CHARGE, SPIN AND MOMENTUM DENSITY

Mn sites. Thus, there is a considerable orbital component to the magnetic moment. Bader topological analysis shows an absence of Mn-Mn bonding and the magnetic ordering is via super-exchange through the oxygen bridges. Formal electron counting suggests mixed valence Mn sites, but this is not supported by the Bader atomic charges, Mn(1)=+0.11, Mn(2)=+0.17. The topological measures show the dominant metal-ligand interactions to be electrostatic, and a simple exponential correlation is derived between Mn-O bond lengths and the values of $\nabla^2 \rho$ at the bond critical points.

 Poulsen R. D., Bentien A., Chevalier M., Iversen, B. B., J. Am. Chem. Soc., 2005, submitted.
Poulsen R. D., Bentien, A., Graber T., Iversen B. B., Acta Cryst., 2004, A60, 382.

Keywords: charge density, physical properties, metallorganic framework

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Charge Density of 1-phenylpropane-1,2,3-triyl Trinitrate

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The organic esters of nitric acid, the nitrates, are a class of products widely used for the treatment of a number of cardiovascular diseases. The prototype of nitrates is glyceryl trinitrate (GTN, nitroglycerin) It is an oil endowed with potent vasodilating and explosive properties. The major limit in its therapeutical use as vasodilator is an early development of the tolerance. 1-Phenylpropane-1,2,3-triyl trinitrate, a phenyl substituted GTN is characterised by having a high lipophilicity and interesting *in vitro* vasodilating profiles assessed on rat aorta strips pre-exposed to GTN.

A multipole analysis has been applied to the low temperature Xray intensities of *threo* form of 1-Phenylpropane-1,2,3-triyl trinitrate (mp 40.5-41°C); the electron density distribution obtained has been analysed using the QTAIM and the topological and energetic parameters of intra- and inter-molecular interactions have been determined. The features of experimental and *ab initio* results are in good agreement and will be discussed.

Keywords: experimental charge density, ab initio calculations, pharmaceuticals

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Ru₃(CO)₁₂. Why D_{3h}?

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 $M_3(CO)_{12}$ (M= Fe, Ru, Os) may, in theory, adopt D₃, D_{3h}, C_{2v} symmetries. Iron carbonyl has a C_{2v} structure, ruthenium and osmium carbonyls have D_{3h} structures and only their derivatives show the least hindered D₃ configuration. The D_{3h} structure of Ru₃(CO)₁₂ has been justified theoretically using steric or electronic parameters. Only a marginal attention was paid to the deformation of axial CO groups, and it has been attributed to steric repulsion among oxygen atoms or to a more efficient orbital superposition between Ru atoms and C atoms. X-ray intensities of a Ru₃(CO)₁₂ crystal have been collected at low temperature and to them a multipole analysis has been applied; the electron density distribution obtained has been studied with the QTAIM and the topological and energetic parameters of intra- and inter-molecular interactions have been determined. Significant C_{ax}···C_{ax} interactions have been detected. The unexpected features of experimental electron density maps will be discussed.

Keywords: experimental charge density, ab initio calculations, ruthenium cluster

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On the (Non)-Planarity of 1,2,4,5-TetramethoxybenzeneChristopheM.L.VandeVelde,EviBultinck,FrankBlockhuys,UniversityofAntwerp,Belgium.E-mail:

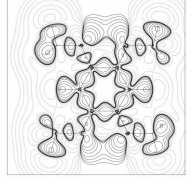
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The conformation of methoxy groups on phenyl rings is a longstanding problem that has been tackled in the past with a variety of structure determination techniques and at different levels of quantum chemical calculations.

1,2,4,5-tetramethoxybenzene has two sets of *ortho*-dimethoxy moieties and displays a nearly planar structure in the crystal. We obtained a high quality data set at 150 K which was suitable for

multipole refinement. The charge density maps and electronic properties in the bond critical points are compared with values from high-level quantum chemical calculations. The latter reveal a large number of energy minima on the potential energy surface.

