DFT+U study of the structures and properties of the actinide dioxides

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ABSTRACT

The actinide oxides play a vital role in the nuclear fuel cycle. For systems where current experimental measurements are difficult, computational techniques provide a means of predicting their behaviour. However, to date no systematic methodology exists in the literature to calculate the properties of the series, due to the lack of experimental data and the computational complexity of the systems. Here, we present a systematic study where, within the DFT+U formulism, we have parametrized the most suitable Coulombic (U) and exchange (J) parameters for different functionals (LDA, PBE, PBE-Sol and AM05) to reproduce the experimental band-gap and lattice parameters for ThO2, UO2, PuO2, AmO2 and CmO2. After successfully identifying the most suitable parameters for these actinide dioxides, we have used our model to describe the electronic structures of the different systems and determine the band structures, optical band-gaps and the Bulk moduli. In general, PBE-Sol provides the most accurate reproduction of the experimental properties, where available.

We have employed diamagnetic order for ThO2, PuO2 and CmO2, transverse 3k antiferromagnetic order for UO2 and AmO2, and longitudinal 3k antiferromagnetic order for NpO2. The Fm 3 m cubic symmetry is preserved for diamagnetic ThO2, PuO2 and CmO2 and longitudinal 3k NpO2. For UO2 and AmO2, the transverse 3k antiferromagnetic state results in Pa3 symmetry, in agreement with recent experimental findings. Although the electronic structure of ThO2 cannot be reproduced by DFT or DFT+U, for UO2, PuO2, NpO2, AmO2 and CmO2, the experimental properties are very well represented when U = 3.35 eV, 6.35 eV, 5.00 eV, 7.00 eV and 6.00 eV, respectively, with J = 0.00 eV, 0.00 eV, 0.75 eV, 0.50 eV and 0.00 eV, respectively.

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1. Introduction

The study of the actinides is primarily due to their importance in the nuclear fuel cycle [1–3], where actinides are commonly encountered in their oxide form (AnO2) [4–6]. Compared to the metal, the oxides demonstrate better thermal and chemical stability [7], which is of paramount importance when we are considering nuclear energy applications [5]. The oxides enable higher operating temperatures and guard against further oxidation of the nuclear fuels, thus limiting the risk of containment failure and thermal excursion, and ensuring the stability of spent nuclear material for long-term storage [3,4,8]. It is, however, known, that under specific conditions, the AnO2 may form peroxides of the form AnO2+1x [9].

In nuclear fission, fissile isotopes of uranium (235U) and plutonium (239Pu) are used as nuclear fuel [3,10]. Over time the composition of nuclear oxide fuels change, either because of the genesis of irradiation-induced defects [11], or because of the formation of fission products [12]. Actinide oxides of interest include thorium dioxide (ThO2), neptunium dioxide (NpO2), americium dioxide (AmO2), and curium dioxide (CmO2). Evaluating the properties of these materials has important implications for the development of new processing strategies and the storage of nuclear material, thus improving the generation and safety of nuclear energy [10,12–16].

The study of AnO2 properties faces several challenges. The actinides are inherently unstable, highly toxic, and, due to the risk of nuclear proliferation, subject to strict regulatory controls [17]. In addition, their experimental investigation is further limited by their low abundance, isolation, and high radioactivity, which has led to
the lack of precise information on fundamental properties [10,18]. Important properties that require assessment include the lattice constants, band gaps, magnetic states, and the bulk moduli. Theoretical chemistry represents a powerful tool to unequivocally determine these properties, although conventional computational methods, based on density functional theory (DFT), often fail due to the relativistic influences and the highly correlated nature of these materials [19]. Hence, treatment by specific methods that are able to assess this type of material is required.

