Core electron excitations in U$^{4+}$: modelling of the $nd^{10}5f^2 \rightarrow nd^{9}5f^3$ transitions with $n = 3, 4$ and 5 by ligand field tools and density functional theory†

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Ligand field density functional theory (LFDFT) calculations have been used to model the uranium $M_{4.5}, N_{4.5}$ and $O_{4.5}$-edge X-ray absorption near edge structure (XANES) in $UO_2$, characterized by the promotion of one electron from the core and the semi-core 3d, 4d and 5d orbitals of $U^{4+}$ to the valence 5f. The model describes the procedure to resolve non-empirically the multiplet energy levels originating from the two-open-shell system with d and f electrons and to calculate the oscillator strengths corresponding to the dipole allowed $d^{10}f^2 \rightarrow d^{9}f^3$ transitions appropriate to represent the $d$ electron excitation process. In the first step, the energy and $UO_2$ unit-cell volume corresponding to the minimum structures are determined using the Hubbard model (DFT+$U$) approach. The model of the optical properties due to the uranium $nd^{10}5f^2 \rightarrow nd^{9}5f^3$ transitions, with $n = 3, 4$ and 5, has been tackled by means of electronic structure calculations based on the ligand field concept emulating the Slater–Condon integrals, the spin–orbit coupling constants and the parameters of the ligand field potential needed by the ligand field Hamiltonian from Density Functional Theory. A deep-rooted theoretical procedure using the LFDFT approach has been established for actinide-bearing systems that can be valuable to compute targeted results, such as spectroscopic details at the electronic scale. As a case study, uranium dioxide has been considered because it is a nuclear fuel material, and both atomic and electronic structure calculations are indispensable for a deeper understanding of irradiation driven microstructural changes occurring in this material.

Introduction

A great deal of research on actinide compounds has been carried out, with both fundamental understanding and possible applications in sight.1–5 Actinide isotopes (such as Th, U and Pu) are radioactive with a wide range of nuclear properties. Their physical

...as well as chemical properties are often investigated because of their industrial use in the process of generating electric energy from nuclear power plants and the safe disposal of the spent nuclear fuel. The interest in actinide compounds is also a matter of tremendous investigations using computational modelling and simulations that help overcome the experimental challenges in handling radioactive materials, and contribute to the fundamental understanding of structural and coordination chemistry of actinide compounds.

Understanding the crystal chemistry and properties of actinide compounds requires a detailed insight into the final electronic distributions and character of the 5f electrons in the ground or excited configurations of the free actinide atoms. The structural, spectroscopic and electronic properties of actinide compounds have been widely studied both theoretically and experimentally.6–10 Obviously, derivation of energy levels from experimental data is often a challenging task due to a multiplet electronic structure. For example the X-ray absorption and/or photoemission spectra of actinide based compounds feature multiple electron correlation effects in a defined configuration.11,12 Therefore it is practically not simple to characterize an optical spectrum without the use of a...
phenomenological model, which is able to illustrate the important features due to electronic transitions.

Ligand field theory is a very successful concept for the description of the bonding regime and the electronic properties of rare-earth ions in coordination chemistry.13 It is a phenomenological model, which at the very beginning employed only empirical parameters.14,15 Nowadays, the premise of ligand field theory has evolved being fully implemented in regular first principles quantum chemistry codes.16–18 Although the principle of the model has not been subjected to any change, keeping its phenomenological origin, the nature of the parameters has been reformed, being derived by a purely non-empirical scheme. The ligand field Density Functional Theory (LFDFT) algorithm is one of these codes, aiming reasonably to compute the multiplet electronic structure, using Density Functional Theory (DFT) based calculations.16–18 In the last three years, the LFDFT has been used to study two-open-shell electronic structures, involving at the same time f and d electrons. The results, when applied to the luminescence of lanthanide compounds, were always comparable with the experimental findings.19–22 Thus LFDFT can be considered as a valuable non-empirical tool to evaluate the electronic structure and optical properties in coordination chemistry.

In the following, we present LFDFT calculations of the electronic structure and the optical properties of U4+ in uranium dioxide (UO2). UO2 is a well-known material extensively used as fuel in the nuclear industry.23–25 Its investigation is therefore of crucial importance and comprises the bulk as well as the intrinsic contributions of fission products in the matrix of irradiated spent fuel etc. In this work, our results will be limited to the description of only fresh UO2 materials. We consider the core and semi-core electron excitations originating successively from the 3d, 4d and 5d orbitals, and their mutual interaction with the strongly correlated 5f (the ground electron configuration of U4+). The multiplet energy levels and the oscillator strength of the electric dipole moment of the inter-configuration nd105f2 → nd35f4 transitions, with n = 3, 4 and 5, are determined. This, in turn, allows us to calculate the X-ray absorption near edge structure (XANES) at the uranium M4,5, N4,5 and O4,5 edges in UO2.

Methodology

The phenomenological two-open-shell ligand field Hamiltonian (H), suitable for describing the nd105f2 → nd35f4 transitions, with n = 3, 4 and 5, combines three independent entities,19–21

\[ H = H_{\text{ER}} + H_{\text{SO}} + H_{\text{EF}} \]  

(1)

where \( H_{\text{ER}} \), \( H_{\text{SO}} \) and \( H_{\text{EF}} \) represent the interactions due to electron–electron repulsion, spin–orbit coupling and ligand field splitting, respectively.