The reason for the planarity of the molecule is not evident from the experimental structure, but the calculated bond orders



in the molecule indicate that the stabilizing factor off-setting the repulsion between the free electron pairs of oxygen is the participation of the latter in the π -electron system of the ring.

Keywords: ab initio calculations, charge density, conformational studies

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On the Conditions Leading to the Gaussian Distribution of the Magnetic Moments in a Spin-glass State

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The unperturbed ideal ferromagnetic system is described by a Hamiltonian of the Heisenberg type with the coupling constants J_{ii} (*i* and *i* numerate the magnetic ions). The presence of dopants (or defects) affects the values of J_{ii} for the magnetic ions surrounded by these dopants (or defects) in the random way. Thus the magnetic properties of such a system will depend on: the number of magnetic ions surrounding a j-th magnetic ion and interacting with it with the coupling constants J_{ij} , the number of dopants (or defects) surrounding a j-th ion, the coupling constants between i-th and j-th ions in the presence of dopants (or defects), and the random probability distributions of appearance of dopants (or defects) between i-th and jth ions. These distributions correspond - after introducing so-called "global" magnetic coupling constant (which turns out to be also random) - to the conditions of the central theorem of the theory of probability (the Lyapunov theorem). Thus the distributions of magnetic moments in such systems are Gaussian. Therefore one can use the Anderson-Edwards model of the spin-glass state in order to describe the system. Several examples of the spin-glass state are described in this approach.

Keywords: spin glass, probability, central theorem of the theory of probability

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Spin Density and Ordered Orbital of YTiO₃ Observed by X-ray Magnetic Diffraction

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CHARGE, SPIN AND MOMENTUM DENSITY

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A technique of X-ray magnetic diffraction (XMD) of ferromagnets was applied to an orbital ordering compound of YTiO₃. XMD, that is nonresonant X-ray magnetic Bragg scattring, is a unique experimental technique which enables one to measure spin- and orbital-magnetic form factor separately. The aim of this study is to observe directly the ordered orbital (wavefunction) in real space through the spin magnetic form factor measured by the XMD.

In $YTiO_3$ which is one of perovskite oxides, a Ti atom exists as a Ti^{3^+} ion, and has one 3d electron in t_{2g} state which shows orbital ordering. Below 30K this compound is ferromagnetic, and spin ordering occurs together with orbital ordering.

The XMD measurement was performed at KEK-PF-BL3C3, and the spin-magnetic form factor was measured for more than 22 reciprocal lattice points. By the Fourier transformation of it the spin density distribution was obtained in real space. As each Ti atom has one 3d electron with spin aligned in one direction, spin density distribution corresponds to 3d electron distribution which is correlated with 3d electron wavefunction. The experimental spin density was compared with calculated electron density base on t_{2g} wavefunction. MEM analysis of spin density is planned and the result will be shown. **Keywords: magnetic X-ray scattering, spin density, titanium oxide compounds**

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Charge Density Study of Cu₃(en)₂(CN)₄·H₂O

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In our recent work we have studied distribution of the electron density in the model compounds for blue proteins. Two model compounds [1] representing the reduced and the oxidized form of the protein contains copper in state +1 and +2. The mixed-valence complex $Cu_3(en)_2(CN)_4$, H_2O [2] contains copper atoms in both oxidation states, and so it should be possible to study electronic structure and particularly copper in both oxidation states not biased by different systematic errors. This provide us interesting comparison with our previous results.

Data were collected at at the beamline F1 at HASYLAB/DESY in Hamburg (T = 100 K, λ = 0.5604 Å). For corrections, integration and data reduction programs SAPRO, SAINT, and SORTAV were used. Multipole refinement was performed with XD software package.

[1] Flanagan S., Dong J., Haller K., Wang S., Scheidt W.R., Scott R.A., Webb T.R., Stanburry D.M., Wilson L.J., *J. Am. Chem. Soc.*, 1997, **119**, 8857. [2] Williams R. J., Larson A. C., Cromer D. T., *Acta Cryst.*, 1972, **B28**, 858. Keywords: charge density, copper(I), copper(II)