12-Amorphous, Imperfectly Ordered and Quasi-periodic Materials

PS-12.01.09 REPEATED ELBOW TWINNING OF Fe₄Al₁₃ SHOWING DIFFRACTION PATTERNS WITH DECAGONAL PSEUDOSYMMETRY. By M. Ellner, Max-Planck-Institut für Metallforschung, Institut für Werkstoffwissenschaft, Seestr. 75, D-7000 Stuttgart 10, FR Germany

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 $\mathrm{Fe_4Al}_{13}$ is formed by peritectic reaction at 1430 K from Fe_2Al_5 and the melt, C2/m, mC101, a = 15.489, b = 8.0831, c = 12.476 Å, β = 107.72° (Black, Acta Cryst., 1955, 8, 43-48 and 175-182). Multiple twinning was observed not only in Fe,Al13, but also in the isotypical compound $\mathrm{Ru}_{\Delta}\mathrm{Al}_{13}$ (Edshammar, Acta Chem. Scand., 1965, 2124-2130) as well as in the ternary isotypical phase Ni₃Zn₆Ga/ (Rajasekharan, Sarah & Schubert, Z. Metallkd., 1982, 73, 526-529). Twins and planar defects were recently observed in Fe,Al, by means of HREM (Tsuchimori, Ishimasa & Fukano, Phil. Mag. B, 1992, 66, 89-108). In the present study $\mathrm{Fe_4^{Al}_{13}}$ was investigated in the concentration range from $x_{A1} = 0.75$ up to $x_{A1} = 0.77$ by means of X-ray diffraction, scanning electron and optical microscopy. In contrast to the neighbouring phase $\mathrm{Fe_2Al_5}$ which does not show any polymorphic transformation, metallographic investigation of the structure of $\mathrm{Fe_4Al_{13}}$ yielded an indication for a phase transformation. Hollow prismatical lamellae were observed in the $\mathrm{Fe_4Al_{13}}$ alloys by the scanning-electron-microscopy investigations. They were interpreted as composite crystals formed by repeated elbow twins with the twin plane (100), (001) and ($\overline{2}01$). An atomistic model showing pentagonal pseudosymmetry is discussed in direct space for the repeated elbow twins of $\operatorname{Fe_4Al}_{13}$. Guinier photographs of splat-cooled alloys $Fe_{100-x}Al_x$ (x = 0.75...0.77) showed diffraction lines of a new phase homoeotypic to Fe_4Al_{13} , oB50, a = 7.733(3), b = 4.021(2), c = 23.688(9) Å, $c/a \approx tan 72$ °. The relationship between this oB = q unit cell and the monoclinic unit cell of Fe_4Al_{13} is $(a,b,c)_q =$ $(a,b,c)_m(1/2,0,1/2; 0,1/2,0;0,0,2)$, i.e. $a_q = a_m/2$ etc. The low-temperature monoclinic structure ${}^{\mathrm{Fe}}_4{}^{\mathrm{Al}}_{13}$ is a displacive variant of the high-temperature orthorhombic phase Fe₄Al₁₃(h). A twinning model is discussed in reciprocal space for structures with orthorhombic face centred Bravais lattices oB (oA, oC) having an axial ratio c/a = tan 72°, (b/c, b/a = tan 72°), i.e. the ratio of the interplanar spacings d_{002}/d_{101} = $(1 + \sqrt{5})/2 = \tau$ $(d_{020}/d_{011}, d_{020}/d_{110} = \tau)$ (golden mean). Composite crystals (repeated elbow twins) of these oB-structures which have the common direction $[010]_q = [010]_m$ and showing the twin plane $\{101\}_q$ i.e. twin planes $\{001\}_m$ and $\{\overline{2}01\}_{m}$ yield diffraction patterns $(h01)_{q} = (h01)_{m}$ of decagonal pseudosymmetry; cf. electron diffraction pattern of twinned $\operatorname{Fe_4Al}_{13}$ (Fung, Zou & Yang, Phil. Mag. Lett., 1987, 55, 27-32, Fig. e).

PS-12.01.10 X-RAY DIFFRACTION PATTERNS OF ONE DIMENSIONAL QUASICRYSTALS. By H. Terauchi, M. Ishida, N. Sano, H. Kasatani and K. Sakaue, Department of Physics, Kwansei-Gakuin University, Nishinomiya 662, Japan.

It is of great interest to study one-dimensional (1D) quasicrystals in which quasiperiodic orders may be established. The most typical 1D quasiperiodic order is found in the lattice with a second-order Fibonacci sequence, $S_T = S_{T-1} + S_{T-2}$. The lattice is fabricated by molecular beam epitaxy techniques with semiconductor materials. The diffraction pattern of the lattice shows many satellite peaks assigned by the golden mean τ related to a diagonal value of a regular pentagon (H. Terauchi et al., J. Phys. Soc. Jpn., 1988, 57, 2416). Of course, quasiperiodic lattices with third-order Fibonacci sequences such as $S_T = 2S_{T-1} + S_{T-2} \cdot S_{T-3}$ are fabricated and here the diffraction peaks are assigned by two diagonal values of a regular heptagon (H. Terauchi et al., J. Phys. Soc. Jpn., 1990, 59, 405).

The other 1D quasiperiodic orders are seen in a Thue-Morse lattice (F. Axel and H. Terauchi, Phys. Rev. Lett., 1991, <u>66</u>, 2223) and a dragon lattice (H. Terauchi *et al.*, J. Phys. Soc. Jpn., 1992, <u>61</u>, 1141), where the sequences are given by $S_r = S_{r-1} + S_{r-1}^*$ and $S_r = S_{r-1} + X + S_{r-1}^*$, respectively. These 1D quasiperiodic lattices give the curious diffraction patterns.

patterns.

The marked differences of the diffraction patterns in the above four quasiperiodic orders are discussed, in terms of the correlation lengths as well as the selfsimilarities.

PS-12.01.11 GEOMETRICAL MODE ANALYSIS OF INCOMMENSURATE RE-ENTRANT MODULATIONS: (C₃H₇NH₃)₂MCl₄ M=Cu, Mn, Cd. By N.Achiwa*, H.Ikeda and Y.Yamauchi, Department of Physics, Kyushu University, Fukuoka 812, Japan.

We have developed a geometrical method to analyse molecular modulation modes in incommensurate crystal structures. From the three dimensional locus of each modulated atom which are pulled back to the original unit cell, that of a special mode, for example translational molecular displacement is subtracted. Then, the remainder figure suggests the next main mode, for example, rotational molecular mode. So we can find successively miner modes orthogonal to the subtracted modes by the trial and error method. The family compounds (C3H7NH3)2MCl4 (M=Cu, Mn, Cd) show the following phase sequence: $\alpha \to \beta \to \gamma \to \delta \to \epsilon \to \zeta$, with the two incommensurate ϵ and γ -phases. These incommensurate structures were extensively studied using superspace symmetry. We have applied the new geometrical method to find the displacive modulation modes in the γ and ϵ phases. Figure 1 and 2 show loci of modulated atoms of (C₃H₇NH₃)₂CuCl₄ in γ (superspace group, Pbca:($\alpha00$)ss0, Doudin and Chapuis, 1990 B46 180.) and ϵ (superspace group, $BP2_1/b11:(0,0.5,\xi)1\overline{1}$, Ikeda and Achiwa) phases. The main modes in the γ and ϵ phases are a translational modulation along the c axis and a twisted intra-molecule mode of propyl ammonium base, respectively, though in both phases each orbit of atomic modulation consists of multi-molecular modes. Finally, the amplitude of each molecular mode can be refined by least squares fitting of satellite structure factors.