The matrix elements of all the entities in eqn (1) are expressed on the basis of single determinants of spin-orbitals. Therefore, the matrix elements of electron–electron repulsion (eqn (2)) consist of the product of the Slater-Condon parameters \( F \) and \( G \) with the coefficients \( f \) and \( g \), which depend on the angular quantum numbers of all the single determinants arising from the ground configuration nd105f2 and the excited nd35f4.

\[
H_{\text{ER}} = \sum_{k=0,2,4,6} F^k (\langle \Pi \rangle) f_k (\langle \Pi \rangle) + \sum_{k=0,2,4} F^k (df) f_k (df) 
+ \sum_{k=1,3,5} G^k (df) g_k (df)
\]  

(2)

The Slater–Condon parameters representing the Coulomb \( F \) and the exchange \( G \) interactions are theoretically defined as integration over a portion of the one-electron radial functions \( \{ k \} \) centred at the position of a U4+ ion (eqn (3)-(5)),26,27 whereas the coefficients \( f \) and \( g \) are independent of \( R \) but particular to any actinide ions belonging to the present nd105f2 and nd35f4 electron configurations.

\[
F^k (\langle \Pi \rangle) = \int_0^\infty \int_0^{r_2} R_{5f}^2 (r_1) R_g^2 (r_2) r_1^2 r_2^2 dr_1 dr_2
\]  

(3)

\[
F^k (df) = \int_0^\infty \int_0^{r_2} \nabla R_{5f}^2 (r_1) R_g^2 (r_2) r_1^2 r_2^2 dr_1 dr_2
\]  

(4)

\[
G^k (df) = \int_0^\infty \int_0^{r_2} \nabla \nabla R_{5f} (r_1) R_g (r_2) R_g (r_1) R_g (r_2) r_1^2 r_2^2 dr_1 dr_2
\]  

(5)

The matrix elements of the spin–orbit coupling interaction (eqn (6)) are obtained as the product of the one-electron spin-orbit coupling constant \( \zeta \) with coefficients \( d \), which depend on the spin and angular momentum operators defined in terms of spherical harmonics of order \( l = 3 \) and \( l = 2 \) for the f and d electrons, respectively.

\[
H_{\text{SO}} = \zeta_{5f} d_5 + \zeta_{4d} d_4
\]  

(6)

The matrix elements of the ligand field splitting are obtained from the product of the Wybourne-normalized crystal field parameters \( B \) and the spherical harmonic tensor operator \( C \) acting on the f and d orbitals of U4+. In total, there is a possible combination of 64 independent \( B \) parameters to compose the ligand field Hamiltonian.26 However in the case of the present cubic symmetry, there are only three parameters, which are taken into account (eqn (7)).

\[
H_{\text{LF}} = B_0^0 (\langle \Pi \rangle) \left[ C_0^{(4)} + \frac{\sqrt{5}}{14} (C_4^{(4)} + C_{-4}^{(4)}) \right]
\]

\[+ B_0^2 (\langle \Pi \rangle) \left[ C_0^{(6)} - \frac{\sqrt{5}}{2} (C_4^{(6)} + C_{-4}^{(6)}) \right] + B_0^4 (dd) \left[ C_4^{(4)} + \frac{\sqrt{5}}{14} (C_4^{(4)} + C_{-4}^{(4)}) \right]
\]  

(7)

The DFT calculations have been carried out by means of the Amsterdam Density Functional (ADF) program package (ADF2014.01).29–31 This code is a quantum chemistry code
enabling the run of average of configuration (AOC) type calculation needed by the LFDFT procedure. The AOC is a restricted DFT calculation imposing fractional occupation numbers on molecular orbitals assigned to d or f parentage. The hybrid functional B3LYP has been used to compute the electronic structure and the optical properties of the cubic cluster (UO$_3$)$_{12}^{2-}$ embedded in UO$_2$. A series of point charges are added in the position of the next nearest neighbour of (UO$_3$)$_{12}^{2-}$ in order to neutralize the high negative charge of the cluster and also to mimic the long-range interaction due to the periodicity of the UO$_2$ structure using a Madelung potential. The molecular orbitals were expanded using a quadrupole-zeta plus polarization Slater-type orbital (STO) functions (QZ4P) for the U atom and triple-zeta plus polarization STO functions (TZP) for the O atom. The self-consistent field (SCF) was set-up to take into account all electrons. The relativistic corrections were treated considering the Zeroth Order Regular Approximation (ZORA) of the Dirac equation as implemented in the ADF program package. The scalar and the spin–orbit parts of the ZORA Hamiltonian were used, which provide an accurate description of both the spin-independent and spin-dependent relativistic effects inherent to heavy elements such as uranium.

The equilibrium structure and energies of the pristine UO$_2$ were theoretically determined by means of DFT based periodical calculation implemented in the Vienna Ab initio Simulation Package (VASP). The exchange and correlation effects are described using the local density approximation (LDA) and the generalized gradient approximation (GGA) functionals. The following valence configurations have been considered: 6s$^2$7s$^2$6p$^6$6d$^2$5f$^2$ and 2s$^2$2p$^4$, respectively for U and O atoms. The interaction between these valence electrons and the core ones was emulated using the projected augmented wave method. A plane wave basis set with a cut-off energy of 520 eV was used. 6-k points were included in each direction of the lattice according to the Monkhorst-Pack scheme.

Results and discussion

The oxidation state of the uranium ion in UO$_2$ is formally +4. The tetravalent uranium ion is prone to a single-open-shell electronic structure, which exists in its ground state by the population of 2 electrons in the seven-fold degenerate 5f orbitals. Although we may believe that UO$_2$ has a metallic character due to this partially filled 5f shell, UO$_2$ belongs to a special class of insulating materials being a Mott-insulator. Experimentally, the band gap of UO$_2$ is known to be circa 2 eV and it is predominantly characterized by the 5f$^2$–5f$^2$ transitions. The 5f orbitals of U in UO$_2$ interact weakly with the ligand environment (the oxygen 2p orbitals). In this perspective, the calculated DFT band structure based on the LDA or GGA functional frequently gives a metallic ground state regardless of the antiferromagnetic interaction observed in the experiments below the Neel temperature. A realistic description of the properties is nonetheless achieved, while the computational set-up incorporates the Hubbard correction via the DFT+U formalism. Recently, it has also been shown that the DFT band structure algorithm operated using a hybrid functional offers a good alternative to represent correctly the ground state properties of UO$_2$. In this work, we realize the optimisation of the lattice parameters of UO$_2$ using the early concept defined in the DFT+U methodology. Based on previous theoretical studies, the Coulomb (U) and exchange (J) parameters have been set to the values of 4.5 eV and 0.5 eV, respectively. These values have been validated, which provide a realistic description of the electronic and magnetic structures of UO$_2$. The definition of the U parameter itself cannot be rationalized, since U is often deduced from the experimental data. On the other hand, we can justify the value of the J parameter with respect to the calculated Slater–Condon parameters that will be described later. Nevertheless, we are going to use the simplified formalism by taking only the difference U–J in the periodical calculation.

