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First-principles DFT+U investigation of charged states of defects and fission gas atoms in CeO2

Lei Shi,1,* Emerson Vathonne,1 Vincent Oison,2 Michel Freyss,1 and Roland Hayn2
1CEA, DEN, DEC, Centre de Cadarache, 13108 Saint-Paul-lec-Durance, France
2Aix-Marseille Universit´e, IM2NP, Campus Scientifique Saint-Jérôme, Case 142, 13397 Marseille Cedex 20, France

I. INTRODUCTION

The investigation of defects and impurities in oxide materials is a subject of high scientific interest, both for fundamental and applied sciences. First-principles studies are of great importance in that research field. For instance, taking native point defects in ZnO as a representative example, the influence of the proper charge state on the defect formation energies was demonstrated by Kohan et al. [1] In our following study, we concentrate on cerium dioxide (CeO2), an oxide of the rare-earth metal cerium, that has recently received great scientific interest due to its technical importance in a large number of applications. Generally, CeO2 is widely used as automobile exhaust catalyst, electrolyte of solid oxide fuel cell (SOFC), oxygen storage device, etc. [2–6]. For experimental radiation damage studies in nuclear applications, CeO2 could largely simulate UO2, having the same crystal structure, similar bulk properties and common features in the electronic structure with the advantage of being nonradioactive. That simplifies considerably any experimental study on radiation damage. Therefore a comparative study of the calculated defect and fission gas properties in UO2 and CeO2 is of particular interest as a first step to confirm the similarities of both oxides regarding radiation damage.

CeO2 is an insulator with a 3 eV band gap [7,8]. The oxidation states of the atomic elements are, respectively, Ce4+ and O2−. In CeO2, the 4f states remain completely unoccupied. However, the hybridization between Ce 4f orbitals and O 2p orbitals in the valence band has been demonstrated in the study of Wuilloud et al. [8]. In the case of nonstoichiometry, i.e., for oxygen deficient samples where Ce4+ changes to the trivalent Ce3+, the chemical reduction of CeO2 leads to the localization of one 4f electron on each Ce3+ ion. Experimentally, in Ohno and Iwase’s study [9,10], Ce3+ ions are observed in the vicinity of oxygen vacancies created under irradiation with swift Xe ions by using extended x-ray absorption fine structure (EXAFS) measurements, and a new peak of normalized intensity in x-ray photoelectron spectroscopy (XPS) spectrum is observed and corresponds to the Ce3+ state in the valence band. Moreover, a ferromagnetic behavior is determined as an intrinsic property of reduced ceria due to the magnetic moments induced by the localized 4f electrons of the Ce3+ ions [11]. This ferromagnetic behavior opens the way to possible spintronics applications of CeO2. The ratio of Ce3+/Ce4+ is shown to be an important factor in the appearance of this magnetic property during irradiation experiments with neon ions [12,13].

To investigate CeO2 including defects and impurities by a first-principles approach, the DFT+U method is used in this study in order to describe properly the localized 4f electrons in contrast to the standard DFT approach, which is clearly insufficient. Previous first-principles calculations have been performed to describe the ground-state properties of bulk cerium oxides (CeO2 and Ce2O3) [14–20] and of CeO2 with defects [21–37].

In order to get insight into the defect behavior in CeO2 and to investigate an eventual similarity to UO2, we report here the results of a detailed first-principles study on defects and fission gases in CeO2 and show that the proper charge state has an important influence, not only on the defect formation energy but also on the incorporation of fission gas atoms (Xe and Kr) in CeO2. The studies of Zacherle et al. [24] and Keating et al. [33] on point defects in CeO2 already treated the influence of the charge states for the defects, but we will show that their results have to be completed by all possible intrinsic charge states. Xiao et al. [38] considered the trapping and diffusion of Xe in CeO2 by taking into account only the neutral defects. In our study, we investigate the trapping behavior of Xe and another fission gas atom Kr in defect sites under various charge states. Furthermore, we improve the DFT+U calculations for charged defects in CeO2 by a thorough use of the occupation matrix control (OMC) scheme. This scheme has to be used for a partially filled f shell as it was for instance demonstrated in the case of UO2 [39,40] to avoid the metastable states, which were reported in several publications based on the
DFT+$U$ approach [41–43]. In detail, we study the neutral and charged single-point defects by calculating their formation energies, and the fission gases Xe and Kr by calculating their incorporation energies in CeO$_2$. We also investigate larger defects such as bound Schottky defects. Three regimes of stoichiometry are considered in the calculations: hypostoichiometric (oxygen-poor), stoichiometric, and hyperstoichiometric (oxygen-rich) conditions. By taking into account the necessary technical improvements in the theoretical description, we are able to obtain results that provide a new insight into the trapping properties of fission gases in CeO$_2$ and similar oxides.

