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Optimized finite difference method for the fullpotential XANES simulations: application to molecular adsorption geometries in MOFs and metal-ligand intersystem crossing transients

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ABSTRACT. Accurate modeling of the X-ray absorption near-edge spectra (XANES) is required to unravel the local structure of metal sites in complex systems and their structural changes upon chemical or light stimuli. Two relevant examples are reported here concerning: (i) the effect of molecular adsorption on 3d metals hosted inside metal-organic frameworks and (ii) light induced dynamics of spin cross-over in metal-organic complexes. In both cases, the amount of structural models for simulation can reach a hundred, depending on the number of structural parameters. Thus, the choice of accurate but computationally demanding finite difference method for the ab initio X-ray absorption simulations severely restricts the range of molecular systems that can be analyzed by personal computers. Employing FDMNES code [Phys. Rev. B, 63, 125120 (2001)] we show that this problem can be handled if a proper diagonalization scheme is applied. Due to the use of dedicated solvers for sparse matrices, the calculation time was reduced by more than one order of magnitude compared to the standard Gaussian method while the amount of required RAM was halved. Ni K-edge XANES simulations performed by the accelerated version of the code allowed analyzing the coordination geometry of CO and NO on the Ni active sites inCPO-27-Ni MOF. The Ni-CO configuration was found to be linear while Ni-NO was bent by almost 90°. Modeling of the Fe K-edge XANES of photoexcited aqueous [Fe(bpy)₃]²⁺with a 100 ps delay we identified the Fe-N distance elongation and bipyridine rotation upon transition from the initial low-spin to the final high-spin state. Subsequently, the X-ray absorption spectrum for the intermediate triplet state with expected 100 fs lifetime was theoretically predicted.

1. Introduction

The knowledge about local atomic and electronic structure of the active center is required for understanding the elementary steps of chemical reaction. Experimental tools such as X-ray absorption near-edge spectroscopy (XANES) or resonant X-ray diffraction (RXD) can extract information about the electronic structure of the absorbing atom of interest, but also about its local coordination environment. The interpretation of these experiments is often impossible without quantum *ab initio* simulations.

Much progress has been made over the past two decades in the understanding of the near-edge features of the X-ray absorption spectra (XANES region).¹ Earlier XANES spectra were used mainly in a qualitative way, considering the presence/absence of certain fingerprint features and comparing the data with the spectra of model compounds.² The appearance of progressively more sophisticated codes, together with the increased computational capabilities, has made XANES a spectroscopy able to quantitatively confirm, or discard a structural model for the environment of the X-ray absorbing atom,^{1i, 3} forming thus a new fundamental diagnostic tool in condensed matter physics and chemistry.⁴ Among such codes multiple scattering *FEFF-9,⁵ XKDQ,⁶ CONTINUUM,⁷ SPR-KKR⁸* and periodic basis set *Wien2k,⁹* and *PARATEC*¹⁰ should be mentioned. Several density functional theory (DFT)-based codes with localized orbitals in a basis set, *ADF*¹¹ and *ORCA*,¹² are able to directly compute the pre-edge region within the time-dependent DFT approximation. TDDFT can provide reasonable values for XANES cross-section up to 40eV above absorption edge¹³ and other techniques are being developed to access higher-lying electronic core-excited states, e.g. algebraic-diagrammatic construction scheme.¹⁴

The Finite Difference Method (FDM) is attractive for calculations of the photoelectron wavefunction up to 100-200 eV above the absorption edge. It avoids in a simple way the muffin tin approximation most often used, for example, in the multiple scattering theory (MST). In the latter approximation, the potential is assumed to be spherical inside the touching (or overlapping) atomic spheres and constant between them.^{1c} However, in some cases the MST scheme might be not fully applicable, especially when the studied compound is sparse or with strong covalent bonds. This is the case for some molecules during photoexcitation¹⁵, or the analysis of defects in semiconductors^{3c}. The FDMNES project¹⁶ aims to provide a user-friendly tool for the accurate XANES and RXD simulations.

In this work we introduce a new version of the FDMNES code where the computational cost of FDM scheme is reduced by more than one order of magnitude and required RAM is decreased by a factor of 2. This implies that the FDM becomes comparable to the MST technique^{4a, 4c, 17} in terms of CPU demand, with the evident advantage of a superior accuracy. This major improvement is due to the use of sparse matrix solvers, which are applied for the first time to XANES calculations, and a new scheme for parallelization. The original code was based on Gaussian method to solve the very large set of linear equations coming with the FDM. More than 95% of elements in the corresponding system matrix are zeros, while the non-zero ones concentrate predominantly around the leading diagonal. To deal with this configuration of the matrix in the most efficient way we have introduced the sparse storage form bundled with the correspondent solvers, which are much better suited for the problem. Diagonalization with Unsymmetric Multi Frontal method, UMFPACK,¹⁷ or with MUltifrontal Massively Parallel sparse direct Solver, MUMPS,¹⁸ speeds up the calculations by a factor of 5 to 40 on a single core, depending on the input and hardware. The latter method is preferable for parallelization on multicore processors, because it requires less RAM. However, the scaling of the method strongly depends on the matrix used for diagonalization.