The following methods have been developed to calculate highly correlated materials: the self-interaction correction method (SIC) [20], modified density functional theory (DFT+U) [20,21], dynamic mean field theory (DMFT) [22], and screened hybrid density functional theory (HSE) [23,24]. Among these methodologies, DFT+U represents one of the more computationally tractable means of study, since it offers the possibility to accurately reproduce the properties of highly correlated materials at a reasonable computational cost [1]. This is achieved by manipulating the so-called Hubbard Coulombic (U) and exchange (J) modifiers, thus minimizing the DFT self-interaction error (SIE). However, U and J have to be fitted to experimental data, and for actinide materials, relativistic effects must also be taken into account. This is the reason why, to the best of our knowledge, few theoretical reports have been published so far that consider SOI and noncollinear magnetic order [25–27]. Hence, determining an accurate value of U and J would represent a significant advance, allowing in the exploration of these materials.

In this paper, we perform a systematic study to determine a set of Hubbard parameters for the DFT+U approximation for different functionals and different materials. Past studies have so far limited themselves to discussion of LDA, PW91, PBE or hybrid functionals. To the authors’ knowledge, this is the first study of the AnO2 systems that investigates the performance of the more current AM05 and PBE-Sol functionals. In previous work in the literature, these functionals have been shown to improve calculations of transition metal complexes in terms of the lattice constant and surface energetics [28,29]. It is thus important that their application to AnO2 systems is evaluated as well. In addition, we have chosen to implement the magnetic state determined from experimental procedure and/or predicted from the one electron crystal field (CF) and Russel-Saunders coupling scheme [25]. Our paper aims to address the issues encountered in the computational modelling of actinide-based materials to provide a basis on which further research can be built.

### 1.1. Magnetic structure

In Fig. 1, under standard conditions, the AnO2 crystallize in the calcium fluoride (CaF2) structure with Fm3 m (No. 225) symmetry [14,30–33]. The lattice resembles that of an actinide face-centred cubic (FCC) structure where oxygen occupies tetrahedral holes. Thus, each actinide (An⁴⁺) cation is coordinated to eight oxygen anions (O²⁻) and each O²⁻ is coordinated to four An⁴⁺.

The electronic structure of these materials can be understood from the crystal field (CF) theory. The spin–orbit interaction (SOI) forces the splitting of the single electron f-orbitals into two different levels, with j = 7/2 and j = 5/2 respectively. Since each actinide cation has a cubic environment, the degeneracy of these levels is subsequently broken due to the crystal field. Although this approximation is technically valid only for the single f-electron case, it is widely used to predict the orbital occupancy [25]. Hence, Th⁴⁺, Pu⁴⁺, and Am⁴⁺ are predicted to have no effective magnetic moment (µeff = 0), whereas U⁴⁺, Np⁴⁺, and Am⁴⁺ present an effective magnetic moment (µeff ≠ 0), as illustrated in Fig. 2.

The display of these magnetic moments in the actinide dioxides leads to different magnetic structures. If the ions show no magnetic moment, the resulting structure is the non-ordered diamagnetic (DM) state. In contrast, if the ions do present an effective magnetic moment, we can have two different situations, depending on whether these magnetic moments are coupled or not. If they are decoupled, there is no ordered distribution and this results in the disordered paramagnetic (PM) phase; if ordered or coupled, either ferromagnetic (FM) or antiferromagnetic (AFM) states are obtained. In addition, if magnetic moments are coupled but they show different µeff, the resulting magnetic structure is known as the ferrimagnetic (FI) state.

Further, the ordered magnetic AnO2 exhibit non-collinear behaviour [34–37], where the magnetic moments of the ions have contributions in more than one direction. Hence, we distinguish 1k (one wave-vector), 2k (two independent wave-vectors), and 3k (three independent wave-vectors) magnetic wave vectors [37,38]. Specifically, AnO2 are non-collinear 3k AFM materials. For the AFM 3k phase, one can distinguish between three different phases: the longitudinal 3k AFM, and the two equivalent transverse 3k AFM domains (A and B), depicted in Fig. 3.

