An opening in the anvil assembly having a 20 angle of 45° (Figure 1) permits, the recording of diffraction lines from lattice planes having spacings less than 1 Å when molybdenum $K\alpha$ radiation is used. The diffraction patterns were recorded on Polaroid Type 57-3000 speed sheet film at a specimen-to-film distance of about 4 cm. The film was held in a Polaroid XR7-Land Diffraction Cassette (#57-1) which has a fluorescent intensification screen. The use of this combination of film and cassette permitted patterns to be recorded in periods ranging from 3-15 hr, whereas standard wet film required exposures ranging from 20-50 hr under the same test conditions. Figure 3 shows the difference between diffraction patterns recorded on the Polaroid film and on a high-contrast, high-speed wet x-ray film. These patterns are of an annealed iron powder sample obtained at ambient pressure. Although the wet film gives an obviously sharper and higher contrast pattern, the difference in exposure times, 8 and 44 hours respectively, is large enough to make the use of the wet film undesirable except for circumstances justifying the long exposure time. The advantage of much shorter exposure time combined with the absence of shrinkage correction, ease of handling, and avoidance of cassette movement during a series of high-pressure x-ray diffraction exposures led to the exclusive use of the Polaroid film.

During the course of these experiments, it was found that the manner in which the sample is placed between the two diamonds and the material used to prevent the sample from extruding from between the diamond surfaces at high pressure were important. Figure 4 shows the sequence used to place a sample in the high-pressure x-ray camera. The procedure for mounting powder samples consisted of taping a piece of 0.03 inch-thick sheet lead over the anvil diamond (4a) and placing both the piston and anvil assemblies together with about 5 lb of force (4b). The cavity thus formed had a thin portion of lead covering the anvil diamond face. The center of this thin portion was removed with a needle, and the cavity was filled with the sample powder (4c). The sample was then compacted at light loads between the piston and anvil diamonds (4d). Thus, a uniform sample, surrounded by a thin gasket of lead and excess sample, was obtained.

Application

Phase-pressure information was obtained by taking a series of diffraction patterns as follows: for ambient and increasing pressures at 10 or 20 kbar increments up to a nominal pressure of 80 kbar; and for decreasing pressures, again at 10 or 20 kbar increments, down to ambient pressure. Transformation pressures and percentages of phases present were noted as a function of nominal pressure. The diffraction patterns were then measured and interplanar spacings calculated. The actual pressures were arrived at by substituting our experimentally determined lattice parameters into the equation developed by Mao et al⁴. As an example of the type of information obtained by these procedures, Figure 5 shows four of a series of twelve x-ray diffraction patterns obtained from iron at various pressures. These patterns and results are typical of the two "pressure-phase" determinations which were made on iron. The actual pressure appears in kbar next to each pattern. The arrows indicate increasing or decreasing pressure. The numbered vertical lines running between the patterns indicate the approximate positions of the lines which are identified in the table below the patterns. The lines for the face-centered-cubic (γ) phase are included only for comparative purposes. No fcc material was detected in this sample. It can be seen that at 70 kbar the sample contains a very small amount of the hcp phase (line 3). Prior to reaching this pressure, no hcp was detected. . As Figure 5 shows, after the pressure was decreased to 10 kbar, only a small amount of hcp remained. All of the hcp present at high pressure had transformed to bcc before ambient pressure was reached. Using a series of twelve diffraction patterns, it was possible to determine that the bcc-to-hcp transition began at 70±10 kbar and that the reverse transition, hcp-to-bcc, began at 27±10 kbar.

Problems of Interpretation

There are two problems of major concern in the use of opposed-anvil high-pressure x-ray techniques in the study of high-pressure martensitic transformations: (1) the existence of a pressure gradient from the center to the edge of the sample, and (2) the combination of factors affecting determination of the fraction of the sample that has transformed.

In considering the first of these problems, we find that there are actually two consequences: (a) variation in pressure across the irradiated portion of the sample, and (b) the difficulty of determining the pressure at any given point in the sample.

Myers et $a1^{22}$ found that the variations in pressure across the center 75% of the sample become negligible when the sample diameter-to-thickness ratio is greater than 20:1. This ratio, together with an x-ray beam diameter less than one-half the sample size, can eliminate most of the pressure variation across the irradiated portion of the sample.

Given the existence of a pressure gradient, it is not possible to calculate the pressure at a given point in the irradiated portion of the sample on the basis of applied gas pressure and the ratio of the area of the piston assembly to that of the piston diamond. The value arrived at on this basis will necessarily be lower than the actual value. However, by using known variations of the bcc iron lattice parameter with pressure⁴ as an internal calibrant, it was found that all pressures could be determined within a one sigma error of 10 kbar. This error is small in comparison with the 50 to 130 kbar range of the previously reported transformation pressure for iron^{1-5,7}. Most of the 10 kbar error was introduced during diffraction pattern measurement.

For work with metals having a cubic crystal structure for which lattic parameter-pressure information is not available, an accurate estimate of the change in lattice parameter with pressure can be obtained from volume-pressure data such as those presented by Bridgman¹³. An alternate method of determining pressure is to mix into the sample a material, such as NaCl, whose pressure-lattice parameter characteristics are well known²⁻⁵.

The second of the two major problems in high-pressure x-ray diffraction work stems from the difficulties involved in estimating the precentage of each particular phase. Since the percentage of each phase is estimated on the basis of relative diffraction line intensity and the structure factor, any factors, in particular preferred orientation and line overlap, that interfere with the correct determination of line intensity will also lead to an incorrect estimate. Preferred orientation is usually present in cold-rolled metals even after annealing. Annealed powdered metals often develop a preferred orientation during deformation during the compaction at low pressures. This preferred orientation is usually not as severe as that found in cold-rolled and annealed foils.

Diffraction line overlap is shown in Figure 6, which is a schematic repressentation of interplanar spacing for the three structures of iron. As this figure illustrates, the different structures of a metal can give diffraction lines at the same points on the diffraction pattern. Overlapping lines occur at about 2.05 Å, 1.26 Å, and 1.08 Å. Because of these overlapping lines, only a few of the diffraction lines can be used in the estimation of phase percent. A minor point illustrated in the figure is that the (100), (102), and (200) lines of the hexagonal closepacked structure are only weakly diffracting planes.

SUMMARY

To sum up, the existing techniques for the use of a commercially available opposed-anvil high pressure x-ray diffraction unit in the study of high-pressure transformations in iron alloys were modified to permit one to more closely monitor these phase changes as a function of pressure. Modifications affected specifically: beam collimation, beam and diamond alignment, and mounting procedures. The use of Polaroid film to record the diffraction patterns was introduced.

Methods of minimizing pressure=determination difficulties associated with the existence of the pressure gradient across the irradiated portion of the sample were described. The procedure for using diffraction lines to determine phase fractions was discussed.

Employing the modified techniques, it was demonstrated that the transformation of bcc iron to the hcp phase occurs at 70 kbar.

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