Two widely discussed structures were chosen to test the accelerated version of the code. The first object is a porous metal-organic framework (MOF) CPO-27-Ni. MOFs are three-dimensional crystalline porous coordination polymers of alternating metal clusters (or single metal atoms) and organic units. They can be used for gas storage,¹⁹ gas separation and purification,^{3b, 20} liquid phase separation,²¹ drug delivery,²² catalysis,²³ molecular sensors,²⁴ and solid state ion conduction²⁵. Moreover, MOFs can represent a new class of optical,^{4b, 26} magnetic^{19h, 27} and semi-conducting²⁸ materials. Although XANES spectroscopy was successfully used in several CPO-27-Ni studies,^{20,} ²⁹ the data were analyzed only qualitatively. Several problems, which are typical for XANES simulations of MOFs in particular and porous materials in general, occur at this point. Very pronounced asymmetry of the environment of the absorbing atom makes the relatively fast muffintin XANES codes less competitive, since the real potential is different from the one used in muffintin approximation. Moreover, due to the high porosity of the material one has to choose a rather large radius for the sphere in which the calculation is performed to include a sufficient number of atoms to reproduce the experimental spectrum. All these factors increase the computational time and the amount of memory required, thus shifting the full-potential XANES calculations for many porous materials way out of reach of personal computers into the dominion of the large-scale computational clusters.

The second example concerns a single metal-organic molecule that undergoes structural changes upon laser excitation. Tris(bipyridine)iron(II) $[Fe(bpy)_3]^{2+}$ is a $3d^6$ spin-crossover complex that undergoes a spin transition from a singlet low spin (LS) ground state to a short-lived high spin (HS) quintet state by visible light irradiation³⁰. Upon 400 nm laser pulse excitation, an electron is excited to a manifold of Metal-to-Ligand ChargeTransfer (MLCT) states of mixed spin multiplicity (mainly singlet and triplet character) and decays on ultrafast timescale to the lowest energy metal-centered HS quintet state.³¹ The relaxation pathway from MLCT state(s) towards HS state involves another extremely short-lived intermediate state of triplet spin, which has been only very recently observed using femtosecond-resolved X-ray emission spectroscopy,³² in agreement with earlier theoretical prediction.³¹ Geometry structure of the HS quintet state was previously analyzed from EXAFS,³³ and molecular dynamics DFT simulations.³⁴ Compared to the ground state the Fe-N distance changes by 0.2Å and bipyridine ligands undergo rotation around twist axis by approximately 7 degrees as shown by Lawson Dakuet al.³⁵ However, there is still no available experimental XANES data for the intermediate triplet state. According to the theory,³¹ its structure, along the Fe-N bond distance reaction coordinate, should differ from the ground state by 0.1 Å, which is significantly shorter than the fully relaxed HS quintet state. Based on the time-resolved Fe K-edge XANES we perform the structural analysis for the high spin state of $[Fe(bpy)_3]^{2+}$ and introduce an option to use external spin-polarized electron density in the FDM procedure for metalcentered singlet, triplet and quintet spin states.

Section 2.1 reports the details of the changes in the code applied to speed up the finite difference procedure. In section 2.2 we present a quantitative comparison between the original and the accelerated version of the FDMNES code in terms of speed and memory demand. Section 2.3 describes a way to introduce a self-consistent electron density to the calculations. Readers that are interested in the application of the code in solid state physics and chemistry are invited to move directly to Section 3.1 where we show the ability of FDMNES to calculate rapidly and accurately the Ni K-edge XANES spectra of CPO-27-Ni MOF. Calculations were performed also with NO and CO molecules adsorbed on Ni sites. The preferred orientation of the gas molecules on the metal centers was determined exploiting the sensitivity of XANES to bond angles. Section 3.2 reports the FDM simulations of the Fe K-edge in the ground, triplet and quintet states of

 $[Fe(bpy)_3]^{2+}$ molecule based on self-consistent electron density obtained in separate DFT calculations.

2. Computational methods

X-ray absorption cross section obeys Fermi golden rule and dipole matrix element for the electron transition can be written as:

$$\sigma(E) \sim \left| \left\langle \psi_f \left| \mathbf{\epsilon} \cdot \mathbf{r} \right| \psi_{1s} \right\rangle \right|^2 \cdot \delta \left(E - E_f - E_{1s} \right)$$
(i)

where ψ_{Is} is a core-electron wave function, ψ_f is a wave function for unoccupied state, ε is a photon polarization. The simulation of the absorption cross section needs the calculation of the states probed by the photoelectron. It requires solving the Schrödinger equation, the relativistic Schrödinger equation or the Dirac equation depending on the nature of the absorber. This task is fulfilled in FDMNES by using either MST or FDM approach, which were mentioned above. FDM has been known for a long time;³⁶ it has been a key part of FDMNES from its beginning.³⁷ In this approach the whole space is divided in three regions: (i) an outer sphere surrounding the cluster of interest, (ii) an atomic region composed by very little spheres around the atomic cores (no more than 0.65 Å of radius, much smaller than the usual muffin-tin sphere) and (iii) the interatomic region where an FDM formulation of the Schrödinger equation (SE) is performed. In the outersphere and in the little atomic spheres classical expansion in spherical harmonics of the wave functions is performed. In the interatomic region, the Laplacian is discretized by approximating the wave function ψ around the point *i* by a polynomial of the 4th order. In this way the SE becomes:

$$\left(-\Delta + V_{Coul} + V_{XC}\right) \cdot \psi = E \cdot \psi \quad \rightarrow \left(-\sum_{j} l_{ij} + V_i^{Coul} + V_i^{XC}\right) \cdot \psi_i = E \cdot \psi_i$$
(ii)

where Coulomb and exchange-correlation potentials (V^{Coul} and V^{XC} respectively) are replaced by their values in a point *i*, and l_{ij} are the Laplacian operators making the connection between the point *i* and its neighbors *j*. For example in an orthogonal frame $l_{ii} = -15/2d^2$ and $l_{ij} = 4/3d^2$ and $-1/12d^2$ respectively for the first and second neighboring points, *d* being the inter-point distance. For a uniform mesh FDM is equivalent to the finite element method. More details and the way to maintain the continuity between the regions of calculation can be found in Ref.^{16a} The potential V_i on the grid of points can be calculated internally with or without a self-consistent procedure. It can also be imported from other code as demonstrated further on. Consequently, one has to deal with a large system of n linear equations, *n* being the number of points in the area of calculation. For any given energy *E* the system of linear equations is solved by matrix diagonalization and one gets the solution, that is the absorbing atom wave function amplitudes, everywhere in the region of calculation. Obtained values are finally injected in the cross-section equation to get the signal response. The smaller is *d*, the more precise will be the simulation, but at the same time the larger will be the system to solve. Regardless of the way to order the points in the system, (for example from the center toward the outer-sphere), one gets a sparse band-like matrix.

2.1 Implementation of the sparse solvers

In the FDMNES code the major part of CPU time is used to solve a large system of linear equations. Non-zero elements concentrate near the leading diagonal of the corresponding matrix. To save computing memory and time, after Gaussian elimination the authors of the original code store only the elements of the lower half band starting from the first non zero element in each matrix line.^{16a} In fact, even before Gaussian elimination this band also contains many zeros (Fig.

1 - yellow points). The diagonal neighborhood on the Fig. 1 seems to be full of non-zeros (blue dots), but quantitative analysis confirms that it consists of 96% of zeros. Therefore sparse storage form, which keeps only non-zero elements and their positions, is better suited for the problem.

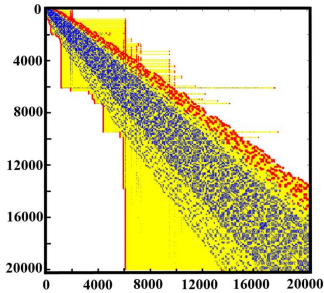


Figure 1.Visualization of the elements in the finite difference matrix. Blue color denotes non-zero elements, red – boundary elements, yellow – zeros inside the diagonal neighborhood.

In order to speedup the computation, the parallel mode was introduced to FDMNES by Hatada, Hayakawa, Wilcke and Joly which divides calculations by energy points, so that each node solves linear system corresponding to its own energy value. The calculations are independent and the decrease of computational time equals the number of used hosts. However this method does not exploit the sparseness of the matrix and therefore for large linear systems the required RAM volume becomes huge and does not allow running two or more process on one multicore host.

Naturally, sparse matrices require the correspondent specialized solvers. A widely used one, UMFPACK, is an Unsymmetric Multi Frontal direct solver for sparse matrices developed by Davis et.al.^{17, 38} It has a sequential algorithm which accelerates the solving of a linear system but does not fully utilize multicore computational resources. Thus the problem with running two or more processes for the solution of linear system on one multicore host remains. Therefore another solver with low memory requirements, which utilizes fully multicore processor is needed. Such are the features of MUMPS. MUltifrontal Massively Parallel Solver is a package for solving systems of linear equations by direct method based on a multifrontal approach.¹⁸

In original FDMNES code single MPI process solves a linear system for a given energy value in Schrödinger equation. We change FDMNES parallel structure so, that several MPI-processes can solve one linear system (see Fig.2). All MPI-processes are now divided into groups of size *p* each. A user passes the size of group to the program by means of an environmental variable. Processes of one group perform parallel calculation for common energy value on common system of linear equations. When the atomic cluster radius is high, the correspondent linear system fills the memory of a personal computer and in case of original FDMNES a user has to run only one MPI-process per computer, which uses only one core and the whole memory. The new FDMNES with MUMPS solver can calculate the linear system in parallel by the processes of one group on the same machine and utilize fully multicore processor without increase of occupied memory. For huge problems, which one computer memory can't hold, we can make the solution possible by running it in parallel on enough number p of cluster nodes.