The non-ordered magnetic systems are relatively simple to calculate compared to their ordered non-collinear magnetic counterparts. Experimentally, for ThO2 [39] and PuO2 [40–44] diamagnetism has been confirmed. It is worth noting, however, that there is a controversy over the exact magnetic ground state for PuO2, where first-principles methods have suggested the possibility of an AFM ground state [25]. However, for this work we have assumed the DM ground state, as supported by the complete active space self-consistent field (CASSCF) method and experimental evidence [40,42–44]. CMox has been reported as PM, although according to CF theory, it should be DM. It is believed that the presence of impurities plus the small energy difference between the DM and the PM phases in CMox are the reason the DM phase has been difficult to determine [12,46]; here, we have again assumed a DM ground state for CMox.

The ordered magnetic AnO2 have attracted considerable interest, in part due to their non-trivial multi-k and multiorbital nature. In UO2, the magnetic ground state is identified as a transverse 3k AFM [47] state at TN = 30.8 K [48] (where TN is the Neel Temperature for an AFM-PM magnetic transition) with a corresponding effective magnetic moment of 1.74 µB/µm. In addition, neutron studies have indicated an internal oxygen displacement; although no external distortion of the cubic structure occurs [49]. One theoretical study has employed SOI to investigated the relative
energetics of the transverse 1-3k AFM states, and has indicated a transverse 3k AFM as the ground state [19], which is the one considered in this study.

In NpO₂ the effective magnetic moment of the ground state remains conspicuously elusive [35, 50–52]. Np⁴⁺ is a Kramers ion [53], an ion with an odd number of electrons, and therefore should have a magnetic moment. ²³⁷Np Mössbauer spectroscopy [50] and muon experiments [35] have thus far established an effective magnetic moment of 0.00 ± 0.15 μB/Np⁴⁺ ion. In the PM phase, T_N = 60 K, the effective magnetic moment increases to 2.95 μB/Np⁴⁺ ion [54]. The longitudinal 3k AFM ground state has been inferred from resonant x-ray scattering [36] and ¹⁷O NMR [55] studies on NpO₂. Thus, the longitudinal 3k AFM state is employed in these calculations.

Finally, experimental information on the magnetic structure of AmO₂ is extremely limited. Magnetic susceptibility measurements have indicated a AFM ground state (T_N = 8.5 K) although the effective magnetic moment still eludes detection. Conversely, in the PM phase, an effective magnetic moment of 1.32 μB/Am⁴⁺ ion at 15–40 K and 1.53 μB/Am⁴⁺ ion at 50–100 K has been measured [56]. Independent Mössbauer and neutron measurements have been unable to provide evidence for a magnetically ordered ground state [57]. In ¹⁷O NMR studies on (Pu₀.⁹¹Am₀.⁰₉)O₂ an effective magnetic moment of 1.38 μB/Am⁴⁺ ion is estimated with an internal distortion of the oxygen ions synonymous with transverse 3k AFM order [58]. To our knowledge no study has examined the energetic differences between the longitudinal 3k AFM and transverse 3k AFM phases of AmO₂. Thus, the relative energetics of the longitudinal 3k AFM and the transverse AFM states are compared to establish the magnetic ground state for AmO₂.