The crystal structure of UO$_2$ is graphically represented in Fig. 1, where the U and O atoms are located at their ideal fluorite positions. The U atoms are eight-fold coordinated with the O atoms (Fig. 1) in a cubic arrangement and these latter ones are four-fold coordinated with the U atoms in a tetrahedral symmetry. Besides, distortion of the oxygen cage is possible due to the Jahn–Teller effect. This is observed in neutron diffraction experiments, but it is neglected in this work conserving the high symmetry of the molecular cluster (UO$_3$)$_{12}^{2-}$ (Fig. 1) embedded in UO$_2$. Within the fluorite structure, the lattice a, b and c constants are all equivalent and the angles α, β and γ are equal to 90°. The unit-cell volume V represents the product a$\times$ b$\times$ c = a$^3$. We use the traditional representation of the unit-cell of UO$_2$ by means of twelve atoms in a cell, including four U and eight O atoms. The magnetic interaction due to the valence electrons of the U atoms is approximated using the collinear 1k antiferromagnetic order alternating U centres along the c-axis of the unit-cell, in line with previous theoretical descriptions of UO$_2$. The iterations in the self-consistent field process are therefore spin-unrestricted but the global spin-polarization is fixed to zero.

The total electronic energy is calculated with respect to the previously described conditions by varying the volume of the...
Table 1 Optimized unit-cell volume $V$ (in Å$^3$), lattice constant $a$ (in Å), and calculated bulk modulus $B$ (in GPa) of UO$_2$ obtained at the LDA+U and GGA+U levels of theory, compared with the available experimental data (Exp.); together with the calculated distances $d$ (in Å) between U and O and two nearest U atoms.

<table>
<thead>
<tr>
<th>LDA+U</th>
<th>GGA+U</th>
<th>Exp.</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V$</td>
<td>162.23</td>
<td>170.94</td>
</tr>
<tr>
<td>$a$</td>
<td>5.4540</td>
<td>5.5499</td>
</tr>
<tr>
<td>$B$</td>
<td>221.42</td>
<td>197.15</td>
</tr>
<tr>
<td>$d_{U-O}$</td>
<td>2.3617</td>
<td>2.4032</td>
</tr>
<tr>
<td>$d_{U-U}$</td>
<td>3.8566</td>
<td>3.9244</td>
</tr>
</tbody>
</table>

$^a$ Taken from ref. 63 and 64. $^b$ Taken from ref. 59.

unit-cell. The energy and unit-cell volume corresponding to the minimum structures are determined via fitting of the DFT results with a second order polynomial function using the Birch–Murnaghan equation of state (see the ESI† Fig. S1). The bulk properties of UO$_2$ obtained in the present DFT calculations are collected in Table 1, together with the corresponding values of experimental results reported in the literature. In general both LDA+U and GGA+U calculations yield a realistic description of the unit-cell of UO$_2$, since the difference in the unit-cell volume, if compared to the experimental results, represents a deviation of less than 5%. The calculated unit-cell volumes are in line with previous theoretical results based on LDA+U$^{23}$ and GGA+U$^{69,73}$ calculations. We note that if we carry out a full relaxation of the structure of UO$_2$ using the present computational details without symmetry constraint, we obtain a tetragonal structure characterized by a shortening of the lattice constant $c$. This point is also often mentioned in the literature, which is essentially due to the approximation of the magnetic interaction of the uranium valence electrons by a collinear $1k$ order antiferromagnetic instead of the non-collinear $3k$ one. The calculated bulk modulus $B$ is comparable with the experimental results, even if it is observed that the LDA+U as well as the GGA+U calculations (Table 1) tend to overestimate and to underestimate it, respectively. We consider the LDA+U results to be the most suitable to represent the bulk properties of UO$_2$, since their deviation with respect to the experiments is the smallest (Table 1). Therefore, the difference in the unit-cell constant $\Delta a$ (between the LDA+U calculation and the experimental results) does not represent a significant change in the U–O and U–U bond distances (see Table 1) in the bulk UO$_2$, which is a crucial prerequisite for the consideration of the molecular cluster (UO$_8$)$^{12-}$ embedded in UO$_2$ described in the next section.

The model of the optical properties due to the uranium $n d^{10}5f^n \to n d^{9}5f^{n+1}$ transitions, with $n = 3, 4$ and 5 is tackled here by means of multiplet energy level calculations based on the ligand field concept with two-open-shell configurations of $d$ and $f$ electrons. It is necessary to take a molecular cluster representing the U centre and its ligand environment instead of the full periodical model. This molecular cluster consists of one uranium centre plus eight oxygen ligands placed in the vertices of a regular cube (see Fig. 1). The bond length between U and O is fixed to the LDA results given in Table 1. Moreover the long-range influence of the UO$_2$ lattice is reproduced by adding a series of point charges placed at the crystallographic positions of the U and O atoms in UO$_2$. We have carried out DFT calculations using the ADF suite of programs.$^{29–31}$

The first step comprises a DFT calculation considering the ground electron configuration of U$^{4+}$, i.e. 5f$^2$. Since the Wyckoff position of the U site is octahedral, the 5f orbitals split by the ligand field interaction (eqn (7)) into three distinct energy levels forming the basis of the $a_{2u}$, $t_{2u}$ and $t_{1u}$ irreducible representations of the $O_h$ point group. The AOC set-up occupies each component of the 5f orbitals with a fractional electron occupation of 2/7.$^{24}$

In this situation, the density belongs to the totally symmetric irreducible representation of the $O_h$ point group under which the ligand field Hamiltonian (eqn (1)) is invariant. The three-fold degenerate $t_{1u}$ level has theoretically the lowest energy, whereas the $a_{2u}$ is strongly destabilized partly due to the specific cubic arrangement of the eight O ligands.

The splitting pattern of the 5f orbitals of U$^{4+}$ in the cluster (UO$_8$)$^{12-}$ embedded in the bulk UO$_2$ is graphically represented in Fig. 2 together with the molecular orbital scheme as a result of linear combination of atomic orbitals.

