07.1-8 INTERGROHTH IN CRYSTALS OF ZSM-5 ZEOLITE, By D.G. Hay, H. Jaeger and K. Wilshier, CSIRD Division of Materials Science and Technology, Clayton, Australia.

The systematic intergrowth observed in ZSM-5 zeolite crystals has been examined with transmission and scanning electron microscopy. The most frequent form of intergrowth is where adjoining crystals are rotated $90^{\circ}$ about a common [001] axis, with intergrowth always initiated on growing [.010] faces. Sections nomal to [001] and [100] of the main crystal show the boundaries are curved and they can be modelled as a series of steps. The model intergrowth boundaries are between (010)/(100) and (001)/(001) planes (where the subscripts refer to planes in the main crystal and intergrowth respectively). A second form of intergrowth, associated with inclusions of impurities in the crystals, is seen as ramps on the (100) faces although the intergrowth is often initiated well below the crystal surface. Observation of both forms of intergrowths has led us to propose the likely building units involved in the formation of well defined crystals.
07.1-9 CRYSTAL GROWTH BY METASTABLE
NUCLEATION. By B.N. Mehrotra, Department of Physics, Sardar Patel University, Vallabh Vidyanagar 388120, Gujarat, India.
single orystals of sodium sulphate III, a modification which is stable only in the temperature range 200 to 228 C, were obtained by metastable nucleation frog aqueous solution of sodium sulphate at $50^{\circ} \mathrm{C}$, and this led to its structure determination (Zeit. Krist., Bd $155(3 / 4) 159 / 163)$ by the author.

Similsrly,crystals of sodium heptahyarate are also grown by metastable nucleation from the aqueous solution at $0^{\circ} \mathrm{C}$. The powder photograph of this substance is obtained on Philips diffractometer using Cuke radiation, and the results are given in the table below :
Table: d-values and relative intensities of sodium sulphate heptahydrate.
S.N. d( $\AA$ ) Int. S.N. d(A) Int. S.N. d(A) Int.

| 1 | 5.48 | 99 | 8 | 2.750 | 40 | 15 | 2.040 | 70 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| 2 | 5.41 | 50 | 9 | 2.697 | 30 | 16 | 1.782 | 15 |
| 3 | 5.27 | 70 | 10 | 2.499 | 25 | 17 | 1.764 | 40 |
| 4 | 4.018 | 50 | 11 | 2.427 | 15 | 18 | 1.631 | 68 |
| 5 | 3.516 | 40 | 12 | 2.255 | 17 | 19 | 1.616 | 35 |
| 6 | 3.030 | 35 | 13 | 2.220 | 15 | 20 | 1.419 | 18 |
| 7 | 2.838 | 40 | 14 | 2.092 | 20 | 21 | 1.369 | 28 |

07.1-10 ON THF ORTENTATION OF $\mathrm{K}_{2} \mathrm{Cr}_{2} \mathrm{O}_{7}$ CRYSTALS GROWING FROM AQUROUS SOIUTIONS BY SELF-HTUCLEATION ON RHE CRYSTALIIZRR BOTMOM By M. Szurgot and K. Sangwal, Institute of Physics, Technical University of zódź, Wólczaŕska 219, 93005 Iódé, Poland.

The growth of single crystals on isotropic or anisotropic substrates and of thin crystalline films of particular crystallographic orientation requires a knowledge of the effect of various factors on $i t$. The orientation of a crystal growing from solutions on the bottom of a crystallizer is detemined by the surfece in contact with the botton. In this work are described and discussed the experimental results on 'contact' planes of $\mathrm{K}_{2} \mathrm{Cr}_{2} \mathrm{O}_{7}$ (KBC) srowing by self-nucleation from aqueous soIutions a.t $5-97^{\circ} \mathrm{C}$.
It was observed that the planes of contact with the bottom of the growth vessel are always habit plancs of the crystal and are affected by both external and internal factors. The external factor affecting the contact planes is growth temperature, while the intermal dominating factor determining the probability $P$ of contact planes is reticular density d of the atomic planes. It was also found that the greater the $d$, the greater the Value of $P_{\text {: }}$ and that the dependence of $P$ on $d$ is exponential. Our experiments with a large MBC crystal on the probability of its falling in air showed that the site for nucleation is, in all probability, the bulk solution rather than the crystallizer bottom.
Uur conclusions are also valid for the data reported on other inorganic and organic crystals amd thin films.

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