2. Computational method

All calculations have been performed using the Vienna Ab-initio Simulation Package (VASP) [19–21], which is based on the density functional theory (DFT) and uses plane wave basis sets. In our convergence tests (Fig. S1) we defined an acceptable kinetic energy cut-off of 500 eV. Our results are derived from the conventional 12 atom unit cell. The description of the reciprocal space was done using the Monkhorst-Pack (5/5/5) k-point mesh, and the integration over the Brillouin zone was performed using the Blöchl tetrahedron method [60]. In optical absorbance calculations the k-point mesh is (15/15/15). For the band structure calculations, we define the k-point pathway for the respective Fm₃m and Pa₃m structures as G/L/W/X/G and G/M/R/X/G respectively. In this study, we have compared the following exchange-correlation functionals: the local density approximation (LDA) [61, 62], and with the general gradient approximation (GGA) of Perdew-Burke-Ernzerhof (PBE) [63], Armiento-Mattsson (AM05) [28, 64, 65], and PBE revised for solids (PBE-Sol) [66]. We have considered the following explicit electrons for each atomic species: O (1s²2s²2p⁴); Th (5f⁶6s²6p⁶6d²⁰); U (5f⁶6s²6p⁶6d⁸7s²); Np (5f⁶5d⁶6p⁶6d⁸7s²); Pu (5f⁶5d⁶6p⁶6d⁸7s²); Am (5f⁶5d⁶6p⁶6d⁸7s²); and Cm (5f⁶5d⁶6p⁶6d⁸7s²). The ion-electron interaction was described using the projector-augmented wave (PAW) approximation [22, 23], also considering in our calculations the relativistic spin-orbit interactions (SOI) and magnetic moments as well [67]. All calculations were performed until self-consistency, with electronic and ionic thresholds of 1·10⁻⁷ eV and 1·10⁻² eV Å⁻¹ respectively, and with the evaluation of the forces performed by the conjugate gradient algorithm [68]. The correction of the electron self-interaction error implicit in DFT calculations, was assessed by...
considering the on-site Coulombic (U) and exchange (J) constant, as described in the rotationally invariant Liechtenstein et al. formulism [69]. These parameters were applied to the f-orbitals of the actinide elements. The juxtaposition of energy levels and the presence of meta-stable states makes the determination of the lowest energy configuration for a given magnetic system challenging [70–73]. Here, by analysing trends in the data, and where necessary considering different initial electronic and ionic configurations, we have ensured as far as possible that the lowest energy configurations were obtained.

3. Results & discussion

3.1. Non-ordered magnetic materials

In our discussion, the performance of the functionals is addressed with DFT+U where a range of U = 0.00–7.00 eV is investigated, with J = 0.00 eV. Next, the effect of exchange is investigated further with PBE-Sol for U = 0.00–7.00 eV and J = 0.00–1.00 eV.

The non-ordered AnO2 materials include ThO2, PuO2 and CmO2. The selection of DFT+U values and the exchange correlation functional influences first on reproducing the electronic structure as described by the band gap, and second the lattice constant. In general, the functional has little effect on the calculated band gap between the respective L and X- valleys, exemplifying the Mott insulator characteristics of ThO2. In ThO2, the Th4+ f orbitals to which the U is applied are unoccupied and the DFT+U correction thus has a marginal effect on the electronic structure. Here, the experimental band gap of 5.90 eV [74,75] is systematically underestimated by DFT and DFT+U (Fig. S2a). The AM05/PBE-Sol functionals provided the closest approximation when U = 6.00 eV with a calculated value of 4.70–4.80 eV. This represent an underestimation of 1.10–1.20 eV but it still offers an improvement over previous DFT studies [76,77]. The lattice constant is reproduced by LDA, AM05 and PBE-Sol when U = 3.00–6.00 eV, but remains overestimated by PBE (Fig. S3a).

In PuO2, the band gap is greatly influenced by the choice of U. In DFT, PuO2 is calculated in the DM state as a metal; increasing U enables the formation of a band gap. In addition, past DFT studies have additionally reported a FM metallic ground state [78]. Our focus, however, remains on replicating the experimentally derived DM condition. Given the limited experimental information, previous DFT+U studies referenced the band gap from the activation energy for electronic conduction at 1.80 eV [16,70,79,80], but recent measurements of optical absorbance on epitaxial films have reported a band gap of 2.80 eV [81,82]. Assuming the correct band gap is 2.80 eV, this is reproduced by all functionals when U = 6.00–6.50 eV (Fig. S2b). In this range, the lattice constant is best represented by AM05/PBE-Sol (Fig. S3b).

No experimental data has been published on the band gap of CmO2, although it is believed to be an insulator [12,18,25]. Our calculations also point in that direction. As U is increased CmO2 transforms from a semi-conductor to an insulator, with a band gap that increases from 0.50 to 2.50 eV for U = 0.00–7.00 eV. Although none of the functionals calculates the experimental lattice constant exactly, an excellent approximation is made by AM05 and PBE-Sol when U = 6.00 eV (Fig. S3c).