The second step consists of carrying out DFT calculations taking into account the core or semi-core electron excitations originating from the 3d, 4d and 5d orbitals of U$^{4+}$ to the partially filled 5f levels. We deal with the electron configurations nd$^3$5f$^n$ with $n = 3, 4$ and 5, respectively. In these situations the AOC includes two-open-shell systems with $d$ and $f$ electrons, which populates 3/7 electrons in each component of the 5f orbitals (Fig. 2), plus 9/5 electrons in each component of the 3d or 4d or 5d orbitals, respectively.$^{21}$ In the $O_h$ point group, these d orbitals transform into the $t_{2u}$ and $e_g$ irreducible representations. We stress that the splitting of the 3d, 4d and 5d orbitals, as obtained theoretically here, is relatively small and will be completely superseded by the effect of the spin–orbit coupling interaction.
To find the ligand field potential in the definition as given by eqn (7), we use the eigenvalues of the Kohn–Sham orbitals with dominant 5f and nd (with n = 3, 4 and 5) character obtained from the AOC calculations. These Kohn–Sham orbital energies match with the quantities \( \langle a_{2u} | H_{\text{LF}} | a_{2u} \rangle \), \( \langle t_{2u} | H_{\text{LF}} | t_{2u} \rangle \), and \( \langle t_{2u} | H_{\text{LF}} | t_{1u} \rangle \) representing the eigenvalues of the 5f ligand field (eqn (7)), and \( \langle t_{2u} | H_{\text{LF}} | t_{2g} \rangle \) and \( \langle c_{g} | H_{\text{LF}} | c_{g} \rangle \) of the nd ligand field. Therefore, one can deduce the ligand field parameters \( B_{0}^{f}(f) \) and \( B_{0}^{d}(dd) \) from DFT as follows:

\[
B_{0}^{f}(f) = \frac{39}{40} \left( \langle a_{2u} | H_{\text{LF}} | a_{2u} \rangle + 4 \langle t_{2u} | H_{\text{LF}} | t_{2u} \rangle \right) \tag{8}
\]

\[
B_{0}^{d}(f) = \frac{36}{11(2 + \sqrt{2})} \left( \langle a_{2u} | H_{\text{LF}} | a_{2u} \rangle + 2 \langle t_{1u} | H_{\text{LF}} | t_{1u} \rangle \right) \tag{9}
\]

\[
B_{0}^{d}(dd) = \frac{21}{10} \left( \langle c_{g} | H_{\text{LF}} | c_{g} \rangle - \langle t_{2g} | H_{\text{LF}} | t_{2g} \rangle \right) \tag{10}
\]

One may note that \( H_{\text{LF}} \) (eqn (7)) is a traceless matrix (i.e. \( B_{0}^{f}(f) = 0 \) and \( B_{0}^{d}(dd) = 0 \)). Therefore, the orbital energies extracted from DFT were adjusted around the barycenter of the 5f orbitals, and also for the nd orbitals. The mixed term parameters due to the ligand field related to the d-f interaction are by definition zero since the f and the d orbitals possess different parity in the \( O_{h} \) point group.

The spin–orbit coupling constants \( \xi \) are obtained by means of ZORA spin–orbit calculations. For the 3d, 4d and 5d electrons, \( \xi_{3d}, \xi_{4d} \) and \( \xi_{5d} \) are determined from the energy difference between the \( d_{1/2} \) and \( d_{3/2} \) components of the d orbitals: the ligand field interaction (eqn (7)) being negligible (see Table 2). For the 5f electrons, \( \xi_{5f} \) is obtained by mapping the DFT based ZORA spin–orbit results with the matrix elements of the spin–orbit coupling interaction (eqn (6)) plus the ligand field potential (eqn (7)). In previous studies, we have presented the details of the mapping algorithm and involved procedures to estimate \( \xi \), illustrating the influence of the ligands on the spin–orbit coupling in particular that the coupling constant may vary notably with the orbital energy. The 5f orbitals in \( O_{h} \) symmetry split into six spinors (spin–orbit terms), as the molecular orbitals \( t_{1u}, t_{2u} \) and \( a_{2u} \) (see Fig. 2), respectively, transform to \( g_{2u}, e_{12u}, e_{52u} \) and \( c_{g} \) irreducible representations of the \( O_{h}^{*} \) double group. Therefore, using symmetry analysis we have obtained a set of spin–orbit coupling constants. Thus, the calculated spin–orbit coupling constant of 5f electrons reported in Table 2 is an averaged value used for the ligand-field calculation. The calculated \( \xi_{5f} \) value in Table 2 is in agreement with the spin–orbit coupling constant reported in ref. 79, which was derived for the linear \( UO_{2} \) molecule (uranium 5f\(^{2}\) configuration) using CASPT2. We also note that the parameter \( \xi_{5f} \) changes from 2 to 3-electron population of the uranium 5f orbitals (see Table 2), in agreement with previous studies based on four component relativistic calculations of americium in \( AmO_{5}^{3+} \) (with 5f\(^{2}\) configuration), \( AmO_{5}^{3+} \) (5f\(^{2}\)) and \( AmO_{5}^{3+} \) (5f\(^{2}\)) molecular systems. The Slater–Condon integrals in eqn (3)–(5) are calculated by means of the radial functions of the Kohn–Sham orbitals with dominant 5f and 3d, 4d, and 5d character. A graphical representation of the radial functions of 5f Kohn–Sham orbitals is shown in Fig. 3. The corresponding results of nd orbitals are available in the ESI† (see Fig. S2). According to eqn (3), there are four independent parameters given by the four different k values: 0, 2, 4 and 6. However according to eqn (4) and (5), there are three independent parameters discriminated by \( k = 0, 2, 4 \) and \( k = 1, 3, 5 \), respectively for the \( F \) and the \( G \) integrals. In total, there are ten Slater–Condon integrals to be considered in the calculation of the \( nd^{10}5f^{2} \rightarrow nd^{9}5f^{3} \) transitions, with \( n = 3, 4 \) and 5. We may consider four different situations, giving rise to the series of mentioned parameters tabulated in the four columns of Table 2.