II. COMPUTATION DETAILS

The DFT calculations were all performed using the Vienna $ab$ initio simulation package (VASP) code [44] in terms of the projector augmented wave method [45,46]. Both generalized gradient (GGA-PBE) [47] and local density approximations (LDA-CA) [48] have been considered to study the defect behavior. The valence electrons (5$s^2$5$p^6$4$f^1$5$d^1$6$s^2$ for Ce and 2$s^2$2$p^4$ for O) were treated using a plane wave basis set truncated by a cutoff energy of 500 eV. The core electrons are frozen and taken into account by using PAW atomic data. The same atomic data are used for the various oxidation states of each element. An additional on-site Coulomb repulsion was considered by including a Hubbard-like term proposed by Dudarev et al. in order to take into account the $f$-electron localization [49]. Based on the study of Castleton et al. [28], showing the $U$-parameter dependence of the Ce 4$f$ electron localization in the presence of the O vacancy, the optimum $U$ value for Ce 4$f$ states in our study was selected to be 6.0 eV for LDA+$U$ and 5.0 eV for GGA+$U$ calculations. We did not take into account the Hubbard term for O 2$p$ states, which was considered in Keating et al.’s study in order to have a better description of localized holes on the O 2$p$ orbitals in presence of the Ce vacancy in CeO$_2$ [32,33]. However, in their study, the influence of the $U$ parameter for O 2$p$ states was found negligible for the formation energies of point defects except the Ce vacancy [32]. It is to be noted that this influence for the Ce vacancy under its formal charge state (4+) is also negligible, because there is no localization of holes on the O 2$p$ orbitals, since the holes are electron compensated completely. In our study, we will show later that we find the charge state 4+ to be the most favorable one for Ce vacancy in CeO$_2$, justifying the fact that we can neglect the Hubbard term for O 2$p$ states.

The defect calculations were performed using a supercell containing 96 atoms, i.e., by doubling the conventional cell along the three lattice parameters, which is a standard supercell also used in the previous defect studies using DFT+$U$ for CeO$_2$ [24,32,33,38] but also for UO$_2$ [50–52], with which we want to compare our results. Let us note that we verified the energy convergence with respect to the supercell size, by comparing the defect formation energies of neutral Ce and O vacancies obtained from calculations using supercells containing 96 and 324 atoms. We found a maximum difference of 0.14 eV on the defect formation energies between the two supercells. Thus we conclude that the calculations using a 96-atom supercell are a good compromise between accuracy and CPU time. In all calculations, the atomic positions and lattice parameters were optimized until forces acting in each atom were less than 0.005 eV/Å and stress components were less than 0.01 kbar.

We first conduct bulk property calculations on the perfect CeO$_2$ crystal in order to have a comparative investigation on the difference of calculated results between the LDA+$U$ and GGA+$U$ functional. Table I presents the experimental and calculated values of some bulk properties of CeO$_2$: lattice constant, bulk modulus, band gap, elastic constants, enthalpy of formation and cohesive energy. The bulk properties calculated using both LDA+$U$ and GGA+$U$ are in good agreement with the experimental values. However, we see that LDA+$U$ reproduces slightly better electronic properties such as the band gap as well as some other bulk properties of CeO$_2$ (lattice constant, bulk modulus, cohesive energy and enthalpy of formation) than GGA+$U$. Therefore, for all the following calculations considering the defect formation energies and fission gas incorporation energies, we will mainly use the LDA+$U$ functional.

In the study of defects, in order to avoid the metastable states yielded by the DFT+$U$ method, the occupation matrix control (OMC) scheme was used following the scheme implemented by Jomard et al. [41] and Amadon et al. [42]. The OMC scheme distinguishes the ground state from metastable states based on the total energy of the system, as detailed in the study of UO$_2$ by Dorado et al. [39]. For CeO$_2$, the electronic ground state of the supercell containing a neutral oxygen vacancy involves the localization of electrons, initially localized on the removed oxygen atom, on two of four adjacent cerium atoms, leading to the formation of two Ce$^{3+}$ ions. A DFT+$U$ calculation of CeO$_2$ with oxygen defects without an adequate preconditioning method yields the distribution of charges on several adjacent Ce ions and fails to reach the system ground state with localized 4$f$ electrons on Ce$^{3+}$ ions. To solve this problem by using the OMC scheme, the proper 4$f$ electron occupation matrix that corresponds to the Ce$^{3+}$ ions on which the 4$f$ electrons are localized should be determined in the first place.