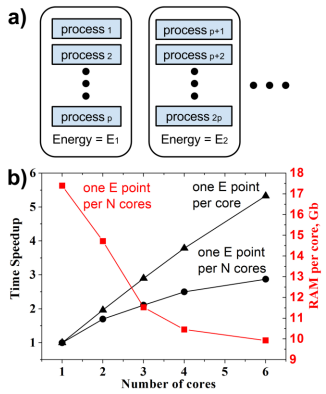


Figure 2.(a) New FDMNES parallel structure. The groups of p processes each solve the linear system for the common energy value. (b) Scaling tests performed for a 6-core processor for two options: when equation is solved for several number of energies, each per one core, or when single energy point is distributed over given number of cores. In the last case the memory required per each core is also reduced.

When Gaussian solver is used in FDMES it occupies up to 98.5% of working time. Sparse solvers changed the time distribution picture (see figure S2 in Supplementary materials) and matrix diagonalization occupies only 22% of working time while the rest is required for the construction of the system of equations. MUMPS scaling depends greatly on the matrix to solve. The solution of finite element Navier-Stokes codes by MUMPS does not scale much beyond 6 processors. MUMPS maximum speedup in the problem of acoustic propagation and radiation of turbo machinery noise³⁹ is about 5 on different multicore architectures with shared memory and stops increasing much after 8 MPI processes. One of the main bottleneck is the possibility for the matrix to be divided into sub matrices with equal solution times. The reached speedup for FDMNES with MUMPS solved is 3 when using 6 cores on one node as shown in figure 2b. In this case the RAM used by each core is reduced almost by a factor of 2. When several nodes are available the best strategy is to calculate different energy points of XANES spectrum on distinct nodes, while within one node to use up to 6 cores for matrix diagonalization.

2.2. Acceleration tests

The main idea of developing the new version of FDMNES was to make XANES calculations beyond the muffin-tin approximation feasible on a standard modern PC, thus decreasing the need in large supercomputer centers for this task. For performance tests we chose a PC with Intel Core i7-3820 processor and 32 GB of DDR3 memory. The program and the libraries were built with the Intel compiler. The structure used for calculation was a low-symmetry cluster of CPO-27-Ni MOF discussed further in Section IV.

Fig.3 shows how the calculated spectrum converges with the increase of cluster radius. In the panel a) the normalized Ni K-edge spectra of CPO-27-NiMOF are presented, while panel b) exhibits the difference spectra with respect to the one calculated with the largest radius R=7 Å. For the cluster size R=4Å the difference amplitude is about 10% from the XAS signal, for R=5 Å and R=6 Å it is smaller than 3%. The data reported in Fig. 3 imply that large clusters are required in order to reach convergence for this class of materials and 3% accuracy can be achieved if we use sphere with radius larger than 5Å for simulations.

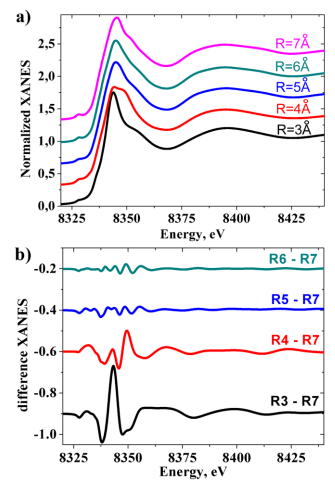


Figure 3. (a) Normalized theoretical Ni K-edge XANES spectra as a function of cluster size used in calculations for the CPO-27-Ni structure. (b) Differences calculated for the spectra from panel (a) with respect to R=7Å. In both panels the spectra are shifted vertically for the sake of clarity.

The time required to calculate XANES spectrum with one core was measured for the clusters of different size. The speedup value was obtained with respect to a time required for the same task by a standard FDMNES code with Gaussian solver. The Fig. 4 shows that FDMNES with MUMPS solver achieves the best speedup, at the same time decreasing the memory requirements. UMFPACK solver is twice as slow, despite the use of the same Gauss transformation ordering method⁴⁰.

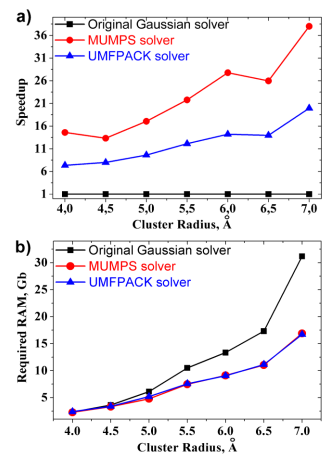


Figure 4.FDMNES speedup (a) and RAM consumption (b) for different solvers as a function of cluster radius.

After implementation of the new sparse solvers we have tested all model examples provided with the FDMNES code, which cover many branches of the X-ray spectroscopy. The output results were compared with those obtained with the standard Gaussian diagonalization scheme and no deviations within the computational accuracy were found (see Supplementary materials). Subsequently in Section 3.1 we have investigated a more sophisticated case, to demonstrate that the accelerated version allows performing simulations of complex systems on personal computers. For this study we also chose CPO-27-Ni MOF, as a prominent example of modern crystalline microporous materials.