In the ground state of \( U^{4+} \) i.e. the single-open-shell 5f\(^{2}\) electron configuration, the Slater–Condon parameters corresponding to the mixed d-f terms are not important. Accordingly, the spin–orbit coupling constant corresponding to the d shell is ignored as well as the parameter for the ligand field potential of

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
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<tr>
<td>( F_{0}(f) )</td>
<td>0.0235</td>
</tr>
<tr>
<td>( F_{2}(d) )</td>
<td>2.9737</td>
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<tr>
<td>( F_{4}(d) )</td>
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<tr>
<td>( H_{2}(d) )</td>
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<td>( H_{4}(d) )</td>
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Table 2 Calculated parameters (in eV) corresponding to the Slater–Condon integrals, the spin–orbit coupling constants and the ligand field potential of \( U^{4+} \) in the molecular cluster \( (UO_{2})^{2-} \) - embedded in \( UO_{2} \); obtained for the ground electron configuration 5f\(^{2}\) as well as the core and semi-core electron excitation 3d\(^{5}5f^{2}\), 4d\(^{5}5f^{2}\), 5d\(^{5}5f^{2}\) configurations

5f Kohn–Sham orbitals are shown in Fig. 3. The corresponding results of nd orbitals are available in the ESI† (see Fig. S2). According to eqn (3), there are four independent parameters given by the four different k values: 0, 2, 4 and 6. However according to eqn (4) and (5), there are three independent parameters discriminated by \( k = 0, 2, 4 \) and \( k = 1, 3, 5 \), respectively for the \( F \) and the \( G \) integrals. In total, there are ten Slater–Condon integrals to be considered in the calculation of the \( nd^{10}5f^{2} \rightarrow nd^{9}5f^{3} \) transitions, with \( n = 3, 4 \) and 5. We may consider four different situations, giving rise to the series of mentioned parameters tabulated in the four columns of Table 2.

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![Graphical representation of U\(^{4+}\) radial functions (5f\(^{2}\) Kohn–Sham orbitals) in UO\(_{2}\) electron configuration 5f\(^{2}\) (in blue), 3d\(^{5}5f^{2}\) (in magenta), 4d\(^{5}5f^{2}\) (in black) and 5d\(^{5}5f^{2}\) (in green). The corresponding result for a free U\(^{4+}\) ion is also shown (in red).](image-url)
the d–d interaction. Then, we have the three situations where one electron is promoted from the 3d, 4d or 5d shells to the 5f. In all three cases, the mixed Slater–Condon parameters do not vanish anymore, as they are important to represent the interaction between the two-open-shell electron configurations. On the other hand, the Slater–Condon parameters representing the f–f terms are not comparable with the parameters obtained for the ground configuration. They must be different because they take into account the presence of three electrons instead of the earlier two ones on the emulation of the radial function of the 5f Kohn–Sham orbitals. In the calculation of the multiplet energy levels, they will be present in the diagonal element of the ligand field Hamiltonian (eqn (1)) corresponding to the \( nd^3f^6 \) (with \( n = 3, 4, \) and 5) matrix interaction.

The phenomenological Hamiltonian in eqn (1) can be represented by a block matrix of 3731 times 3731 elements. By definition, it is a complex and Hermitian matrix composed by a sub-block diagonal of 91 times 91 elements corresponding to the ground \( 5f^6 \) configuration of \( U^{4+} \) and a sub-block diagonal of 3640 times 3640 elements suitable for the \( d^95f^3 \) configuration. The sub-block of the \( 5f^6 \) interactions is parameterized by only the ground state values in Table 2, whereas the sub-block of the d–f interactions may take different values in Table 2 according to \( n = 3, 4 \) and 5, respectively.

Considering the ground state configuration, in terms of Russell-Saunders coupling, two electrons in the 5f orbitals induce seven spectral terms with different energy levels. They are constituted by the high spin \( ^3P, ^3F \) and \( ^3H \) states as well as the low spin \( ^1S, ^1D, ^1G \) and \( ^1I \) states. Hund’s rules allow the determination of the ground state, in this case \( ^3H \). By inclusion of the spin–orbit coupling interaction (eqn (6)), all the spin-triplet states split into energy levels discriminated by their \( \tilde{J} \) values namely, \( ^1P_0, ^1P_1, ^1P_2, ^3F_0, ^3F_2, ^3H_4, ^3H_5 \) and \( ^3H_6 \). Together with the earlier spin-singlet states, it results in total thirteen states, in which the ground state becomes \( ^3H_4 \) by applying Hund’s rules once more. Finally the ligand field interaction, which accounts for the presence of oxygen ligands, splits all the spectral terms into various ligand field levels. The present octahedral coordination allows in total forty energy levels.

The values of the parameters in Table 2 are comparable with the experimental findings, although it is observed that the DFT slightly overestimates some values. We can refer to the experimental results in ref. 81 and 82 corresponding to the spectroscopy of the free \( U^{4+} \) ion. The best reproduction of the energies of the thirteen spectral terms suitable for the free ion is achieved by the parameters (in eV): \( F^4(\mathbf{ff}) = 6.44, F^4(\mathbf{ff}) = 5.30, F^4(\mathbf{ff}) = 3.44 \) and \( \zeta_{\mathbf{ff}} = 0.24 \). These parameters will be reduced by the nephelauxetic effect when \( U^{4+} \) is being coordinated with ligands, for instance in the present UO\(_2\) case. In this context we can refer to the magneto-optical experiments of UO\(_2\) in ref. 84 and 85, where the following Slater–Condon and spin–orbit coupling parameters (in eV): \( F^4(\mathbf{ff}) = 5.34, F^4(\mathbf{ff}) = 4.57, F^4(\mathbf{ff}) = 3.64 \) and \( \zeta_{\mathbf{ff}} = 0.22 \) are deduced. The overestimation due to DFT is mainly visible for \( F^4(\mathbf{ff}) \), which is also accompanied by an underestimation of \( F^4(\mathbf{ff}) \) and \( F^4(\mathbf{ff}) \) in ref. 86, the experimentally deduced ligand field parameters are \( -3.080 \text{ eV} \) for \( B^4(\mathbf{ff}) \) with a fixed value for the ratio \( B^4(\mathbf{ff})/B^4(\mathbf{ff}) \). On the other hand, the inelastic neutron scattering experiment in ref. 87 yields \( B^4(\mathbf{ff}) = -0.984 \text{ eV} \) and \( B^4(\mathbf{ff}) = 0.424 \text{ eV} \). It is clear that the reproduction of the experimental \( B^4(\mathbf{ff}) \) parameter by DFT (Table 2) is realized, whereas \( B^4(\mathbf{ff}) \) is in the interval of two different experimental values. The calculated parameters corresponding to the \( d^35f^6 \) configurations (Table 2) are comparable to the values related to the d–f transitions obtained from Hartree–Fock calculations of \( U^{5+} \) in ref. 88.

