and so on. For example, in the case of polyoxymethylene, synchrotron X-ray diffraction gave us the observed reflections more than 700, from which the details of the chain conformation and chain packing mode were derived accurately. The 2-dimensional neutron diffraction data collected for both of H- and D-polyoxymethylene samples allowed us to extract the H and D atomic positions exactly as shown in Figure. The similar results were obtained also for the other polymers listed above.

Keywords: deuterated polymers, neutron diffraction data, hydrogen atoms

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Invarions for the DNA

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The application of the invariom database [1] of pseudoatom scattering factors that are transferable from one molecule to another (invarions) in the refinement of single crystal X-ray diffraction data has so far been limited to amino acids / oligopeptide molecules and to a number of organic compounds. Nucleotide bases and the macromolecules they are part of form another class of compounds that are ideally suited for a database approach. An advantage of a scattering-factor database built by purely theoretical methodology is that the database can be extended to any chemical environment with ease. However, scattering-factor (invariom) assignment based on experimentally determined geometries is sometimes not unambiguous. This problem does occur more often for nucleotide bases, where the distinction of bond orders from the bond distance is more difficult. We present a new algorithm that circumvents these difficulties and have extended the invariom database also to nucleotide bases. Charge-density quality single-crystal X-ray diffraction data on thymidine [2], adenosine and the watsontwick base pair 1-methyl-adenine-1-methyl-thymine provide real-life examples to validate these database entries. To highlight their applicability to larger systems, initial refinement results of X-ray data of a DNA-porphyrin complex [3] are also presented. Diffraction data to 0.86 Å resolution were retrieved from the Protein Data Bank (1EM0).


Keywords: charge density studies, nucleic acid complexes, databases

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Charge density and X-ray absorption spectroscopy of Mo-W complexes

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The heterobimetallic phosphido-bridged complexes, CpW(CO)3(μ-PPh2)Mo(CO)3 (1) with W-Mo distance 3.1723(4)Å and CpW(CO)3(μ-PPh3)Mo(CO)3 (2) with W-Mo distance 4.510(4)Å have been reported with special chemical reaction properties because of the interaction between W and Mo: complex 2 can be converted into 1 after irradiation with UV or heating at reflux temperature. In order to correlate how the chemical bond of Mo-W affect the chemical reaction in complex 1 and 2, the accurate single crystal diffraction data of both complexes at 100 K are collected and the experimental electron density in terms of multipole model are derived to investigate the electron density distribution and chemical bonds. All chemical bonds will be characterized based on atoms in molecule theory, and classified by the location of the bond critical point (BCP) and its associated topological properties. The bonding characters will be complemented with x-ray absorption spectroscopy of Mo K-edge. All experimental observations will be compared with the density functional theory calculation. A reasonable explanation will be given in relation with the different chemical reaction properties between 1 and 2.

Keywords: charge density, X-ray absorption spectroscopy, density functional theory

P14.03.01

Generalized library of experimental multipolar atoms

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A library of average multipole populations describing the electron density of common chemical groups is being built. The library values are obtained from several small peptide or organic compound crystal structures refined against available ultra-high-resolution X-ray diffraction data [1]. The atom types are defined on the basis of their chemical environment and local symmetry. New local coordinate axes systems have been defined in special cases. The introduction of optimal constrains and restraints allows for reduction of the number of multipolar parameters. As a consequence more meaningful results and stable refinements are gained. We will present the latest advancements of the crystallographic software MoPro suite [2] for the estimation of protein-ligand electrostatic interaction energy. Aldose reductase (hAR) is an enzyme involved in diabetes complications and the inhibition of the protein is a therapeutic way to treat them. The electrostatics interactions of the Fidarestat inhibitor with the enzyme active site have been characterized using the library electron density modelling. This information is useful to understand the affinity and specificity of Fidarestat with hAR compared to other inhibitor.
inhibitors.