In general, the experimental data is best represented by the AM05 and PBE-Sol functionals. Thus, PBE-Sol is used further to investigate the influence of the exchange modifier (J) on the band gap and the lattice constant (Figs. S4–S5). In ThO2, the introduction of J is negligible with respect to the band gap and only has a minor effect on the lattice constant. In PuO2, the J value detrimentally affects the band gap, with a reduction of 0.50 eV for every 0.25 eV of J. In fact, the best band-gap fitting is obtained when J = 0.00 eV, with negligible implications to the lattice parameter. Finally, in CmO2, the J modifier increases the band gap by a maximum of 0.75 eV when J = 0.00–1.00 eV, and again, it has barely any influence on the lattice parameter.

In conclusion, for the non-ordered magnetic AnO2, we have observed that the PBE-Sol/AM05 functionals, in a combination of high U values with J = 0.00 eV, provide the most accurate reproduction of both band-gaps and lattice constants. Hence, assuming the PBE-Sol functional, we have selected the following U and J parameters for an accurate description of the non-ordered magnetic AnO2: ThO2 (U = 6.00 eV, J = 0.00 eV), PuO2 (U = 6.35 eV, J = 0.00 eV) and CmO2 (U = 6.00 eV, J = 0.00 eV). It is worth noting that the U and J choice for CmO2 is based upon the better fitting to its lattice parameter, thus predicting a bandgap of 2.50 eV. Using these determined values, we have calculated the following variables: band structure, density of states, optical absorbance spectra, and bulk modulus.

The band structures of ThO2, PuO2 and CmO2 are presented in Fig. 4. In ThO2, the calculated indirect band gap of 4.63 eV occurs between the respective L and X- points of the valence band maximum (VBM) and conduction band minimum (CBM), compared to the experimental band gap of 5.90 eV [74,75]. In PuO2, with respect to the band structure, we report a X- centred direct band gap of 2.81 eV. For CmO2, an indirect band gap of 2.50 eV is predicted between the respective L and X points. It is worth noting that the valence band maximum is set to 0 eV. For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
PuO₂ and CmO₂, the conduction band is primarily formed from the An(f) states, from which we predict these materials to be charge-transfer insulators where electronic transitions occur across the oxygen and actinide ions.

Whereas the fundamental band gap represents the transition between the VBM and CBM the optical band gap is restricted by orbital symmetry. In recent studies on In₂O₃ [83], AgCuS [84], PbO₂ [85], TiO₂ [86], and SrCu₂O₂ [87], it has been shown that the fundamental band gap and optical band gap may differ in value. Our calculated optical absorbance spectra for ThO₂, PuO₂ and CmO₂ are shown in Fig. 6, from which we found that the direct band gap and the optical band gap respectively differ by 0.11 eV, 0.01 eV and 0.19 eV for ThO₂, PuO₂ and CmO₂, as shown in Table 1. Thus, optical absorbance measurements are only representative of the direct band gap in PuO₂.

Finally, the bulk modulus is notoriously difficult to obtain experimentally, which has led to a large range of values in the literature [32,95]. However, our calculated values are in excellent agreement with the experimental data for ThO₂ and CmO₂. For PuO₂, where the reported experimental values for the bulk modulus range from 178 to 379 GPa, our value of 217 GPa provides a useful point of comparison for experimentalists.

3.2. Ordered magnetic materials

Experimental studies of UO₂ and NpO₂ indicate a respective transverse 3k AFM and longitudinal 3k AFM ground state, whereas for AmO₂, to our knowledge, the domain of the 3k AFM ground state is undetermined. Initially, the relative energetics of the transverse 3k AFM and longitudinal 3k AFM states were calculated with PBE-Sol when U = 0.00–7.00 eV and J = 0.00 eV. DFT calculates a longitudinal 3k AFM ground state, but DFT+U (U = 1.00–7.00 eV) calculates a transverse 3k AFM ground state (Table S1). Thus, the transverse 3k AFM ground state is used for further calculations of AmO₂.