The energies of the multiplet levels arising from the \( 5f^6 \) configuration of \( U^{5+} \) in UO\(_2\) using the cluster model (UO\(_8\))\(_{12}^+\) are calculated and graphically represented in Fig. 4. We present three series of energy levels in Fig. 4 representing first the influence of the inter-electron repulsion \( H_{ER} \) (eqn (2)), secondly the influence of the sum of \( H_{ER} \) and the spin–orbit coupling interaction \( H_{SO} \) (eqn (6)) and finally the influence of the sum of \( H_{ER} \) and \( H_{SO} \) and the ligand field interaction \( H_{LF} \) (eqn (7)). This is important in order to show the evolution of the spectral term energy with respect to the three components, which constitute the Hamiltonian in eqn (1). The highest energy level obtained for the \( 5f^6 \) corresponds to the single degenerate \( ^3S_0 \) state with a calculated energy of 5.504 eV (Fig. 4). For information, in free-ion spectroscopy this state is experimentally found to have an energy of 5.407 eV. The spectral terms are identified and labelled for the \( H_{ER} \) multiplets as well as for the \( H_{ER} + H_{SO} \) (Fig. 4). However, the labelling of the spectral terms due to \( H_{ER} + H_{SO} + H_{LF} \) is omitted for clarity in Fig. 4. The forty states obtained at this stage form the basis of the following representations: \( A_1 \) (seven times), \( A_2 \) (three times), \( E \) (nine times), \( T_1 \) (nine times) and \( T_2 \) (twelve times). Selected energy levels corresponding to them are collected in Table 3 (see also the ESI† Table S1 for the whole manifold of the ligand field multiplet of the \( 5f^6 \) configuration).

In Table 3, the calculated multiplet energy levels of the uranium \( 5f^6 \) configuration in UO\(_2\) are compared with some...
Table 3: Selected theoretical (Calc.) energy levels (in eV) arising from the ground 5f² electron configuration of U⁴⁺ in the cluster (UO₄)₂²⁻ embedded in UO₂, together with three reference data (Exp.) based on fit to the experiments.

<table>
<thead>
<tr>
<th>5f² levels</th>
<th>Calc.</th>
<th>Exp.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1⁰</td>
</tr>
<tr>
<td>T₂</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>E</td>
<td>0.1805</td>
<td>0.1504</td>
</tr>
<tr>
<td>T₁</td>
<td>0.2714</td>
<td>0.1667</td>
</tr>
<tr>
<td>A₁</td>
<td>0.3195</td>
<td>0.1748</td>
</tr>
<tr>
<td>T₁</td>
<td>0.6434</td>
<td>—</td>
</tr>
<tr>
<td>E</td>
<td>0.7484</td>
<td>—</td>
</tr>
<tr>
<td>T₂</td>
<td>0.7640</td>
<td>—</td>
</tr>
<tr>
<td>E</td>
<td>0.8518</td>
<td>—</td>
</tr>
<tr>
<td>T₂</td>
<td>0.9375</td>
<td>—</td>
</tr>
<tr>
<td>T₂</td>
<td>0.9979</td>
<td>—</td>
</tr>
<tr>
<td>T₁</td>
<td>1.1083</td>
<td>—</td>
</tr>
<tr>
<td>A₁</td>
<td>1.1708</td>
<td>—</td>
</tr>
<tr>
<td>T₁</td>
<td>1.2112</td>
<td>—</td>
</tr>
<tr>
<td>A₁</td>
<td>1.2153</td>
<td>—</td>
</tr>
</tbody>
</table>

a Taken from ref. 87. b Taken from ref. 84. c Taken from ref. 89.

Theoretical and experimental results available in the literature. The multiplet energy levels in the second column of Table 3 have been computed by diagonalizing the ligand field Hamiltonian (eqn (1)) with a ground state (lowest T₂ state) energy correction. Therefore, these energy values represent the vertical excitation energies of the ground T₂ state, hence, can be compared with the corresponding excitation energies measured in the experiments. The measured energy values in the third column of Table 3 are associated with T₂ → E, T₂ → T₁, and T₂ → A₁ transitions reported in ref. 87. These energy values have been obtained by comparing the neutron energy gain or loss spectra (acquired in inelastic high-resolution neutron scattering experiments) with the detailed ligand field calculations. The fourth column in Table 3 shows the transition energies as determined using ligand field calculations, which also corroborate the magnetic susceptibility measurements in UO₄. The last column in Table 3 lists the excitation energies obtained from optical absorption data measured by probing the intra 5f² → 5f² transitions and multi-phonon excitations in UO₂. Although the ligand field model has always been considered to interpret the experimental data in the studies highlighted above, it becomes also evident from Table 3 that all energy values are not throughout in good agreement with each other. In particular, we note that the energy values in ref. 84 and 89 of T₂ → T₁ and T₂ → A₁ transitions differ significantly when compared to those presented in ref. 87. The exact reason for this discrepancy is unclear. We believe that it is the result of methodological differences between studies under different experimental conditions. Moreover, the data in third to fifth columns of Table 3 are estimates of a model’s parameters based on the evaluation of fits to experimental data with theoretical functions. Therefore, it seems more reasonable to consider the results of the high-resolution neutron scattering experiment in the study by Amoretti and co-workers, which would be best suited to validate our LFDF results. The most important aspect is the splitting of the ground state of the free ion (H₄⁺) in terms of ligand field energy, where by group theoretical considerations we obtain four levels: A₁, E, T₁, and T₂. The calculation designates T₂ as the ground state (Table 3) in line with the above referenced studies.

Considering the d⁹f² electron configuration, we address the problem as the product of the multiplets of f⁹ configuration with a d electron. This is equivalent to the product of f⁹ with a d hole, i.e. d⁹ in this work. The parent f⁹ configuration leads to spectral terms with quartet and doublet spin multiplicity, forming the basis of 364 single determinants. When we add the d electron (or equivalently the d hole), we obtain 158 spectral terms, which form the basis of 364 single determinants. The resulting spectral terms of the d⁹f² configuration are then: 5S, 5P (two times), 5D (four times), 5F (three times), 5G (four times), 5H (three times), 5I (two times), 5K, 5L, 5S (three times), 5P (seven times), 5D (eleven times), 5F (twelve times), 5G (thirteen times), 5H (eleven times), 5I (nine times), 5K (six times), 5L (four times), 5M (two times), 5N, 5S (two times), 5P (five times), 5D (seven times), 5F (nine times), 5G (nine times), 5H (eight times), 5I (seven times), 5K (five times), 5L (three times), 5M (two times) and 5N.