2.3 FDM coupled with molecular orbital DFT

Implementation of the finite difference method for XANES relies on the *a priori* knowledge of the exchange correlation V^{XC} and Coulomb V^{Coul} potentials in equation (ii) which, in turn, requires the knowledge of the electron density. In first approximation a total density could be obtained as a superposition of atomic densities. It works well in many cases, *e.g.* those shown in Section 3.1. However, XANES calculations of different spin states of 3d metal complexes require a self-consistent way to obtain V^{XC} and V^{Coul} . The self-consistent procedure can be realized for a MST approach but within muffin-tin approximation for the shape of the potential. Alternatively Schrödinger equation can be solved self-consistently within the finite difference method.⁴¹ The latter approach is however highly inefficient because too many trial energy points should be used

before the energy eigenvalue is found. Thus an optimal option to get a self-consistent density for FDM is to import it on a 3D grid of points from a separate quantum chemistry simulations. For periodic structures, FMDNES offers a possibility to import electron density from the full-potential Wien2k calculations which utilizes augmented plane waves as a basis set. For molecules, we implement a possibility to use electron density obtained in DFT simulation based on molecular orbitals (MO-DFT) in ADF program package.^{11a, 42} In such a way, we exploit all the advantages of the MO-DFT method to obtain a self-consistent electron density and finite difference method for XANES calculations. Section 3.2 demonstrates the procedure applied for tris(bipyridine)iron(II) $[Fe(bpy)_3]^{2+}$ complex in different spin states.

3. Case studies

3.1 Interaction of porous CPO-27-Ni MOF with CO and NO gases

This microporous compound with a formula Ni₂(dhtp)(H₂O)₂·8H₂O has a three-dimensional honeycomb-like structure that is formed by nickel-oxygen chains connected by the 2,5-dihydroxyterephthalic acid linkers.⁴³ It crystallizes in the trigonal $R\bar{3}$ space group, with a cell volume 3900 Å³. As-synthesized material contains a lot of water: one molecule is coordinated to each Ni site and the rest is distributed in the channels. Upon the activation in vacuum at 120°C for one hour the water is removed from the material, leaving Ni ions with five oxygens in the first coordination shell at slightly varying Ni-O distances (1.885, 1.945, 1.991, 2.074, and 2.120 Å,⁴³ see Fig.5a). Many potential applications of MOFs involve interaction with gases, so gas adsorption studies constitute an important part of their characterization. CPO-27-Ni was tested in H₂, CO, NO N₂,C₂H₄, H₂S environments and changes of the crystalline structure and local environment of Ni were monitored by EXAFS, XRPD, UV-Vis, FT-IR and X-ray emission spectroscopy^{20, 29, 43-44} 20, 29, 43-44

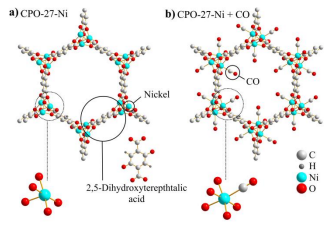


Figure 5. The structure of CPO-27-Ni MOF in dehydrated form (a) and upon activation with CO (b). Color code: Ni (cyan), C (light gray), O (red), H (dark, gray).

For XANES simulation we used a crystal structures of CPO-27-Ni solved by powder X-ray diffraction,⁴³ supported by EXAFS analysis, UV-Vis and IR data^{20, 29},^{20, 29} and optimized by means of density functional theory using B3LYP-D*/TZVP functional in the periodic approach of CRYSTAL-09 code⁴⁵ as described in the works of Valenzano et al.^{45c, 46}

Fig.6a shows the simulated Ni K-edge XANES of the dehydrated CPO-27-Ni compared to the experimental data. The agreement is very good since the calculation was performed within the sphere with the radius of 6 Å without imposing any symmetry restrictions. Slight mismatch in the peak positions may be explained by an overestimation of the bond distances in the optimized

theoretical structure used for calculation (around 0.03Å for Ni–O and 0.05 Å for Ni–Ni distances compared to EXAFS data^{45c}). Calculation time for 160 energy points was 2 hours at a PC with the 4-core Intel Core i7-3820 processor. Such task with the previous version of FDMNES would have required twice more memory and more than 2 days to be completed.

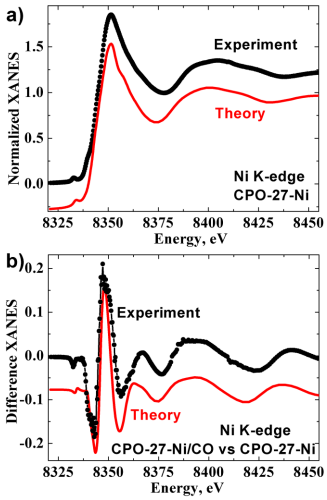


Figure 6. (a) Comparison of experimental and simulated Ni K-edge XANES data for the dehydrated CPO-27-Ni. (b) Difference spectrum (CPO-27-Ni–CPO-27-Ni/CO) highlighting the changes that occur in the Ni K-edge XANES spectrum of the dehydrated CPO-27-Ni upon the absorption of CO. The simulations were performed using the corresponding structures displayed in Figure 5 and a cluster radius of 6Å. In both panels the spectra are shifted vertically for the sake of clarity.