As with the non-ordered magnetic AnO₂, the choice of functional has a minor effect on the calculated band gap (Fig. S6) and on the effective An⁴⁺ magnetic moment (Fig. S7). The sole exception is the band gap of AmO₂: here, when U = 7.00 eV, the LDA functional differs considerably from the GGA functionals. The choice of functional, however, has a pronounced effect on the lattice constants (Fig. S8). The U modifier is the dominating factor for the band gap and the lattice constant, whereas the magnetic moment is only marginally influenced. In order to be consistent with the analysis of the non-ordered magnetic materials, the influence of J was only assessed for the PBE-Sol functional.

In UO₂, the experimental band gap of 2.00–2.50 eV [98–101] is reproduced when U = 3.00–4.00 eV, for all functionals (Fig. S6a). In this range, the experimental lattice constant of 5.470–5.473 Å [32,88,102] is perfectly matched by the AM05/PBE-Sol functionals. The experimental effective magnetic moment of 1.74–1.80 μ₈/U⁴⁺ ion [48,103] is underestimated by all functionals. In DFT (U = 0 eV) the calculated magnetic moment is found between 0.05 and 1.01 μ₈/U⁴⁺ ion. In DFT+U, when U = 1.00 eV, it is improved to 1.04–1.41 μ₈/μ₈ ion. Unfortunately, further increases in U have minimal impact. The introduction of J, only considered for the PBE-Sol, has minimal impact on the band gap (Fig. S5a) and the lattice constant (Fig. S11a). However, J has a detrimental influence on the effective

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**Fig. 5.** The calculated density of states for a) ThO₂, b) PuO₂, c) CmO₂. The valence band maximum is set to 0 eV.

**Fig. 6.** Calculated optical absorption for a) ThO₂, b) PuO₂, c) CmO₂. The black and red lines respectively represent α (the optical absorption coefficient) and (αhv)² (the optical absorption). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Table 1
Direct band gap (ΔE₀, eV) and optical band gap (ΔEₒ, eV), lattice constant (aₜ in Å) and bulk modulus (B₀ in GPa) for ThO₂, PuO₂, and CmO₂, as calculated in this paper (Calc.) compared to the reported experimental values (Exp.). U and J are expressed in eV and all materials have been calculated as diamagnetic.

<table>
<thead>
<tr>
<th>AnO₂</th>
<th>DFT+U</th>
<th>ΔE</th>
<th>aₜ</th>
<th>B₀</th>
</tr>
</thead>
<tbody>
<tr>
<td>ThO₂</td>
<td>6.00</td>
<td>0.00</td>
<td>4.63</td>
<td>4.74</td>
</tr>
<tr>
<td>PuO₂</td>
<td>6.00</td>
<td>0.00</td>
<td>2.81</td>
<td>2.82</td>
</tr>
<tr>
<td>CmO₂</td>
<td>6.00</td>
<td>0.00</td>
<td>2.50</td>
<td>2.69</td>
</tr>
</tbody>
</table>

Table 2
Magnetic moment (μ, in μₜ), direct band gap (ΔE₀, eV) and optical band gap (ΔEₒ, eV), lattice constant (aₜ in Å), bulk modulus (B₀ in GPa) for UO₂, NpO₂, and AmO₂, as calculated in this paper (Calc.) compared to the reported experimental values (Exp.). U and J are expressed in eV. The transverse 3k AFM state is calculated for UO₂ and AmO₂, whereas, the longitudinal 3k AFM state is calculated for NpO₂.