In order to find this occupation matrix corresponding to the Ce$^{3+}$ ions, we used an approach proposed by Zacharle et al. [24] in the calculation of an electron polaron. The
additional electron is well localized on the chosen Ce ion if we displace all the neighboring atoms (both Ce and O atoms) as a preconditioning before the structure relaxation. With this approach, we succeed to obtain the occupation matrix corresponding to the Ce$^{3+}$ ion with one 4f electron localized. By applying this matrix in the calculation of the CeO$_2$ supercell containing one neutral oxygen vacancy, we find that the 4f electrons are well localized on two of the four first neighboring Ce ions in the vicinity of the oxygen vacancy. And the corresponding system energy is about 13 meV/(CeO$_2$) lower than the one when the electrons are distributed on the adjacent Ce ions without forming Ce$^{3+}$. In addition, by using the OMC scheme, we could even control the positions of the Ce$^{3+}$ ions on which the electrons are localized. Our study shows that the localization of 4f electrons on either two of the four first neighboring Ce ions results in the ground state of CeO$_2$ with one neutral oxygen vacancy.

III. DEFECT FORMATION AND FISSION
GAS INCORPORATION

A. Formation energies of charged defects

We first investigate the formation of charged single defects in CeO$_2$. The defect formation energy for a defect X in the charge state $q$, $E_{\text{form}}(X^q)$, is obtained using the standard expression:

$$E_{\text{form}}(X^q) = E_{\text{tot}}(X^q) - \left( E_{\text{tot}}(\text{bulk}) - \sum_i n_i \mu_i \right)$$

$$+ q(E_{\text{Fermi}} + E_{\text{VBM}}) + E_{\text{corr}},$$

where $E_{\text{tot}}(X^q)$ is the total energy of the supercell containing the charged defect, and $E_{\text{tot}}(\text{bulk})$ is the total energy of the perfect bulk supercell. $n_i$ is the number of atoms of type i removed from ($n_i > 0$) or added to ($n_i < 0$) the bulk cell to form the vacancy and interstitial defects, respectively, and $\mu_i$ is the corresponding chemical potential. $E_{\text{Fermi}}$ is the Fermi energy in the range of the band gap, which corresponds to the electron chemical potential. The zero energy of $E_{\text{Fermi}}$ is fixed at the valence-band maximum ($E_{\text{VBM}}$) of the perfect supercell. An energy correction term, $E_{\text{corr}}$, is proposed by Taylor and Bruneval for the calculation of charged defect formation energies in a finite-size supercell as follows [57]:

$$E_{\text{corr}} = \Delta E_{\text{el}} + q\Delta V.$$  

Where $\Delta E_{\text{el}}$ is the Madelung term [58,59], which corrects the spurious long-ranged electrostatic interaction between the charged defect and its periodic images. $\Delta V$ is a potential alignment term, which corrects the shifted band structure of defective supercells relative to the perfect supercell. For the moment, the computational resources available do not allow to take into account a sufficiently large supercell in order to neglect these two corrections. $\Delta E_{\text{el}}$ and $\Delta V$ are expressed as

$$\Delta E_{\text{el}} = \frac{\alpha q^2}{2\epsilon_0 L}.$$  

and

$$\Delta V = \langle \psi_{KS}^{\text{bulk}} \rangle - \langle \psi_{KS}^{\text{defect}} \rangle,$$

where $\epsilon_0$ is the static dielectric constant chosen as the experimental value 23 for CeO$_2$ [60]. $L$ is the length of the supercell edge. $\alpha$ is the Madelung lattice constant, which is 2.519 for a fluorite structure [61]. $\psi_{KS}$ is the average Kohn-Sham potential in the perfect supercell and $\langle \psi_{KS}^{\text{defect}} \rangle$ is the average Kohn-Sham potential in the supercell with defects. These potentials are the results of the DFT+$U$ calculations.