Even more impressive results can be obtained when calculating the difference spectra of the dehydrated CPO-27-Ni and the material after the adsorption of CO gas, at an equilibrium pressure resulting into a 1:1 Ni-CO adduct. Fig.6b shows the experimental and theoretical difference XANES obtained subtracting the spectrum of the dehydrated material from the one measured in presence of CO. All the changes, which are qualitatively described in the work of Chavan et al.^{29b}, are reproduced, thus confirming the geometry of the adsorption site.

EXAFS is a good technique in the characterization of the geometry adopted by diatomic molecules AB adsorbed on metal M sites exhibiting one (or more) coordination vacancies.⁴⁷ Indeed, the M-A distance is usually determined within 0.01-0.02 Å accuracy. If the M-A-B angle approaches 180°, the focusing effects of the central A atom enhances the multiple scattering

contributions. So, usually it is easy to discriminate with EXAFS between linear and bent absorption geometries. However, among bent geometries it is difficult to determine the M-A-B angle when it is in the 160° - 90° interval. In the particular case of CPO-27-Ni, the situation is even more complex, because the second shell single scattering M-B contribution falls in the same R-interval of the very strong, two-fold degenerated Ni-Ni contribution along the *c* direction^{20, 29, 20, 29} Moreover, the flexibility of the CPO-27-Ni framework, results in a stretching of the Ni–Ni distance from 2.89 Å to 2.97 (2.94) Å upon CO (NO) adsorption. This fact prevents to fix the framework contributions in the EXAFS fit of the material in presence of adsorbate, making the determination of the geometry of the CO (NO) molecule difficult, even with high quality data analyzable up to 18 Å⁻¹. In case of CO the value of Ni-C-O angle of 180° resulted in a reasonable EXAFS fit.^{29b} In the case of NO authors performed fits where the Ni-N-O angle was fixed to 180°, 160° and 130°.^{29a} In the three cases very similar fits were obtained (R-factor = 0.045, 0.043 and 0.052, respectively) and the authors stated the inability to determine with sufficient confidence the Ni–N–O angle.^{29a} Successively, periodic ab initio calculations optimized the Ni–C–O and Ni–N–O angles to 172° and 123.2°, respectively.^{45c, 46}

It is evident that an experimental determination of the Ni-C-O and Ni-N-O angles is needed and it can be provided by XANES spectroscopy comparing the experimental spectrum and the simulations performed at different angles. To demonstrate this, XANES has been calculated for both CO and NO molecules for Ni-C-O and Ni-N-O angles in the 70°-280° interval.^{45c, 46} The integral over absolute difference between theoretical and experimental difference spectra was used for quantitative comparison (Fig.7)

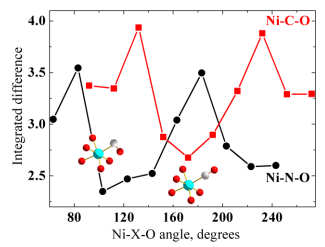


Figure 7. The discrepancy between experimental XANES data and theoretical models as a function of Ni-C-O and Ni-N-O angles. Red (black) curve corresponds to CPO-27-Ni with CO (NO) absorbed.

As expected, the minimum for CO is around 180°, in agreement with EXAFS and DFT results^{20, 46}.^{20, 46} For NO the XANES simulation rejects the linear geometry, which might look plausible from EXAFS. In agreement with theoretical findings^{45c} bent configuration with Ni-N-O angle around 100-120° is favored over the linear one.

3.2 Ground and excited states of the 3d metal complexes.

Iron 3*d* states in $[Fe(bpy)_3]^{2+}$ are split by nearly octahedral crystal field into e_g and t_{2g} manifolds. In a ground state bonding t_{2g} -like orbitals are fully occupied, while antibonding Fe 3*d* (e_g)-like orbitals are vacant. Upon laser excitation and subsequent series of electron transitions the molecule ends up in the HS state with populated $3d (e_g)$ manifold. The presence of electron density on the antibonding orbital initiates distortions of the Fe local atomic structure.DFT structural optimization with standard local exchange-correlation functionals can predict the geometry of the molecule in a ground- and high-spin states in good agreement with EXAFS and molecular dynamics.³⁵ We have used MO-DFT ADF program package, the TZ2P basis set and GGA-PBE exchange correlation potential to calculate the geometrical structure, electron density and corelevel energy for the singlet, triplet and quintet spin states of the[Fe(bpy)₃]²⁺. The core hole on Fe 1s level was introduced for XANES simulations.

