperature dependence of the quadrupole splitting for non-crystalline (A), intermediate (B), and crystalline (C) phases of FeBr\(_2\).

**Fig. 6.** — Temperature dependence of the quadrupole splitting for the non-crystalline (A) and crystalline (C) FeF\(_2\).

Samples of FeCl\(_2\) and FeBr\(_2\) were also prepared using a relatively high vapour pressure (\(\sim 10^{-3}\) torr) in the molecular beam oven. In this case the spectra showed only the quadrupole doublet B; that is, the samples were produced directly in the intermediate phase with no evidence for the existence of the non-crystalline phase A which is the result of preparation a lower vapour pressure. For FeF\(_2\), changes in vapour pressure had no effect on the formation of the non-crystalline phase A which also was always produced in pure form.

3. Discussion. — For FeF\(_2\) the quadrupole splitting of the non-crystalline form differs only little from that of the crystalline phase [5, 6] (see Fig. 6). This is consistent with the fact that the crystal is formed of linear molecules of FeF\(_2\) arranged roughly end to end in lines forming more or less discrete layers. The molecular group is dominant and this is expected also in the non-crystalline form. The transition from one form to the other therefore is thought to involve little structural change.

The quadrupole splittings of all three halides in the non-crystalline form are about equal. This suggests a similarity of the non-crystalline structure; presumably more or less discrete molecular units are present in all three cases. The crystal structures of FeCl\(_2\) and FeBr\(_2\) are (in contrast to FeF\(_2\)) not of this form. This is also apparent from the much smaller absolute value of the quadrupole splitting and the relative stronger variation of quadrupole splitting with temperature in the crystalline forms of FeCl\(_2\) and FeBr\(_2\); both facts suggest a much closer spacing of the two lowest crystal field levels. A greater separation of the orbital states could be expected when the axial field strength is more pronounced as it is presumably the case in the non-crystalline form.

For FeCl\(_2\) and FeBr\(_2\) there is then a substantial change in structure between the non-crystalline and the crystalline forms. The appearance of an intermediate phase can thus be understood. Unfortunately little can be said at present about the actual structure of this phase B. It is suggestive, however, that this phase is formed directly from the high pressure vapour in which case it is known from mass spectroscopy that FeCl\(_2\) and FeBr\(_2\) are present to a considerable amount (\(\sim 10^{-20}\) \%) in the form of dimers [8, 9]. In this form the strongly axial structure of the single molecule is less pronounced and the quadrupole splitting would be expected to be smaller. This is indeed the case in the intermediate phase.

One curious feature of the present results is the suppression of the appearance of the magnetic transition in the non-crystalline phase by the presence of only a small amount of the intermediate phase. It is unlikely that the transition itself no longer occurs and it must be suggested that the observation of the magnetic hf spectrum is prevented by some type of relaxation of the spins. It is possible that the intermediate phase is
formed preferentially along lines which effectively divide the layer into small unconnected regions of the non-crystalline phase. Each of these regions could be of a size that would exhibit superparamagnetic effects [10, 11] that is a relaxation of the macroscopic spin of the whole region. It is well known that in such circumstances the effective magnetic field at the nucleus may average to zero within the characteristic time span for Mössbauer spectroscopy.

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