In Eq. (1), the chemical potentials $\mu_i$ correspond to the cerium and oxygen chemical potentials $\mu_{\text{Ce}}^{\text{CeO}_2}$ and $\mu_{\text{O}}^{\text{CeO}_2}$ in CeO$_2$. They cannot be obtained directly from the DFT+$U$ calculations because they depend, in particular, on the stoichiometry and the partial pressure of oxygen [$P$(O$_2$)]. To calculate the values of $\mu_{\text{Ce}}^{\text{CeO}_2}$ and $\mu_{\text{O}}^{\text{CeO}_2}$, accessible under different stoichiometry conditions, we use a method based on the study of Na-Phattalung et al. [62] and Hong et al. [63] considering the standard enthalpy of formation for various phases of the Ce-O system. For CeO$_2$, the standard enthalpy of formation is defined as

$$\Delta H_f^0(\text{CeO}_2) = \mu_{\text{Ce}}^{\text{CeO}_2} - \mu_{\text{Ce}}^{\text{Ce}^{3+}} - 2\mu_{\text{O}}^{\text{O}_2}.$$  

We introduce two variables $\Delta \mu_{\text{Ce}}$ and $\Delta \mu_{\text{O}}$ of the chemical species with respect to their standard states, defined as

$$\Delta \mu_{\text{Ce}} = \mu_{\text{Ce}}^{\text{CeO}_2} - \mu_{\text{Ce}}^{\text{Ce}^{3+}},$$

$$\Delta \mu_{\text{O}} = \mu_{\text{O}}^{\text{CeO}_2} - \mu_{\text{O}}^{\text{O}_2},$$

Let us note that the values $\Delta \mu_{\text{Ce}}$ and $\Delta \mu_{\text{O}}$ should be negative otherwise the CeO$_2$ crystal would decompose. The enthalpy of formation of CeO$_2$ can then be expressed as

$$\Delta H_f^0(\text{CeO}_2) = \Delta \mu_{\text{Ce}} + 2\Delta \mu_{\text{O}}.$$  

This gives an accessible range of the values of both chemical potential variables $\Delta \mu_{\text{Ce}}$ and $\Delta \mu_{\text{O}}$:

$$\Delta H_f^0(\text{CeO}_2) < \Delta \mu_{\text{Ce}} < 0$$

and

$$\frac{1}{2} \Delta H_f^0(\text{CeO}_2) < \Delta \mu_{\text{O}} < 0.$$  

In our LDA+$U$ calculations, the value of $\Delta H_f^0(\text{CeO}_2)$ is $-11.64$ eV, as reported in Table I. Thus the accessible ranges of $\Delta \mu_{\text{Ce}}$ and $\Delta \mu_{\text{O}}$ are determined as

$$-11.64 \text{ eV} < \Delta \mu_{\text{Ce}} < 0 \text{ eV}$$

and

$$-5.82 \text{ eV} < \Delta \mu_{\text{O}} < 0 \text{ eV}.$$  

In order to further narrow the ranges of $\Delta \mu_{\text{Ce}}$ and $\Delta \mu_{\text{O}}$, we consider the other oxide phase of the Ce-O system, Ce$_2$O$_3$, and put a constraint on $\Delta \mu_{\text{Ce}}$ and $\Delta \mu_{\text{O}}$ to take into account the relative stability of CeO$_2$ compared to Ce$_2$O$_3$. In analogy
to Eq. (9), the standard enthalpy of formation of Ce$_2$O$_3$ is expressed as follows:

$$\Delta H_f^0(\text{Ce}_2\text{O}_3) = 2\Delta \mu_{\text{Ce}} + 3\Delta \mu_{\text{O}}. $$

(10)

The calculated value of $\Delta H_f^0(\text{Ce}_2\text{O}_3)$ is $-20.57$ eV using LDA+U, which is close to the experimental value of $-18.60$ eV [55]. In Fig. 1, we determine the relative stability of each phase by plotting $\Delta \mu_{\text{Ce}}$ as a function of $\Delta \mu_{\text{O}}$ for CeO$_2$ and Ce$_2$O$_3$, respectively. And we can see that CeO$_2$ is more stable than Ce$_2$O$_3$ when $\Delta \mu_{\text{Ce}}$ is lower than $-6.23$ eV and $\Delta \mu_{\text{O}}$ is larger than $-2.70$ eV. This stability of CeO$_2$ compared with Ce$_2$O$_3$ determines a range accessible for $\Delta \mu_{\text{Ce}}$ and $\Delta \mu_{\text{O}}$ as follows:

$$-11.64 \text{ eV} < \Delta \mu_{\text{Ce}} < -6.23 \text{ eV}$$

and

$$-2.70 \text{ eV} < \Delta \mu_{\text{O}} < 0 \text{ eV}.$$  

The calculated values of $\mu_{\text{Ce}}^{\text{O-men}}$ and $\mu_{\text{O}}^{\text{O-men}}$ are $-4.55$ and $-5.24$ eV, respectively, by our LDA+U calculations. Note that in our study, we did not apply any correction for the DFT calculated binding energy of the O$_2$ molecule [64], which come into play in the term $\mu_{\text{O}}^{\text{O-men}}$. The well-known misestimation of the O$_2$ binding energy can be corrected, as suggested by Lee et al. [65], Wang et al. [66] and Grindy et al. [67], by fitting a correction to the calculated formation enthalpy of several metal oxides. However, Grindy et al. showed that the energy correction is small using the LDA functional (0.25 eV/O$_2$) [67]. And it was further emphasized by Lee et al. that applying the DFT+U method reduces even more the error from pure DFT in the calculation of formation enthalpies of oxides [65].

After applying our values of $\mu_{\text{Ce}}^{\text{O-men}}$ and $\mu_{\text{O}}^{\text{O-men}}$ into Eqs. (7) and (8), we could finally determine the accessible ranges for the chemical potentials $\mu_{\text{Ce}}^{\text{O}}$ and $\mu_{\text{O}}^{\text{O}}$ in CeO$_2$:

$$-16.19 \text{ eV} < \mu_{\text{Ce}}^{\text{O}} < -10.78 \text{ eV}$$

and

$$-5.24 \text{ eV} > \mu_{\text{O}}^{\text{O}} > -7.94 \text{ eV}.$$  

In this study, we have considered three limiting cases, depending on the stoichiometry: (1) the oxygen-rich conditions, where $\mu_{\text{Ce}}^{\text{O}}$ is maximum and $\mu_{\text{O}}^{\text{O}}$ is minimum; (2) the oxygen-poor conditions, where $\mu_{\text{Ce}}^{\text{O}}$ is minimum and $\mu_{\text{O}}^{\text{O}}$ is maximum; and (3) the stoichiometric conditions, where both $\mu_{\text{Ce}}^{\text{O}}$ and $\mu_{\text{O}}^{\text{O}}$ take their mean values.

The corresponding values of $\mu_{\text{Ce}}^{\text{O}}$ and $\mu_{\text{O}}^{\text{O}}$ calculated by both LDA+U and GGA+U functionals under these three stoichiometry conditions are reported in Table II. The GGA+U results obtained with the calculated enthalpies of formation, $\Delta H_f^0(\text{Ce}_2\text{O}_3) = -10.52$ eV and $\Delta H_f^0(\text{Ce}_2\text{O}_3) = -18.8$ eV using GGA+U, are very identical to those obtained by Zacherle et al. [24].

### B. Incorporation of fission products in charged defects

In order to investigate the trapping properties of gaseous fission products (FP) in CeO$_2$, we determine their incorporation energies in charged defect sites. The incorporation energies are given using the following general expression:

$$E_{\text{inc}}(FP,X) = E_{\text{tot}}(FP,X) - E_{\text{tot}}(X) - E_{\text{tot}}(FP), $$

(11)

where $E_{\text{tot}}(FP,X)$ is the total energy of the supercell containing a fission product incorporated in the defect site $X$; $E_{\text{tot}}(X)$ is the total energy of the supercell containing the defect site $X$; and $E_{\text{tot}}(FP)$ is the total energy of the isolated fission product.

In order to consider the various charge states of the defect site in CeO$_2$, the incorporation energy is expressed as a function of the formation energy of the defect site and the solution energy of the fission product in the defect site. This gives

$$E_{\text{inc}}(FP,X) = E_{\text{sol}}(FP,X^q) - E_{\text{form}}(X^q), $$

(12)

where $E_{\text{sol}}(FP,X^q)$ is the solution energy of the fission product in defect site $X$ with the most stable charge state $q'$:

$$E_{\text{sol}}(FP,X^q) = E_{\text{tot}}(FP,X^q) - \left( E_{\text{tot}}(\text{bulk}) - \sum_{j} n_j \mu_j \right) + q'(E_{\text{Fermi}} + E_{\text{VBM}}) + \Delta E_{\text{el}}' + q' \Delta V'.$$