<table>
<thead>
<tr>
<th>AnO₂</th>
<th>DFT+U</th>
<th>μ</th>
<th>ΔE</th>
<th>aₜ</th>
<th>B₀</th>
</tr>
</thead>
<tbody>
<tr>
<td>UO₂</td>
<td>3.35</td>
<td>0.00</td>
<td>1.35</td>
<td>1.74–1.80 [48,103]</td>
<td>3.11 [39]</td>
</tr>
<tr>
<td>NpO₂</td>
<td>5.00</td>
<td>0.75</td>
<td>1.87</td>
<td>0.00–0.15 [35,50]</td>
<td>3.08 [82]</td>
</tr>
<tr>
<td>AmO₂</td>
<td>7.00</td>
<td>0.50</td>
<td>3.15</td>
<td>1.38 [58]</td>
<td>1.31</td>
</tr>
</tbody>
</table>

U⁴⁺ magnetic moment (Fig. S10a). Thus, a better correlation between the band gap and the effective U⁴⁺ magnetic moment is obtained in the absence of J. Hence, for PBE-Sol, U = 3.35 eV and J = 0.00 eV are the appropriate values to describe UO₂, Table 2. Our value of U and J for the transverse 3k AFM ground state is considerably lower than the calculated U = 5.7 eV and J = 0.4 eV values obtained explicitly from the constrain random phase approximation (crPA) method considering collinear 1k AFM order [104]. It is generally accepted that the AnO₂ crystallise in the Fm3m motif. Therefore, in UO₂, internal distortion of the oxygen ions had been reported by a number of studies [37,49,103]. Only recently, rather than oxygen distortion, an alternative Pa₃m symmetry was suggested from neutron diffraction measurements [105]. In our calculations, the transverse 3k AFM ground state is successful in reproducing this subtle Pa₃m symmetry as the lowest-energy structure, in contrast to previous theoretical studies where collinear 1k AFM order was modelled which results in either Fm3m or Pa₃m symmetry or a distorted Fm3m symmetry. Here, the distortion of 0.016 Å is identical to that derived from neutron diffraction measurements [105].

For PuO₂, the experimental band gap of 2.85–3.10 eV [82,106] is well reproduced by all functionals when U = 5 eV (Fig. S6b), and the experimental lattice constant of 5.434 Å [88,93] is reproduced by PBE (U = 0.00 eV), LDA (U = 3.00–4.00 eV), and AM05/PBE-Sol (U = 6.00–7.00 eV). The effective Np⁴⁺ magnetic moment, however, remains enigmatic. Our calculated effective Np⁴⁺ magnetic moment ranges from 2.14 to 2.77μₜ/Np⁴⁺ ion (Fig. S7b) depending on the functional rather than on the U parameter. However, these values are not compatible with the experimental values since, as indicated by several experimental papers, the magnetic moment is only 0.00–0.15 μₜ/Np⁴⁺ ion [35,50,51,107–109]. Interestingly, our value does compare with the effective paramagnetic moment of 2.95 μₜ/Np⁴⁺ ion, determined at 60 K [54].

In AmO₂, the experimental band gap of 1.30 eV [106] is well reproduced by all the functionals when U = 4.00 eV. Additionally, the band gap is also calculated by the GGA functionals when U = 7.00 eV (Fig. S6c). However, the experimental lattice constant of 5.372–5.379 Å [14,30,93,112] is reproduced only by the AM05/PBE-Sol when U = 4.00–7.00 eV, (Fig. S8c). The experimental effective magnetic moment of 1.38 μₜ/Np⁴⁺ ion [58] is overestimated by ~3.00 μₜ/Np⁴⁺ ion regardless of the U value, as depicted in Fig. S7c. The magnetic moment of AmO₂ is small when contrasted against the spin moment. Consequently, we have examined the orbitals, Table S2. Here, the f orbital is the major contributor to the total magnetic moment of the Am⁴⁺ ion. It is, however, unclear what causes the reduction in the orbital moment. In the specific case of the PBE-Sol functional, the introduction of J (Figs. S9c–11c) provides a better correlation between the band gap, lattice constant...
and the estimated effective $\text{Am}^{4+}$ magnetic moment when $U = 7.00$ eV and $J = 0.50$ eV. Although the calculated effective $\text{Am}^{4+}$ magnetic moment still presents a significant overestimation, it offers an improvement over previous calculations\[13,113\]. In these DFT$+$U constraints, the lattice constant is in excellent agreement with the experimental information (see Table 2). Finally, it is noted that the transverse $3k$ AFM structure produces an internal distortion of the oxygen ions with amplitude 0.082 Å, which is substantially greater than observed in UO$_2$ (Table 2). The resultant structure therefore possesses $\text{Pm}$$\bar{3}$ symmetry. Unfortunately, the lattice distortion in AmO$_2$ has not yet been experimentally investigated.

