FA5-MS01-P30

Structural Study of Negative Thermal Expansion in Yttrium and Rare Earth Molybdates. <u>Cristina</u> <u>González-Silgo</u>^a, Manuel E. Torres^b, Javier González-Platas^{a,c}, Javier Campo^d. V. Sánchez Fajardo^b, Diego Lozano-Gorrín^c, Candelaria Guzmán-Afonso^b, Nanci Sabalisck^b. ^aDepartamento de Fisica Fundamental II. Universidad de La Laguna. España. ^bDepartamento de Física Básica. Universidad de La Laguna, España. ^cServicios Generales de Apoyo a la Investigación. Universidad de La Laguna. España. ^dInstituto de Materiales de Aragón. CSIC. España. E-mail: metorres@ull.es

Molybdates with formula $A_2(MoO_4)_3$ show a very intense negative thermal expansion (NTE) above 450K, when A=Y, Ho-Lu with $Sc_2(WO_4)_3$ -type structure (Pbcn) [1]. There is not a detailed structural investigation at lower temperatures because they are hygroscopic [2]. Three samples $Y_2(MoO_4)_3$, $Er_2(MoO_4)_3$ and $Lu_2(MoO_4)_3$ have been prepared by the conventional solid-state synthesis with preheated oxides. We have analysed the diffraction data of dehydrated compounds (A= Y, Er, Yb and Lu) from an ILL experiment (5-24-360) at the D2B powder diffractometer. We have found that all compounds show negative thermal expansion (NTE) behaviour in the temperature range of 150-400K. All the structures have been well refined by multipattern procedure [3], through X-ray powder diffraction data too, obtained on the laboratory equipment.

Given an accurate and precise crystal structure we have measured the exact scope polyhedral distortion with the temperature and ionic radii and the role of the transverse vibration of binding A-O-Mo oxygen atoms in the structure, since several authors have related such vibrations with the NTE behavior [4]. Also, we have studied the structural changes depending on the temperature and the ionic radii using the symmetry mode amplitude formalism [5], by comparing the distortions of lower temperatures respect to those at the highest temperature.

[1] Imanaka, N. et al. Chem. Mat., 2000, 12, 1910. [2] Manrinkovic,
B. A., Jardim, P. M. De Avillez, R. R., Rizzo, F. Solid State Sci.,
2005, 7, 14. [3] FullProf Suite: http://www.ill.eu/sites/fullprof/ [4]
Ravidran, T.R., Arora, A. K., Mary, T. A. Phys. Rev. Letter, 2000,
84, 3879. [5] Bilbao Crystallographic Server: http://www.cryst.ehu.es/

Keywords: molybdates; neutron and X-ray diffractometry; thermal expantion

FA5-MS01-P31

Crystal Structures and Properties of Novel Inorganic-Organic Materials. <u>Wieslaw Łasocha^{a,b}</u>, Piotr Putaj^a, Alicja Rafalska-Łasocha^a, Bartlomiej Gaweł^a, Wojciech Nitek^a.Faculty of Chemistry, Jagiellonian ^aUniversity, ul. Ingardena 3, 30-060 Kraków, Poland.^bInstitute of Catalysis and Surface Chemistry PAS, ul. Niezapominajek 8, 30-239 Kraków, Poland. E-mail: <u>lasocha@chemia.uj.edu.pl</u>

25th European Crystallographic Meeting, ECM 25, İstanbul, 2009 *Acta Cryst.* (2009). A**65**, s 329 The synthesis of new hybrid inorganic-organic materials is a growing field of research. The main purpose of designing such new structural types is to combine the functional properties of organic and inorganic structures to create materials with desired/tailored properties for new challenging applications. Such materials can be used interchangeably with zeolites, as molecular reactors or for the storage of various gases.

Recently, we have synthesised and investigated the complexes of $MeX_2(Me=Cd \text{ or } Zn, X=Cl \text{ or } Br)$ with different amines.

Coordination polymers (1-3) were obtained when ethylenediamine or triethyleneamine were used in the synthesis. These compounds are built of straight, inorganic chains of "Cd"X₂"Cd" and organic groups. In the case of ethylenediamine inorganic chains were joined by diamine molecules forming two-dimensional layers (1,2), whereas in the case of triethyleneamine (a bigger molecule, one nitrogen atom) double inorganic chains with the formula $Cd_2Cl_5(H_2O)$ were formed (3). The chains in this compound are isolated from each other by organic groups which are connected to the chains by H-bonds only. The same type of structures has been recently described in the literature [1].

Isolated, butterfly-like moieties (4) were obtained when $ZnCl_2$ reacted with 3,5-dimethylaniline (a big monovalent ligand).

Crystallographic data for the investigated compounds (space group, lattice parameters [Å], V [Å], Z)

(1) $[Cd(C_2H_8N_2)Cl_2]$, Pbam (55), 9.9032(3), 7.819(4), 4.0692(5), 315.09(17), 2.

(2) $[Cd(C_2H_8N_2)Br_2]$, Pbam (57), 10.415(2), 7.8653(5), 4.2085(8), 344.75(10), 2.

(3) $[Cd_2Cl_5(H_2O) ((C_2H_5)_3NH)] P2_1/n (14), 11.271(1), 19.089(3), 7.648(2), 96.93(3), 1633.9(4), 4.$

(4) $[ZnCl_2 {(CH_3)_2C_6H_3NH_2}_2] C2/c$ (15), 30.029(5), 4.8990(8), 12.067(2), 103.52(2), 1726.0(5), 4.

The crystal structures of compounds 1 and 2 were solved by powder diffraction methods, whereas the structures of 3 and 4 were solved by single crystal methods.

[1] Thorn A, Willett R.D, Twamley B, Cryst. Grow and Design, 2006, 6(5),1134.

Keywords: crystal engineering; inorganic-organic coordination polymers; powder diffraction

FA5-MS01-P32

Ba₂Ni₃ - A New Phase in the Binary System Ba-Ni. <u>Peter Höhn</u>^a, Stefan Hoffmann^a, Rüdiger Kniep^a. *^aMPI-CPfS, Dresden, Germany.* E-mail: <u>hoehn@cpfs.mpg.de</u>

For investigation of the formation of intermediate phases in the ternary (and quaternary) system Ba-Ni-N(-C) pelletized mixtures of the elements and binary alkaline-earth nitrides were preferentially used [1]. In order to prevent impurities of elemental Ni in the samples, alternate routes of preparation via intermetallic precursors were evaluated.

The binary system Ba-Ni was first investigated in 1966 [2]. According to this study, no intermediate phases exist [3]. Our recent investigations [4] in the system Sr-Ni did