molecules assumed as a protection again spontaneous crystallization in organisms? Acknowledgements: The support by the Grant Agency of AS CR under grant no. 1A10010915 is gratefully acknowledged.


Keywords: molecular packing, space-group frequencies, crystallization

MS17.P21  

Stacking interactions between phenanthroline ligands in square-planar complexes  
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Noncovalent interactions of π-systems, including stacking interactions, have been extensively studied in recent years. Stacking interactions are generally studied between aromatic organic molecules or fragments. However, we showed that other planar molecules and fragments can be also involved in stacking interactions. Stacking interactions between chelate and C₆- aromatic rings were recognized by analyzing the data in crystal structures of square-planar transition metal complexes from the CSD. In the crystal structures there are mutual slipped-parallel orientation of these rings, similar to the orientation of two benzene rings. Recently we showed the existence of chelate-chelate stacking interactions in crystal structures from CSD [1]. Geometrical analysis of stacking interactions with aromatic nitrogen-containing ligands show that nitrogen-containing aromatic rings usually form slipped parallel alignment where most of the ring-plane area overlaps [2].

Phenanthroline (1,10-phenanthroline-N,N') (phen) molecule coordinating to a metal ion forms large planar system of four rings: two pyridine fragments, one C₆-ring and one chelate rings. This planar system has propensity to form stacking interactions with the π -system of various aromatic groups. Tendency for stacking interactions is important for using these complexes in biochemistry, supramolecular and medicinal chemistry.

To understand better stacking interactions of phen complexes, here we analyze the geometry of stacking interactions between phen square planar metal complexes in crystal structures from the CSD.

In the Cambridge Structural Database (CSD) 61 structures with 172 stacking interactions of phen square-planar complexes were found [3]. The results of crystallographic analysis show that in most of the structures two interacting ligands are oriented “head-to-tail”, with the large area of phen ligand involved in the overlap. Phen complexes show large range of different overlap geometries in stacking interactions, however, short metal-metal distances were not observed. Stacking interactions of phen square-planar complexes form chains and dimers in crystal structures.

MS18.P01  

Structure and magnetic properties of nanoparticles Ni₈-Zn₆-Cr₄-Fe₉-O₁₈  
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Nanostructure ferrite of composition Ni₈-Zn₆-Cr₄-Fe₉-O₁₈ was prepared by sol-gel, co-precipitation, citrate-gel, flash and oxalate precursor methods. Structural and micro structural analysis of all the samples were carried out ( XRD, IR, TEM, SEM and AFM ). The saturation magnetization at room temperature was also studied. Lattice constant and particle size were determined by different methods. The average values of the crystal size for different method were in nanometric range (5 - 75 nm). It was observed that the lattice parameter of cubic spinel was not varied and the positions of the tetrahedral and the octahedral band are not shifted which give the indication that the cation distribution are the same. So, the results of magnetic hysteresis were explained in terms of the relation between grain size and magnetic properties in addition to surface randomizing effect in nanocrystals leading to superparamagnetism state.

Keywords: nanocrystals, synthesis, ferrite.

MS18.P02  

Identification of single photoswitchable molecules in nanopores of silica xerogels using neutron powder diffraction  
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Single photo-switchable molecules of (CN,H₃)₃Fe(CN)₅(NO) (GuNP) are embedded into nanopores of a SiO₂ xerogel. We show that it is possible to identify the structural motif (fingerprint) of the embedded complex by analyzing neutron powder diffraction data in a limited Q range (Q<37 nm⁻¹) using the Debye approach. The structural study reveals that the pores are occupied by GuNP-monomers with a fill factor of 60-80%. The mutual arrangement of the anion and cation in the GuNP monomer is slightly changed (~ 1% elongation) while the bond lengths within anion and cation are changed by less than 0.2% with respect to the single crystalline form of GuNP.

This example demonstrates that the structure of single molecules in nanopores and their distance to nearest neighbors of the host network can be analyzed using a standard neutron powder diffractometer with limited Q-range such as DMC@SINQ. From the analysis of the small angle range one obtains an estimate for the average pore size and the fill factor of pores with embedded complexes. The distance between atoms of the host network and the guest complexes can be determined from the intermediate scattering range. Assuming a sufficient scattering contrast between host and guest this method can be extended also to the x-ray case, such that these experiments can be performed on laboratory