The aforementioned spectral terms originate from the inter-electron repulsion represented by eqn (2), but they are split by the spin–orbit coupling (eqn (6)) and the ligand field interaction (eqn (7)) in the same manner as the process already described for the 5f² configuration (see Fig. 4). We note that the variation of the Slater–Condon parameters F⁹(ff) in the series 3d⁹5f², 4d⁹5f² and 5d⁹5f² configurations is not significant (Table 2). However the mixed term Slater–Condon for d–f interactions increases in the same series showing the relative interaction between the core d shells and the valence 5f. This interaction is relatively weak considering the 3d–5f situation, whereas it becomes globally important in the cases of 4d–5f and 5d–5f. Therefore, multiplet levels are the main contributors to the XANES spectra of UO₂ at the uranium M₄,₅, N₄,₅ and more importantly at the O₂₃ edges, their spectral features may not be reflected in the experimental spectrum.

The multiplet energy levels of the 3d⁹5f², 4d⁹5f², and 5d⁹5f² configurations of U⁴⁺ in UO₂ are calculated by diagonalizing the ligand field Hamiltonian (eqn (1)) with a ground state energy correction corresponding to the lowest T₂ state of the 5f² configuration (see Table 3). The results are graphically represented in Fig. 5, 6 and 7, respectively (in the ESI, the numerical data corresponding to these energy levels are available in the Tables S2–S4).

The DFT calculations yield relatively larger values of 5d₄ and 5d₄ (see Table 2), as has also been underlined previously in the literature. The d orbitals split into two distinct components: d₃/₂ and d₅/₂. This, in turn, leads the multiplet energy levels separated into two blocks as illustrated in Fig. 5 and 6. The two blocks, which are shown in the upper-panel and lower-panel of the figures, represent the excitations of one electron from the d₃/₂ and d₅/₂ of U⁴⁺, respectively. In Fig. 7, the value of 5d₄ (see Table 2) is not large enough to induce the same earlier
graphical representations insofar as all the multiplet levels are
gathered in the same energy interval.

The core d level X-ray absorption mechanisms are in principle
initiated by the $d^{10}f^2 + h\nu \rightarrow d^9f^3$ transitions, which we have
resumed here by the simpler notation: $d^{10}f^2 \rightarrow d^9f^3$. Because of
the mixing between the d and f electrons, the $d^{10}f^2 \rightarrow d^9f^3$
transitions fulfil the electric dipole moment selection rule, giving
rise to strong line intensities in the absorption spectra. It is also
possible to include the electric-quadrupole transitions and/or the
magnetic-dipole transitions in our calculations. But one may note
that such transitions may result in very small intensities relative
to the electric dipole allowed transitions in a measured X-ray
absorption spectrum. For UO$_2$, quadrupole transitions
involve electron transfers $3d^{10}5f^2 \rightarrow 3d^{10}5f^26s^1$ and $3d^{10}5f^2 \rightarrow
3d^{10}5f^6d^1$ in three-open-shell configurations. In this work, our
calculations are restricted to the systems with only two-open-shell
of d and f electrons. In future work, we plan to extend the LDFDFT
approach to include the calculations of electron configurations
with more than two non-equivalent electrons (i.e. ligand field
with three-open-shell systems). The multiplet energy levels arising
from such multiple electron configurations will also be able to
predict for instance possible shake-up satellites in the absorption
spectra.

To model the line intensities, we use the matrix element of
the electric dipole moment operator $D$ (eqn (11)). $D$ consists
of the product between a radial integral, involving the radial
functions $R$ of Kohn–Sham orbitals of dominant f and d
characters, and an angular integral, which can be associated
with the Clebsh–Gordan coefficients.

$$D_{\mu} = \sqrt{4\pi/3} \langle R_{\text{rad}} \mid R_{\text{hl}} \rangle \langle Y_{2,\mu} \mid Y_{1,\mu} \rangle Y_{3,\mu}$$

$D$ is important for the computation of the oscillator strengths
related to the $d^{10}f^2 \rightarrow d^9f^3$ transitions, by distributing its
elements over the whole manifold of the multi-electronic
configuration interactions obtained from the ligand field
Hamiltonian (eqn (1)). The oscillator strengths are calculated
taking as the initial state the three-fold degenerate $T_2$ level
originating from the $^3H_4$ state (Fig. 4) representative of the

![Fig. 5](image1.png)

Calculated multiplet energy levels (in blue colour) of the $3d^95f^3$
electron configuration of U$^{+4}$ in UO$_2$ (see also the ESI,† Table S2),
together with the intensities of the absorption $3d^{10}5f^2 (T_2) \rightarrow
3d^{10}5f^3$ transitions, i.e. oscillator strength (in black colour) representing
the uranium M$_5$ and M$_4$ edge X-ray absorption spectrum of UO$_2$ in the energy intervals 732–742 eV
(upper-panel) and 774–784 eV (lower-panel), respectively. The green curve
represents the oscillator strengths convoluted with Gaussian bands with a
width of 0.5 eV.

![Fig. 6](image2.png)

Calculated multiplet energy levels (in blue colour) of the $4d^95f^3$
electron configuration of U$^{+4}$ in UO$_2$ (see also the ESI,† Table S3),
together with the intensities of the absorption $4d^{10}5f^2 (T_2) \rightarrow
4d^{10}5f^3$ transitions, i.e. oscillator strength (in black colour) representing the uranium N$_5$ and N$_4$
edge X-ray absorption spectrum of UO$_2$ in the energy intervals 732–742 eV
(upper-panel) and 774–784 eV (lower-panel), respectively. The green curve
represents the oscillator strengths convoluted with Gaussian bands with a
width of 0.5 eV.

![Fig. 7](image3.png)
In Fig. S3, S4 and S5 (ESI†) the 5d\(^{9}\)5f\(^{3}\) transitions, respectively, in Fig. 5, 6 and 7. The green curve represents the oscillator strengths convoluted with Gaussian bands with a width of 0.5 eV.

