Ultrafast X-ray Absorption Studies of the Structural Dynamics of Molecular and Biological Systems in Solution

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Abstract: We review our recent studies of excited state structures and dynamics of chemical and biological systems with pico- and femtosecond X-ray absorption spectroscopy in the liquid phase.

Keywords: X-ray absorption spectroscopy

1. Introduction

Chemical reactivity and biological function are intimately linked to structure and structural dynamics on the molecular level. The advent of ultrafast spectroscopy in the mid-1980s has allowed the probing of chemical reactions in real-time. However, these studies are carried out using probe pulses in the optical domain, which only delivers information on the electronic structure. Therefore, over the past 10–15 years a huge effort was made aimed at using electron or X-ray probe pulses that can deliver geometric structural information such as bond lengths and bond angles, via scattering, diffraction, or X-ray spectroscopy which additionally delivers information about the electronic structure. Our primary efforts have mostly been focussed on developing and using time-domain X-ray absorption spectroscopy (XAS). X-ray absorption spectra are characterized by absorption edges, which are due to excitation of the core electrons of a given atom to the ionization threshold. An absorption edge consists of two regions: the X-ray absorption near edge structure (XANES) at energy just below and just above (typically <50 eV) the edge and the extended X-ray absorption fine structure (EXAFS), which lies at higher energies (typically >50 eV) above the edge. For an atom embedded in a molecule, crystal or even a liquid, both XANES and EXAFS exhibit modulations, which are due to the multiple and single scattering of the X-ray excited photoelectron onto the nearest neighbours.

Our choice of XAS is motivated by the fact that: a) it is element specific; b) it delivers information about the local structure (coordination numbers and bond distances) around the atom of interest, c) just below the edge bound-bound transitions occur, which contain information about the electronic structure (such as oxidation state, occupancy of valence orbitals, charge transfer, etc.) of the system; d) last but not least, it can be implemented in any phase of matter, and in particular in liquids, which are the natural environment of biology and of most of chemistry.

Time-resolved XAS requires tunable X-radiation with sufficiently short pulses, which can be synchronized to an external light source (e.g. a pump laser). Storage rings, such as Swiss Light Source (SLS) at the Paul-Scherrer-Institut in Villigen, offer the required specificities in terms of flux, energy range and stability, but the synchrotron X-ray pulses have widths of 50–100 ps. Therefore, in order to push into the femtosecond time domain, which is the fundamental time scale in chemistry, materials science and biology, the slicing scheme was implemented in the hard X-ray range at the MicroXas beamline of the SLS. It provides 100 fs hard X-ray pulses albeit at the cost of a dramatic reduction in flux. In all cases (ps or fs) the methodology is based on recording the transient (difference spectra of the excited minus the unexcited sample) absorption on a pulse-to-pulse basis. In the following sections, we illustrate the methodology on a few examples and how it helped answer specific scientific questions.

2. Intramolecular Charge Transfer in Metal Complexes

Transition metal complexes play an important role in chemistry and in many cases in biology. They are also of importance for solar energy research and for the development of molecular devices. Their visible absorption spectrum is dominated by the metal-to-ligand-charge-transfer (MLCT) bands. For the MLCT states, there is an issue of structural changes as a result of charge transfer. We addressed this question in a picosecond XAS study of aqueous ruthenium(ii)-tris-2,2'-bipyridine, ([Ru(bpy)3]2+). The system was excited in the 'MLCT band, which leads to population of the 'MLCT state on ultrafast time scales, (L1 (2p_{1/2}-4d) and L2 (2p_{3/2}-4d) edges of ruthenium were probed, delivering information on the electronic structure, i.e. the occupancy of the d(t2g) and d(e_g) orbitals in the excited triplet state. Information about the excited state structure was obtained via the electronic structure (based on the relationship between ligand field splitting and coordinating structure) and via the EXAFS at the L1 edge. Both showed that the structure change is a small (~0.037 ± 0.0135 Å) contraction of the Ru–N bonds. This relatively weak bond contraction is due to a balance between strong attractive forces between metal and ligand after charge transfer, and steric effects resulting from the three bipy ligands being in a constrained geometry, already in the ground state. Our results were later confirmed by quantum chemical calculations.
Similar studies have also been carried out on halogenated rhenium-carbonyl polypyridine complexes by probing the electronic and geometric structure at both the L-edges of rhenium and the K-edge of the halogen ligand. These will be discussed in a forthcoming publication.