Figure 8 shows the changes in the $[Fe(bpy)_3]^{2+}$ total potential $(V^{Coul} + V^{XC})$ after self-consistent procedure was applied to the initial superposition of atomic densities (the procedure used by default in the FDMNES code). The biggest differences are observed between carbon atoms and between Fe and N atoms indicating density redistribution and chemical bond formation (Fig 8, bottom).

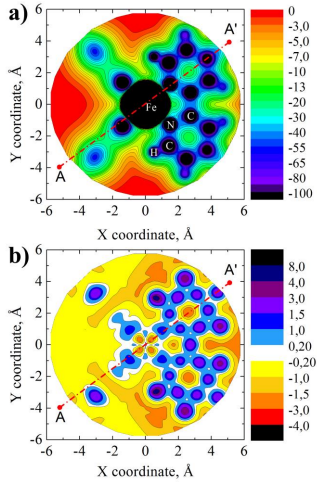


Figure 8. (a) The two dimensional potential cut for the $[Fe(bpy)_3]^{2+}$ molecule. (b): The difference between potentials obtained as superposition of atomic orbitals and after self-consistent MO-DFT calculation. A-A' is a guideline to highlight that largest differences are observed not in atomic spheres but between them in the area of covalent bonds. This line intersects iron, nitrogen and carbon atoms successively.

Three approaches to simulate Fe K-edge XANES for the ground state of the complex were compared (Fig. 9a). Finite difference method was used with two different options. In the first one, the total potential in equation (ii) was calculated on a basis of electron density from superposition

of atomic densities without self-consistency (FDM*non-SCF). In the second one, a self-consistent electron density was used from separate MO-DFT simulation (FDM*MO-DFT). The third approach was the Green-function-based multiple scattering method in muffin-tin approximation (GF-MTA). All spectra were calculated for the crystallographic structure⁴⁸ and equal broadening parameters were applied to compare with experimental data. As expected in the energy region above 80eV from absorption edge all methods are similar. However within first 50 eV FDM*non-SCF provides superior agreement with experiment than GF-MTA, while FDM based on selfconsistent MO-DFT electron density shows superior agreement with experiment than non selfconsistent FDM. We attribute this effect mainly to the local redistribution of the electron density around absorbing iron atom. To prove this we started from the FDM calculations based on non self-consistent superposition of atomic densities (blue curve FDM*non-SCF in Fig. 9a) and manually introduced changes in the electron configuration of Fe atom. Figure 9b shows how the Fe K-edge XANES intensities change along with number of Fe d-electrons. The d-shell occupation number was varied stepwise from 6 to 7. In order to conserve the charge of the molecule the electron from the d-orbital was transferred to a delocalized iron 4s orbital. The main changes are observed in the beginning of XANES spectrum: the intensity of peaks A and B decreases relative to the peak C, while higher-energy region of spectrum remains practically unchanged.

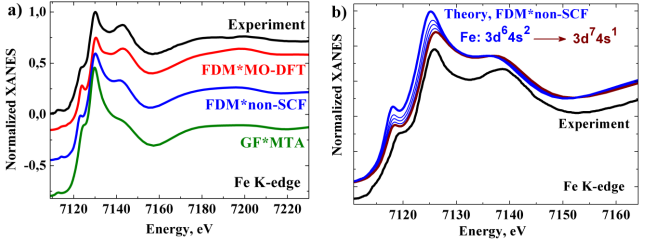


Figure 9. (a) Experimental Fe K-edge XANES (black) compared to the theoretical calculations within three different approaches: green curve is a Green-function method within muffin-tin approximation (GF*MTA). Blue curve is a non self-consistent finite difference method (FDM*not-SCF) with an electron density calculated as a superposition of atomic orbitals. Red curve is a finite difference method with an electron density imported from molecular orbital DFT calculation (FDM*MO-DFT). (b) FDM*not-SCF simulations for different number of electrons on a Fe d-level.

Based on FDM*MO-DFT we compare the calculated signal for quintet state with experimental one measured at a time delay $\Delta t = 100$ ps after irradiation with a 10 ps laser pulse at 532 nm. Fe-N distance in the relaxed HS structure is 2.18Å that is 0.21Å longer than in a singlet state. For a triplet state 2.07 Å distance was obtained that is 0.1 Å longer. Relative chemical shift of the Fe 1s level for HS state compared to the LS state was calculated as 0.5 eV, and 0.1 eV for a triplet state compared to the LS state. Theoretical transient spectrum for the high spin state reproduces all trends of the experimental data, such as intensities and energy positions of peaks. The blue curve of calculated transient for the triplet state can serve as a guideline for future pump-probe experiments with a better time resolution since the lifetime of the triplet state was predicted theoretically less than 50 fs.³¹

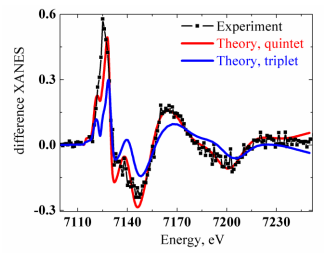


Figure 10. Experimental Fe K-edge transient signal for the $[Fe(bpy)_3]^{2+}$ and theoretical calculations for the high spin quintet state and intermediate triplet state with relaxed geometries.