In summary, for the ordered magnetic AnO$_2$ the AM05/PBE-Sol functionals provide the closest representation of the experimental band gap, effective magnetic moment of the An$^{4+}$ ions and the lattice constant. Thus, from the PBE-Sol functional, the band structure, density of states, optical absorption and bulk modulus are calculated. In contrast to their non-ordered magnetic counterparts, the degeneracy of the bands is perturbed for ordered magnetic AnO$_2$ systems. In UO$_2$, an indirect band gap of 2.06 eV occurs between $\Gamma$ and $R$ of the VBM and CBM. For NpO$_2$, a direct band gap of 3.08 eV occurs at the L point. For AmO$_2$, an indirect band gap of 1.31 eV occurs between respective $\Gamma$ and $\Gamma$-$R$ of the VBM and CBM respectively.

The DoS for UO$_2$, NpO$_2$ and AmO$_2$ is illustrated in Fig. 8. Unlike NpO$_2$, AmO$_2$ and the non-magnetic actinide oxides, in which the valence band is dominated by the O(p) states, UO$_2$ presents a valence band dominated by U(f) states, clearly indicating that UO$_2$ is a Mott \[100,114\] insulator. In each compound the An(f) states contribute heavily to the conduction band. The An(d) states feature more prominently when the energy is greater than 4.00 eV.

The calculated optical absorption spectra for UO$_2$, NpO$_2$ and AmO$_2$ are shown in Fig. 9. The direct band gap and optical band gaps respectively differ by 0.14 eV, 0.03 eV and 0.14 eV for UO$_2$, NpO$_2$ and AmO$_2$, as shown in Table 2. Thus, optical absorbance measurements can only be fully relied upon for the fundamental electronic structure of NpO$_2$.

Finally, the calculated bulk modulus is in excellent agreement with the experimental data. For AmO$_2$, the calculated value of 196 GPa offers a point of comparison against the large experimental range of 205–280 GPa.

4. Conclusions

In this paper we have presented a systematic and comprehensive computational study of the actinide oxides: ThO$_2$, UO$_2$, NpO$_2$, PuO$_2$, AmO$_2$, and CmO$_2$. We have compared the performance of the LDA, PBE, AM05 and PBE-Sol functionals in combination with the DFT$+$U methodology. The choice of functional has little effect on the calculated band gap, but it has a major influence on the lattice constant, bulk modulus and the magnetic moment. In the majority of cases AM05 and PBE-Sol behave in an identical fashion providing the best estimate of the lattice constant. We conclude that either
AM05 or PBE-Sol would provide the best performance in future research of these oxide materials.

The selection of U and J primarily focused on the replication of the magnetic ground state of all oxides to ensure the correct modelling of the magnetic properties as well as the band gap and lattice constants. Where no experimental data was available for AmO2 we have proposed the longitudinal 3k AFM as the magnetic ground state. For the magnetic ordered oxides, UO2, NpO2, AmO2, U and J were also parametrized to reproduce the effective An4 magnetic moment of the materials. In terms of the lattice, displacement of the oxygen ions results in a Pa-3m structure which is only observed in the transverse 3k AFM structure employed in the calculation of UO2 and AmO2 but no such distortion is recorded from the longitudinal 3k AFM for NpO2.

With the DFT–U parameters determined for each compound, we have also studied other properties like the band-structure, density of states, and the optical band gap. Only for PuO2 and NpO2 is excellent agreement found between the fundamental and experimental data. However, at the moment, this work offers as good a set of parameters and the electronic properties. As such, this paper offers a systematic study of the electronic properties using screened hybrid density functional theory.

References