(13)

Herein $E_{\text{sol}}(FP,X^q)$ is the total energy of the supercell containing the fission product incorporated in the defect site $X$ under charge state $q'$. $\mu_j$ is the chemical potential of the species $j$ (fission product atoms, Ce and O atoms) added or removed with respect to the perfect supercell of CeO$_2$. $\Delta E_{\text{el}}'$ is...
the Madelung term and $\Delta V'$ is the potential alignment term as defined in Eqs. (3) and (4), respectively. The term $E_{\text{form}}(X^q)$ is the formation energy of the defect site $X$ in its charge state $q$ obtained from Eq. (1). Thus the incorporation energy of the fission product in the charged defect site is expressed as a function of the Fermi energy as

$$E_{\text{inc}}(FP, X) = E_{\text{tot}}(FP, X^q) - E_{\text{tot}}(X^q) - n_{FP} \mu_{FP} + (q' - q)(E_{\text{Fermi}} + E_{\text{VBM}}) + (\Delta E_{\text{F}} - \Delta E_{\text{C}}) + (q' \Delta V' - q \Delta V).$$

In this equation, $\mu_{FP}$ is the chemical potential of the fission product in the standard isolated state. Let us note that the chemical potentials $\mu_{\text{CeO}_2}$ and $\mu_{\text{Ce}^3+}$ do not appear explicitly. Therefore $E_{\text{inc}}(FP, X)$ does not depend on the stoichiometry conditions.

We introduce a term $\Delta q = q' - q$. When $\Delta q = 0$, which means that the most stable charge state of the defect site $X$ does not change after trapping the fission product, the incorporation energy becomes independent of the Fermi energy $E_{\text{Fermi}}$ and can be expressed as

$$E_{\text{inc}}(FP, X) = E_{\text{tot}}(FP, X^q) - E_{\text{tot}}(X^q) - n_{FP} \mu_{FP} + q \Delta V' - q \Delta V.$$  

In this study, two gaseous fission products are considered, Xe and Kr. The corresponding results are presented in Sec. V.

IV. RESULTS ON THE DEFECT FORMATION

A. Density of states of CeO$_2$ containing the neutral oxygen vacancy

In order to have an insight into the influence of defects on the electron structure of CeO$_2$, we firstly compare the calculated total density of states of perfect CeO$_2$ with the one including a neutral O vacancy by LDA+$U$ in Fig. 2. The valence band mainly consists of O 2$p$ states, whereas the conduction band mainly consists of Ce 4$f$ states. In bulk CeO$_2$, the band gap is found to be 2.7 eV, in good agreement with the experimental value of 3 eV.

When a neutral O vacancy is created, the two extra electrons of the removed O atom are localized on two of the four adjacent Ce atoms. This forms two Ce$^{3+}$ ions, which present a ferromagnetic order, and affects the density of states. An additional electronic state appears in the band gap 1.3 eV above the maximum of the valence band. Consequently, the Fermi level is shifted by 1.4 eV towards the conduction band, indicating that O neutral vacancies involve n doping of CeO$_2$.

However, we should note that when considering a charged O vacancy as well as a charged Ce vacancy under their formal charge states +2 and −4, respectively, the electronic doping is canceled due to the compensation of charges. Thus the resulting density of states remains nearly the same as the one of bulk CeO$_2$.

B. Formation energies of charged point defects in CeO$_2$

We present here the results on the formation energies of defects under various charge states, considering three types of stoichiometry conditions. The formation energies are calculated as a function of the Fermi energy in the range of the band gap according to Eq. (1).

1. Small point defects

We first study the small point defects: the monovacancies (V$_O$ and V$_{Ce}$), the monointerstitials (I$_O$ and I$_{Ce}$) and divacancy (V$_O$+V$_{Ce}$). Figure 3 presents the comparison between the LDA+$U$ and GGA+$U$ results under the stoichiometric conditions. For each defect, the most stable charge state depends on the Fermi energy value in the band gap and is given above the curves. Overall good agreement is found between these two functionals in the calculated formation energies of each defect. The oxygen vacancy has the lowest formation energy and is the most stable defect. We can see that the charge states shown on the curves of each defect are quite similar between LDA+$U$ and GGA+$U$, and the most stable charge states of each defect for the Fermi level in the middle of the band gap are consistent to the ionic picture: V$_O^{2+}$, V$_{Ce}^{4+}$, I$_O^{2+}$, and I$_{Ce}^{2+}$. They are the formal charge states of the corresponding defects.