3. Bond Formation in Bimetallic Complexes

The triplet excited states of binuclear d^8-d^8 platinum, rhodium, and iridium complexes (bridged by various ligands) have attracted much attention owing to their remarkable photophysical and photochemical properties, which are strongly determined by their structure. We investigated the tetrakis-μ-pyrophosphitodiplatinate(ii) [Pt4(P2O5H2)12]4− ion, which is the most studied binuclear metal complex due, among other reasons, to its photocatalytic activity. In the ground state the two metal atoms are not chemically bound, but excitation of the first singlet state results in a bond owing to the promotion of an electron from the antibonding d0* (d^4-derived) to the bonding p(σ-derived) orbitals, which leads to a contraction of the metal–metal bond. The system then undergoes intersystem crossing to the long lived (~1 µs) lowest triplet state.[10] Using time-resolved XAS, van der Veen et al. resolved its geometric[11] and electronic[12] structure via EXAFS and XANES, respectively. Fig. 1 shows the ground state Pt L_3-edge XAS spectrum (black trace) as well as the transient spectrum, integrated from 0 to 150 ns to improve the signal-to-noise ratio. The inset shows the XANES region for the ground-state complex and its transient spectrum, wherein dramatic changes appear. In particular, a new absorption shows up at 11.574 keV, just below the absorption edge, which is due to the creation of a hole in the 5dσ* orbital upon laser excitation that can then be accessed from the 2p_{3/2} core orbital (Ls, edge).[13] Clear changes are visible in the EXAFS region (Fig. 1b), reflecting structural modifications between the ground and excited triplet states.

From the transient EXAFS spectrum (Fig. 1b), the magnitude of the Pt–Pt bond contraction as well as, for the first time, the changes affecting the ligand bonds were extracted. In particular (see scheme in Fig. 1b), while the Pt–Pt bond contracts, the Pt–P bonds slightly elongate, in agreement with theoretical predictions.[13] This work underscores the ability to retrieve details of the excited state structure of a rather complex molecular system in solution. However, the X-ray determined structure deviates somewhat from that obtained from optical spectra (the system happens to be highly harmonic in both the ground and the excited state).[10] and we believe that this may stem from the neglect of the solvent contribution in modeling the EXAFS. These effects and eventually coordination of solvent molecules to the excited complex is an aspect which we are now investigating in detail (see below).

4. Spin Cross-over Complexes

One of the fascinating features of Fe(tii)-polypyridine complexes is their ability to undergo dramatic spin changes under temperature, pressure or irradiation. They have therefore been named spin crossover complexes (SCO complexes).[14] In the predominantly octahedral field of the ligands, all electrons are in the lower t2g sub-shell forming the low spin (LS) ground state, while transferring electrons to the e_g orbitals increases the spin state. Because the e_g orbitals derive from the d_{2g} and d_{3z} orbitals, they are antibonding in six-fold coordinated complexes, which leads to a striking metal–ligand bond elongation in the high spin (HS) state. A generic diagram of the potential energy curves of Fe(tii) SCO complexes is shown in the inset to Fig. 2, as a function of the Fe–N bond length. The ligand field states ^1T, ^3T and ^3E have their equilibrium distances elongated by ~0.1 Å, ~0.2 Å, and ~0.3 Å, respectively, relative to the ground state and MLCT bond distance. Light excitation into the 'MLCT state leads to population of the lowest quintet state ^5T, with unity quantum yield.[14] For the [Fe(tii)(bpy)]^3+ complex, which is the smallest of the family of Fe(tii)-complexes, the quintet state lifetime is the shortest, ~650 ps at room temperature. As a consequence, the quintet state structure could not be determined by quasi-static methods (X-ray diffraction or X-ray absorption spectroscopy). One of the questions that arises is whether the quintet lifetime is determined by the Fe–ligand bond distance, bearing in mind that nearly all SCO complexes exhibit the same bond elongation (typically ~0.2 Å) in the HS state.[17] We determined its HS state structure by recording the Fe K-edge XAS of the system with 70 ps X-ray pulses.[18] Fig. 3 shows the XANES of the molecule in the ground state, the transient (difference) spectrum at 50 ps time delay and the XAS spectrum of the quintet state, as retrieved from the ground state and the difference spectra and from the photolysis yield determined in laser-only experiments (and convoluted to match the much longer X-ray probe width). The structural analysis of the excited state delivered an Fe–N bond elongation of 0.203 ± 0.008
tion experiments confirmed this conclusion and identified the vibrational relaxation dynamics in the quintet state, thus providing a complete picture of the photocycle of aqueous [Fe(II)(bpy)_3]^2+, which is summarised in Fig. 2.