The choice of exchange-correlation functional affects the electron energy level diagram,⁴⁹ especially in the vicinity of the HOMO-LUMO gap. This is particularly important for the unoccupied levels probed in the pre-edge region of XANES spectra. Currently FDMNES offers the choice between Hedin-Lundquist, Perdew and Wang and X-alpha options to calculate energy dependent exchange correlation functional from the electron density. We compare them in figure S4 of Supplementary materials for the Fe K-edge XANES of the high spin state of iron trisbipyridine. It is the aim of our future research to implement other possibilities, such as hybrid functionals with different exchange contributions.

5. Conclusions and Outlook

We have presented a computational scheme that makes accurate but time-consuming finite difference method available for XANES simulations on personal computers. For large non-symmetric systems the matrix diagonalization method adapted for the sparse matrices shows up to 40 times speedup in the calculation compared to the original Gaussian scheme. Thus we narrow the gap between the accurate but time-consuming finite difference method and the fast, but not always applicable multiple scattering approach, which utilizes muffin-tin approximation. The new improved method was tested on a metal-organic framework CPO-27-Ni in dehydrated state as well as after CO and NO absorption. XANES simulation allowed identifying the adsorption geometry of the Ni–CO and Ni–NO as linear and bent (with a Ni–N–O angle 100–120°) respectively. Using the self-consistent electron density from the MO-DFT simulations we were able to reproduce both the shape and chemical shift of the XANES spectrum for high spin structure of tris(bipyridine)iron(II) complex. Based on this we have calculated theoretical spectrum for a short-lived triplet state of $[Fe(bpy)_3]^{2+}$ (<100fs), which has only recently been indicated by theoretical simulations and x-ray emission experiments. This calculation can be used as a guideline for further pump-probe XANES experimental investigations on x-ray free electron lasers.

This accelerated version will be of great help in the fully relativistic simulations of the XANES spectra that are very time-consuming and require using of spinors and Dirac equation. The code will be interesting also for correct simulation of the XANES spectra of disordered systems such as ions in solution⁵⁰ and metallo-proteins.⁵¹ In this case a proper description of the thermal and structural fluctuations is mandatory to reproduce the XANES experimental signal. Indeed the

measured XANES spectrum can be obtained from the microscopic description of the system derived from molecular dynamics simulations allowing one to determine the different local configurations and their fractional population along a suitable time period. This approach needs to compute several different XANES spectra (one for each local configuration) and to average them with the corresponding probability weight. However even accelerated finite difference method would hardly compete with existing multiple scattering methods in the energy region higher that 100 eV above the absorption edge where both methods provide similar accuracy. We believe that future ideal method for theoretical analysis of XAS would combine FDM for the first 100 eV above absorption edge and MST for the higher energy region.

ASSOCIATED CONTENT

Supporting Information. Accuracy tests for the sparse solvers, Ni K-edge simulation for the model NiO compound, scaling details of the FDMNES and the effect of different exchange-correlation functional on Fe K-edge XANES can be found in the supplementary materials. This information is available free of charge via the Internet at http://pubs.acs.org/

The software FDMNES (including Fortran code and executables) can be downloaded free of charge from http://neel.cnrs.fr/spip.php?article3137

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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. S.A. Guda and Y. Joly introduced sparse solvers to the FDMNES code, A.A. Guda and G. Smolentsev supervised the import of the electron density to the finite difference method from the MO-DFT, A.A. Guda, K.A. Lomachenko, A.L. Bugaev and M.A. Soldatov performed the testing of the new version of code and calculated XANES spectra presented in the paper, C. Lamberti and A.V. Soldatov provided experimental data and supervised the simulations on CPO-27-Ni MOF, W. Gawelda and C. Bressler provided experimental data and supervised the simulations on different spin states of $[Fe(bpy)_3]^{2+}$.

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ABBREVIATIONS

DFT, density functional theory; MO - molecular orbitals; FDM, finite difference method; XANES, X-ray absorption near-edge structure; SCF, self-consistent field simulations; GGA-PBE, generalized gradient approximation, Perdew, Burke, Ernzerhof; LS, low-spin state; HS, high spin state; MPI, Message Passing Interface; HOMO, highest occupied molecular orbital; LUMO, lowest unoccupied molecular orbital.

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<u>Synopsys</u>: A sparse matrix diagonalization scheme was applied to the FDMNES code [*Phys. Rev. B*, **63**, 125120 (2001)]. The required calculation time for XANES spectrum was reduced by more than one order of magnitude compared to the standard Gaussian method and efficient multicore parallelization is now available. Thus structural analysis of complex systems can be performed routinely on a modern personal computer. We analyzed the bending angle between CO and NO molecules attached to Ni active site in the CPO-27-Ni metal-organic framework and the ultrafast low spin (LS) to high spin (HS) conversion of photoexcited aqueous $[Fe(bpy)_3]^{2+}$, which - next to the initial and final states - also includes the XANES calculation for the recently identified transient triplet state.