In the vicinity of a neutral oxygen vacancy, by using the occupation matrix control (OMC) scheme, we managed to localize the 4$f$ electrons on the Ce ions to form Ce$^{3+}$ ions in both LDA+$U$ and GGA+$U$ functional calculations. And in the case of a neutral cerium vacancy, the extra positive charge of the removed Ce atom is distributed on the neighboring O atoms and creates holes on the 2$p$ orbitals. As shown by Keating et al., the hole localization is expected when the $U$ parameter for O 2$p$ states is considered $[32,33]$. However, there is no such hole localization in the presence of a Ce vacancy under its formal charge state (−4), because the holes are fully compensated by electrons. And as seen in both Figs. 3 and 4, (4−) is the most stable charge state of Ce vacancies for the Fermi level in a wide range of the band gap from 0.6 to 3.0 eV by both LDA+$U$ and GGA+$U$ calculations. The hole localization on O 2$p$ states in the presence of Ce vacancies under nonformal charge states (different from 4−) would thus occur for Fermi energies close to the valence-band maximum, where the formation energies
FIG. 3. Formation energy of charged point defects (O vacancy $V_O$, O interstitial $I_O$, Ce vacancy $V_{Ce}$, Ce interstitial $I_{Ce}$, and divacancy) calculated using GGA+$U$ and LDA+$U$ as a function of the Fermi energy for CeO$_2$ under stoichiometric conditions.

(a) GGA+$U$

(b) LDA+$U$

FIG. 4. Formation energy of charged point defects (O vacancy $V_O$, O interstitial $I_O$, Ce vacancy $V_{Ce}$, Ce interstitial $I_{Ce}$, and divacancy) calculated using LDA+$U$ as a function of the Fermi energy for CeO$_2$ under O-rich and O-poor conditions, respectively.

For the O interstitial, it is to be noted that we only considered the $I_O$ in the octahedral site rather than the peroxide O interstitial (O dumbbell) in CeO$_2$. That is because our study focuses on the similarities of defect properties between CeO$_2$ and UO$_2$, and the O dumbbell in UO$_2$ is less stable than a single O interstitial according to the study of Middleburgh et al. [68].

2. Bound Schottky defects

The bound Schottky defects (BSD) are considered to investigate the formation of larger point defects. The BSD consists of one CeO$_2$ vacancy in the supercell (one Ce and two O adjacent vacancies). There are basically three BSD configurations, which depend on the positions of oxygen vacancies in the Ce-centered oxygen cage as shown in Fig. 5.

The formation energies of the bound Schottky defects are presented in Fig. 6 under stoichiometric conditions since they are stoichiometric defects. They do not depend on the values of the O and Ce chemical potentials. We can see that the formation energies of all three bound Schottky defects are in the range between 5.0 to 5.5 eV for the Fermi level near the middle of the band gap. The BSDs with oxygen vacancies along the (110) and (111) directions (BSD2 and BSD3) give the lowest energy. However, it is worth noting that the BSDs form in both the (110) and (111) directions with lower formation energy than the BSD1 with oxygen vacancy along the (100) direction.
formation energies for a wide range of Fermi energy near the middle of the band gap. These are the two configurations of BSDs with the two oxygen vacancies that are farther away from each other. The ionic character of CeO$_2$ is also confirmed. The neutral charge state is favored for the most stable BSDs (BSD2 and BSD3) for the Fermi level up to around 2 eV, which is again consistent with the ionic character of CeO$_2$. Between 1.5 and 2.0 eV also the nonformal charge state ($-1$) of BSD1 becomes as stable as the neutral states of BSD2 and BSD3. And the formation energies of all the three BSDs are becoming similar with each other with a charge state $-2$ for a Fermi level position in the vicinity of the conduction-band minimum.

3. Comparison with UO$_2$

The results obtained for the defect formation using LDA+$U$ for CeO$_2$ are compared here with the ones obtained in UO$_2$ using GGA+$U$ in Vathonne et al.’s study [50]. This comparison between the results obtained using the two different LDA+$U$ and GGA+$U$ approximations is made possible by the overall good agreement between the LDA+$U$ and GGA+$U$ results on point defect formation in CeO$_2$ (shown in Fig. 3). Viewing the defect formation energy graphs for CeO$_2$ and UO$_2$ (Ref. [50, Fig. 3]) under stoichiometric and nonstoichiometric conditions, we observe great similarities in the formation energy of each defect between those two compounds. The relative stability of the defects as a function of the Fermi energy follows the same trends for CeO$_2$ and UO$_2$. Moreover, the most stable charge states of each defect in CeO$_2$ are all in good agreement with the ones in UO$_2$.

