covering, for example, the utilisation and protection of ground water, the foundations of skyscrapers, the sealing of waste deposits, canalisation systems, railway and road construction, and the redevelopment of waterways and the soil. The determination of the crystal structure of senaite has progressed to the point where a trial model consisting of all atoms has been defined. Plumboferrite has been examined and found to have the magnetoplumbite substructure. The structure of CoGeO3 has been refined in detail and the occurrence of polytypism in this compound is readily explained by the approximation to closest packing.

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Keywords: General Geoscience, Inorganic crystallography, Geosciences

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Thickness dependent crystallographic transition in Fe/ Ni superlattices

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The 3d transition metals have historically got a huge attention due to both interesting physical properties as well as the applications of systems containing such elements. FeNi alloys, in all concentration ranges, are of particular interest due to industrial applications with emphasis on the so called Invar [1]. While body centered cubic (bcc) Fe is ferromagnetic at room temperature, face centered cubic (fcc) Fe can posses rich variety of magnetic phases. Mossbauer spectroscopy, due to its sensitivity to magnetic and structural environment, is a powerful technique for studying such systems and also because it can isolate Fe magnetism. In contrast to previous studies, where a thick Ni layer has been used as buffer layer and then a thin Fe layer has been establish as fcc structure on that, here another approach has been tried. By starting with higher thickness of Fe than Ni, a bcc structure is established. By increasing the Ni thickness, the structure should transit gradually from bcc to fcc. This transition is however smooth, i.e. there will be coexistence of both faces. Several sample series of Fe/Ni superlattices with varied x ML of Fe and y ML of Ni were grown by dc magnetron sputtering method onto MgO substrate. In the first sample series the ratio x/y was kept constant, while the bilayer thickness was changed. In the second sample series, the ratio x/y was also changed. The study of the first series showed that the bcc structure persisted and was almost insensitive to changes of the interface density. The result from the study of the second sample series, where the ratio x/y was changed, i.e. thickness of the Ni layer increased while the thickness of Fe layer was unchanged, showed a gradual transition from bcc to fcc.

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Keywords: Mossbauer Spectroscopy, Fe/Ni Superlattices, Magnetism

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Combined use of crystallography and mass spectrometry for ligand characterization and drug design

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AliX has developed FAMASS (Fragment Analysis by MASs Spectrometry), a proprietary approach for structure-based drug discovery combining X-ray crystallography and non-covalent mass spectrometry. We routinely use non-covalent electrospray ionization mass spectrometry to check the purity and the functionality of proteins used in [protein-ligand] complex crystallization, and also to screen ligands for binding to given targets. This is especially interesting when a functional assay is not available, as in the case of orphan nuclear receptors. Non-covalent mass spectrometry is an efficient secondary screening method to discard false positives from a high-throughput primary screening. We have now extended this approach to fragment-based drug discovery. Fragments selected by in silico screening are validated for physical binding to the target by non-covalent mass spectrometry. Validated fragments are then cocrystallized with the target and their mode of binding elucidated by crystallography, paving the way toward the design of new scaffolds.

Keywords: protein-ligand complex, mass spectrometry, fragment-based drug discovery

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Teaching how to simply replace the independent atom model - the example of Bergenin

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In four years the crystallographic community will celebrate 100 years of X-ray diffraction. While in the first 50 years the method was applicable only in exceptional cases, the developments in the second half of the last century increased the number of published X-ray structures dramatically, so that the number of CCDC entries is quickly approaching half a million. Nevertheless, in almost all cases the independent atom model (IAM) was applied that uses spherical scattering factors which do not allow a proper description of chemical bonding. To replace the IAM with an aspherical scattering model we recently introduced the concept of Invarioms, generated an Invariom library and corresponding software for a straightforward application [1]. Since the needed X-ray and computer equipment is far from expensive up-to-date standard and since we will show that the use of the Invariom model can easily be taught, the aspherical Invariom model is best suited to be used by groups also in less developed countries. This is demonstrated by the example of Bergenin, a compound of traditional Asian medicine, which is known for anti-HIV and other biological activities. On a more than 20 years old diffractometer a low resolution (sin(theta)/lambda= 0.60 A-1) data set was measured at room temperature that was properly suited to obtain an improved molecular geometry compared to the IAM and a variety

of electronic properties. Since the simple experimental equipment mentioned above is probably available world wide and since the software is free of charge an unlimited use of the procedure we will teach here, seems possible.