Keywords: databases, charge density, electrostatic calculations

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**Topological properties of hydrogen bonds: Charge density studies by the maximum entropy method**

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The Maximum Entropy Method (MEM) is applied to low-temperature single crystal x-ray diffraction data of amino acids and peptides. The resulting electron density maps are analyzed according to Bader’s Atoms in Molecules theory (Bader, 1994) towards the determination of the electron density and its Laplacian at bond critical points (BCPs). Energy densities at BCPs are calculated according to Abramov (1997), employing the local virial theorem. Geometrical, topological and energetic properties of 52 hydrogen bonds are studied. The electron densities, their principal curvatures and the energy densities at BCPs are found to depend systematically on the distance between hydrogen and acceptor atom. These relations appear different from the corresponding relations as described by Espinosa et al. (1999) for multiple densities. MEM electron densities at the BCPs of hydrogen bonds are higher than the multiple densities. The difference between MEM and multiple densities is more pronounced for stronger hydrogen bonds. The contribution of the prior density (overlapping spherical atomic electron densities) to the MEM density is studied. The prior density seems to contribute a large part to topological and energetic properties at BCPs of hydrogen bonds. However, the properties at BCPs of covalent bonds of MEM densities differ significantly from the prior. Thus, for research into chemical bonds an experimental electron density should not be substituted by a promolecule.

References


Keywords: charge density, maximum-entropy method, peptides

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**Avoiding multiple diffraction for accurate charge density measurement using synchrotron radiation**

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Although contamination by multiple diffraction (MD) to the diffracted intensity is neglected in many cases of structure determination, its influence is not negligible for accurate measurement of charge density in crystal, especially with heavy elements, and must be avoided. Since in the researches with Synchrotron Radiation (SR) incident beams are highly collimated, it is generally believed that the MD has little effect on structure factor measurements. The aim of the present study is to see whether MD can be neglected or not. The authors measured integrated intensity from Yttrium Iron Garnet (YIG) crystal by a four-circle diffractometer with avalanche photo diode detector at Photon Factory BL14A. Two types of data collection were carried out, one at ordinary bisecting position and the other at the position avoiding MD by psi-rotation of the crystal (MD-free). In MD-free measurement, the perturbation due to MD in each structure factor was simulated from the orientation program controlling the diffractometer. 9910 reflections up to sin(theta)/lambda < 1.24 (wave length = 0.75Å, 2theta < 137 degrees) were measured in both cases. In the MD-free measurement only 91 reflections were measured at the bisecting position. The XAO analysis of the two data sets exhibited a significant improvement of the MD-free experiment compared to the intensity data measured at the ordinary bisecting setting.

Keywords: charge density, multiple diffraction, synchrotron radiation

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**Topology of the electrostatic potential in the analysis of molecular reactivities and hydrogen bonds**

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The topology of the electrostatic potential (EP) has been studied for single molecules using geometries and electron distributions determined from high-resolution single-crystal X-ray diffraction experiments. Local maxima, corresponding to the nuclei, and local minima are associated to electrophilic and nucleophilic sites, respectively. Space can be partitioned in primary bundles, which are volumes formed by field lines with the same start and end. Influence zones of electrophilic and nucleophilic sites can be built by joining primary bundles with a common start (nucleophilic) or end (electrophilic). Zero flux surfaces and saddle points can be used for delimiting influence zones, as they appear on the surface of the primary bundles. In the bonding regions, the topologies of EP and the electron density (ED) are similar, appearing a saddle point of EP analogous to the bond critical point of ED. As seen from a theoretical study on hydrogen bonded complexes, the region between the saddle points of EP and ED is populated by electrons of the acceptor atom which are electrostatically attracted toward the hydrogen atom. As the topological properties at both saddle points show a similar behavior, the analysis of EP is useful for the characterization of hydrogen bonds. In another theoretical study, the effect of the chemical and crystallographic environment on the hydrogen bond has been simulated by external electric fields. The perturbations induced on both the topologies of ED and the interaction energy are related.

Keywords: electrostatic potential, topology, hydrogen bonds