Fig. 5 shows the multiplet energy levels of the 3d\(^{10}\)5f\(^{2}\) (T\(_{2}\)) → 3d\(^{9}\)5f\(^{3}\), 4d\(^{10}\)5f\(^{2}\) (T\(_{2}\)) → 4d\(^{9}\)5f\(^{3}\) and 5d\(^{10}\)5f\(^{2}\) (T\(_{2}\)) → 5d\(^{9}\)5f\(^{3}\) transitions, respectively, in Fig. 5, 6 and 7.

In these figures, the intensity plots along the ordinate are shown in arbitrary units, which are also normalized to unity either for the oscillator strengths or for the Gaussian convolution. The choice of the Gaussian convolution is motivated by the fact that we shall not intend to perform a full simulation of the X-ray absorption profile. Indeed such a simulation will require a special treatment of the broadening effect of the oscillator strengths, which depends on different factors, which are not always tractable using a pure first-principles method. Thus the simulation of the spectral profile consists of pure Gaussian broadening of the oscillator strengths, taking different values for the half-width parameter. Special treatment of the broadening effect of the oscillator absorption profile. Indeed such a simulation will require a special treatment of the broadening effect of the oscillator strengths, which depends on different factors, which are not always tractable using a pure first-principles method.

The three separate components of the total ligand field Hamiltonian (eqn (1)) are superimposed and summed to produce the total oscillator strength for the 5d\(^{10}\)5f\(^{2}\) (T\(_{2}\)) → 5d\(^{9}\)5f\(^{3}\) transitions. The described results demonstrate a dominant role of the inter-electron repulsion (ER) in determining the O4,5 edge XANES of UO\(_{2}\) (see the ESI† Fig. S5). They are situated in the energy scale at 96.0 eV, 104.6 eV and 112.5 eV. The highest energy peak possesses the most intense absorption (Fig. 7). The influence of the spin–orbit coupling interaction of the 5d electrons is weak and the spectrum is mainly governed by the electrostatic inter-electron d–f repulsion. This is furthermore illustrated in Fig. S6 provided in the ESI†. The three separate components of the total ligand field Hamiltonian (eqn (1)) are superimposed and summed to produce the total oscillator strength for the 5d\(^{10}\)5f\(^{2}\) (T\(_{2}\)) → 5d\(^{9}\)5f\(^{3}\) transitions. The described results demonstrate a dominant role of the inter-electron repulsion (ER) in determining the O4,5 edge XANES of UO\(_{2}\) (see the ESI† Fig. S6). Therefore it is impossible to ascribe with energy ranges the O3 and O4 edges in the absorption spectrum. The lowest intense peak is strongly dependent on the spin–orbit coupling constant for 3d electrons. In ref. 101, X-ray absorption spectroscopy and electron energy-loss spectroscopy have been employed to probe the electronic transitions from the 5d semi-core level of uranium in UO\(_{2}\). It is observed that the O4,5 absorption edges of UO\(_{2}\) consist of three intense peaks, in line with our theoretical findings. Although we notice a slight shift in the energy scale (~2 eV) from the DFT analysis, overall agreement between computational and experimental results is very encouraging.
The LFDFT results of UO$_2$ presented in this paper addressing the two-open-shell ligand field concept complement our recent work on the lanthanide systems, which shed light on the understanding of the relationship between electronic states and the XANES spectrum of actinide bearing compounds, and set the foundation for further work. A detailed study of uranium L$_{2,3}$ edges XANES spectra would require a three-open-shell ligand field concept involving p, f and d orbitals (i.e. the 2p$^5$3f$^6$ $\rightarrow$ 2p$^5$3f$^5$6d$^1$ transitions), will be the subject of a forthcoming report with the aim of gaining deeper insights into the structural as well as electronic properties of irradiated UO$_2$ fuels.

Conclusions

The demand for a theoretical description of the chemical, physical and crystallographic properties of uranium dioxide (UO$_2$) emerges from the fact that UO$_2$ is an actinide-isotope containing material for nuclear fuel applications. Besides, the electronic structure of UO$_2$ attracts scientific attention because of many quantum effects, associated with the f electrons in the conduction process, which are often not fully understood. The purpose of this work, yet independent of any experimental verification, is to simulate the uranium M$_{4,5}$, N$_{4,5}$ and O$_{4,5}$ edge XANES of UO$_2$ exploiting the power of the LFDFT approach for electronic structure calculations in relation to the X-ray absorption mechanisms in the separate (low to high) energy regime. XANES is a fingerprint of the electronic structure, more specifically of the valence state of the absorbing atoms.

To achieve a quantitative understanding of this development and appraisal of the corresponding XANES profiles, we have drawn some points on the description of the core and semi-core d electron excitation process through its interaction with the valence electrons of U$^{4+}$ in UO$_2$. The relevant model, which we use to solve the electronic structure problem, is based on the LFDFT algorithm. The LFDFT is a computational code merging the power of non-empirical DFT-based calculations with a phenomenological model. With this theoretical modelling, we have obtained parameters corresponding to the Slater-Condon integrals, the spin-orbit coupling interaction and the ligand field potential, which allowed computation of the multiplet energy levels arising from the ground 5f$^6$ and the excited nd$^5$5f$^5$ electron configurations of U$^{4+}$ in UO$_2$ with n = 3, 4 and 5. We have additionally calculated the oscillator strength due to the dipole allowed nd$^{10}$5f$^5$ $\rightarrow$ nd$^9$5f$^6$ transitions and the corresponding XANES profiles of UO$_2$ at the uranium M$_{4,5}$, N$_{4,5}$ and O$_{4,5}$ edges. Our calculated absorption bands are in good agreement with experimental spectra. Calculations for the uranium L$_{2,3}$ edges are underway.

Acknowledgements

This work was supported by Swissnuclear. We are indebted to the travel grant from the Scientific Computing & Modelling (www.scm.com) in Amsterdam. An anonymous reviewer is also thanked for providing useful comments and suggestions to improve this presentation.

Notes and references

1 M. Dolg, Computational Methods in Lanthanide and Actinide Chemistry, John Wiley & Sons, United Kingdom, 2015.
The molecular cluster (UO$_2$)$_{12}^{-}$ is surrounded by 122 point charges corresponding to 80 oxygen ions with $Q = -2$ and 42 uranium ions with $Q = +4$, which allow simulation of the Madelung potential representing the long range interaction due to the periodicity of the UO$_2$ crystal structure.

