### m30.p13 New 1D, 2D and 3D chiral MOF's based on cyclohexane-1,2-diamine

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Cyanide-bridged bimetallic systems, prepared from assembling cyanometallates and transition metal complexes, have been shown to exhibit fascinating structures with interesting magnetic, electrochemical, magneto-optical and zeolite-like properties [1]. The design and synthesis of nanoporous metal-organic frameworks (MOF's), mimicking zeolities, is drawing a great deal of attention due to the application of these materials as catalysts, adsorbents and ion exchangers. Chiral structures are a new target for the lattice architecture in the field of magnetic materials. There are only a very few examples where cyclohexane-1,2-diamine has been incorporated into cyanide-containing complexes resulting in the formation of ferromagnets [2], [3], [4]. We will present some new one-, two- and three-dimensional chiral cyanide-bridged bimetallic MOF's.  $[M1(chxn)_{n}]_{n}[M2(CN)_{6}]_{m}$  (where M1 = Cu, Ni; M2 = Ru, Cr, Co; n = 2, 3; m = 1, 2; chxn = trans-1S,2S or 1R,2R-cyclohexane-1,2-diamine), and their structural features, magnetic and zeolite-like properties will be discussed.

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## m30.p14

#### Atomic and domain structure of the low temperature phase of barium metagermanate

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The crystal structure of the low temperature form of barium metagermanate (BaGeO<sub>3</sub>) has been determined from laboratory X-ray powder diffraction data collected at 25.5(5) °C. The compound is monoclinic with space group C2/c and unit cell parameters of a = 13.1877(2) Å, b = 7.6222(1) Å, c = 11.7190(1) Å,  $\beta = 112.295(1)$ °, V = 1089.9(1) Å<sup>3</sup> and Z = 12. The structure was found to consist of alternating layers of barium atoms and three-membered [GeO<sub>3</sub>]<sub>3</sub>-rings showing a twofold positional disorder. Apart from the disorder the structure is very similar to the one of SrGeO<sub>3</sub> [1]; both are six-layer structures build from the same type of layers. The same structure type also appears in pseudo-wollastonite, forming four- or six-layer polytypes [2]. Careful studies of possible structural variations using simulated annealing [3] did not reveal any deviation from symmetry C2/c.

The disorder of the ternary rings can be summarized as follows: ternary  $[GeO_3]_3$ -rings appear in only one out of two possible orientations and occupy only 2/3 of the positions compatible with this orientation. Following [2], for six-layered structures of this type there is only one distinct polytype with the inherent symmetry C2/c. There are, however, two non-congruent enantiomorphs of this structure. The disorder observed in BaGeO<sub>3</sub> can be described as the simultaneous appearance of both possibilities.

Electron microscopy reveals in SAED patterns the existence of twins; the twin law can be described as a mirror plane normal to **a**. Diffuse scattering within the lattice rods parallel  $c^*$  indicates disorder within the layers of the structure. HRTEM shows fully ordered domains, which are separated by narrow disordered regions; the domain borders observed are normal to  $c^*$ . Larger disordered regions are observed as well.

Based on these observations the disorder in the structure can be addressed as stacking fault disorder, a behaviour also found in polytypes of pseudo-wollastonite [2]. The stacking, however, is not random but can be rationalized by a twinning mechanism mapping the enantiomorphic polytypes onto each other.

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