5. Hydrophobicity

XAS is an ideal tool for probing the solvent shell structure around atomic solutes. For halide ions in aqueous media, a dramatic change of solvent shell structure is expected in going from the halide to the halogen case, in what has been termed hydrophilic and hydrophobic solvation, respectively. For the case of aqueous iodide, the electron can be abstracted by UV excitation below 260 nm or by multi-photon excitation at 400 nm. The latter approach was used by us and the formation of the I_0 atom was probed via the L_1 (2s-5p) edge, which delivers electronic information, while the probing at the L_3 edge delivers structural information about the solvent shell via EXAFS.

We recently investigated the transition from hydrophilic to hydrophobic behaviour by femtosecond XAS and both classical and quantum mechanics/molecular mechanics (QMMM) molecular dynamics (MD) simulations. The general picture that emerges is that formation of a hydrophobic cavity takes typically 4 ps to be completed. This is captured by the QMMM calculations.
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The above ps results were all carried out on systems of fairly high concentrations (10’s of mM) using a pump laser at 1 kHz, while the synchrotron delivers X-ray pulses at 1.04 MHz. In order to exploit this higher repetition rate, and therefore increase the S/N ratio, we implemented a new laser set-up running at 520 kHz, and tested it on the aqueous [Fe(bpy)3]2+ sample. This set-up allows us to perform experiments at lower concentration samples, such as biological ones. As anticipated, an increase of the S/N by a factor of 20 was derived, which allows the study of biological samples. As a proof of principle, Fig. 6 shows the first ps pump-probe Fe K-edge transient of Myoglobin-CO (MbCO) in a physiological solution obtained at different time delays upon abstraction of the CO from the Fe atom. Because of the long lifetime of the system, these data were not collected on a pulse-to-pulse basis, but rather using the so-called asynchronous mode. Ps XAS transients have also been obtained on the MbNO system and will be discussed in a forthcoming publication.

7. Time-resolved Soft X-ray Studies of Liquid Systems

Extension of time-resolved XAS to the soft X-ray regime offers very exciting new advantages: a) most L-edge spectra of transition metals lie in the 200–2000 eV region; b) it allows the probing of light elements such as C, N, O, S, P, etc. which are very important in biology. The limitation of soft X-ray spectroscopies of high vapour pressure samples has been recently overcome by the use of high speed liquid jets in vacuum chambers or of cells equipped with soft X-ray transparent windows (silicon nitride windows). Recently, time-resolved soft X-ray absorption spectroscopy have entered the picosecond time domain by probing transient changes of liquid water heated by an ultrashort infrared laser pulse or the photoinduced changes in SCO complexes in solution in cells equipped with SiN windows. As a first step towards time-resolved studies of biological systems and in collaboration with the BESSY synchrotron in Berlin, we demonstrated the feasibility of soft X-ray absorption spectroscopy of biological systems in physiological solutions. Parallel to this, and for metallic and atomic and molecular ions in solution, we identified features in the soft X-ray excitation spectra of the total X-ray fluorescence that show up as dips, rather than bands. These dips reflect a strong interaction with the aqueous solvent and are due to the fact that the X-ray fluorescence is quenched by electron transfer from the d-orbitals of the metal to the oxygen of the water molecules. The time scale is estimated to be on the order of a fs based on the core-hole lifetime. These observations are promising to what we may call attosecond radiation chemistry, as they offer an exciting subject of investigation with high harmonic generation sources of attosecond pulses. The dips are also a powerful and simple diagnostics of the interactions of the solute with the water solvent molecules. When transposed into the time domain, which is one of our aims within MUST, these features will turn out to be extremely useful to probe solvent changes, along with the ultrafast XANES and EXAFS in the hard X-ray domain.

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