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Properties of BiFeO₃ and Bi_{0.7}La_{0.3}FeO₃ nanopowders obtained by mechanosynthesis

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The search for materials exhibiting both electric and magnetic long-range order at room temperature, driven by the perspective applications in information storage and sensors, renewed the interest in BiFeO₃. At room temperature BiFeO₃ has rhombohedrally distorted cubic perovskite cell (R3c) [1] and the antiferromagnetic properties are related to G-type ordering with a cycloid modulation (62 nm) apparent down to 5 K [2]. Due to the due to long-range spin arrangement one may expect the magnetic properties of BiFeO₃ to be size-dependent. We expected that the decrease in the particle size below the periodicity of cycloid modulation will modify considerably the magnetic properties. Recently nanosize powders (obtained by glycine combustion synthesis) were reported to exhibit ferromagnetic hysteresis loops at room temperature [3]. We studied the magnetic properties of BiFeO₃ and Bi_{0.7}La_{0.3}FeO₃ nanosize powders produced by mechanical synthesis from respective oxides at room temperature. In Bi_{0.7}La_{0.3}FeO₃ the ferroelectric Curie temperature T_C is close to the magnetic Néel temperature ($T_N \approx 640$ K) [4], whereas $T_C \approx 1100$ K for BiFeO₃. The nanopowders were prepared by mechanically triggered room temperature synthesis from commercially available oxides Bi₂O₃, Fe₂O₃, La₂O₃ (99% purity from Aldrich) in a SPEX 8000 Mixer Mill. The batch contained \sim 6g of the oxide powders in stoichiometric ratio. The weight ratio of the stainless steel balls to the oxidesamounts to 2:1. The process was controlled by XRD and the powder was characterized by TEM. Magnetic properties were measured with Oxford Instruments Ltd. MagLab 2000 magnetometer in the temperature range 2 - 350 K. The temperature variation of the magnetization was measured in zero field cooling (ZFC) and field cooling (FC) experiments, whereas the magnetization loops were recorded at constant temperature with magnetic field applied step by step. The powders obtained after 120 h milling of the oxides were found to exhibit rhombohedrally distorted perovskite structure and TEM observations show that the powder forms irregular 100-150 nm loosely packed agglomerates composed of oval 10-30 nm grains. The nanograins exhibit core-shell structure with crystalline core surrounded by a disordered shell (1-2 nm thick). Thermal treatment (1 h at 500°C) results in a disappearance of the shell and an increase in the grain size to 40-50 nm. BiFeO3 and Bi_{0.7}La_{0.3}FeO₃ nanopowders with core-shell structure of the grains were found to exhibit fielddependent $1/\chi(T)$ anomaly at \sim 8K in ZFC measurements. A continuous increase in values was observed on FC-measurements down to 2K. For annealed BiFeO3 powder (grains without disordered shell) a broad $1/\chi(T)$ anomaly at ~94K was found in ZFC experiments; for Bi_{0.7}La_{0.3}FeO the anomaly was found to be shifted towards lower temperature. A continuous increase in χ value was observed in FC experiments down to 2K, whereas for Bi_{0.7}La_{0.3} FeO₃ the χ values slightly change with temperature. To check whether the disturbance in the long-range cycloid spin arrangement in BiFeO3 type multiferroics can result in an increase in magnetization and in a nonlinear field dependence of magnetization we studied the field dependences of magnetization M(H) for the prepared nanopowders. Magnetic hysteresis loops were observed at low temperatures for BiFeO3 and Bi0.7La0.3FeO3 nanograins both as-prepared and annealed. The magnetization loops were not really saturated which points to basic antiferromagnetic nature. We found that the annealing decreases considerably the magnetization. Moreover, the coercive field of Bi_{0.7}La_{0.3}FeO₃ was found to be higher than that of BiFeO₃. The improvement of magnetization in nanosize particles of BiFeO3 and Bi_{0.7}La_{0.3}FeO₃ obtained by mechanical activation of respective oxides we would like to relate both to:

i) An incomplete rotation of the spins along the direction of the wave vector (62nm modulation length of the cycloid),

ii) An increase in the spin canting due to the strain extended by the grain shell (which gives rise to ferromagnetism).

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