Table III presents the calculated formation energies of defects in both CeO$_2$ and UO$_2$ under three different stoichiometry conditions and for the Fermi energy in the middle of the band gap. We can see that the most stable charge state of each defect for the Fermi level in the middle of the band gap is consistent to the ionic character of both compounds (except the uranium interstitial in UO$_2$), which means that there are no additional electrons localized or holes formed on the cation $f$ orbitals or oxygen $p$ orbitals, respectively. The formation energy values of oxygen vacancies and cation interstitials ($V_O$ and ICe) in CeO$_2$ are quite similar to the ones in UO$_2$ with differences between 0.2 and 0.5 eV. However, when considering other types of defects ($I_O$, V$_{Ce|U}$, and divacancy), the differences rise to a range between 1.6 and 2.8 eV. That can be explained by the fact that it is more easy to stabilize O-rich compounds in UO$_2$ than in CeO$_2$. For instance, there exist stable compounds $U_3O_8$ and $U_4O_9$ besides UO$_2$. However, this is not the case for Ce-O compounds, where only CeO$_2$ and Ce$_2$O$_3$ are stable oxides [69]. In other words, $U$ can have the valency up to $U^{6+}$ in UO$_3$, which is not the case for Ce. This is the fundamental difference between CeO$_2$ and UO$_2$, which leads

![FIG. 6. Formation energy of charged bound Schottky defects (BSD) calculated using LDA+$U$ as a function of the Fermi energy for CeO$_2$.](image)

TABLE III. Formation energies (eV) and the most stable charge states of various defects for the Fermi energy in the middle the band gap in CeO$_2$ using LDA+$U$ under various stoichiometry conditions and the comparison with UO$_2$ using GGA+$U$.

<table>
<thead>
<tr>
<th>Defects</th>
<th>O-rich</th>
<th>Stoichiometric</th>
<th>O-poor</th>
<th>Charge state</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CeO$_2$</td>
<td>UO$_2$</td>
<td>CeO$_2$</td>
<td>UO$_2$</td>
</tr>
<tr>
<td>$V_O$</td>
<td>2.90</td>
<td>2.58</td>
<td>1.56</td>
<td>1.29</td>
</tr>
<tr>
<td>$V_{Ce</td>
<td>U}$</td>
<td>2.23</td>
<td>0.00</td>
<td>4.94</td>
</tr>
<tr>
<td>$V_{Ce</td>
<td>U+V_O}$</td>
<td>3.40</td>
<td>0.68</td>
<td>4.75</td>
</tr>
<tr>
<td>$I_O$</td>
<td>1.60</td>
<td>0.06</td>
<td>2.95</td>
<td>1.35</td>
</tr>
<tr>
<td>ICe$^{\prime\prime}$</td>
<td>10.39</td>
<td>10.70</td>
<td>7.68</td>
<td>8.12</td>
</tr>
<tr>
<td>BSD1</td>
<td>-</td>
<td>-</td>
<td>5.49</td>
<td>3.25</td>
</tr>
<tr>
<td>BSD2</td>
<td>-</td>
<td>-</td>
<td>4.91</td>
<td>2.47</td>
</tr>
<tr>
<td>BSD3</td>
<td>-</td>
<td>-</td>
<td>5.00</td>
<td>2.60</td>
</tr>
</tbody>
</table>

*Reference [50].
to higher formation energies of oxygen interstitials and cation vacancies (but also divacancy) in CeO$_2$ than in UO$_2$.

For bound Schottky defects, the BSD2 and BSD3 are the most stable configurations (with the BSD2 having a slightly lower formation energy than the BSD3) in both UO$_2$ and CeO$_2$ for the Fermi energy near the middle of the band gap. The formation energies of all the BSDs in CeO$_2$ are about 2.2–2.4 eV higher than the ones in UO$_2$, which can be linked to the higher formation energies of cation vacancies in CeO$_2$ compared to UO$_2$.

Now we compare the point defects under neutral and nonformal charge states between CeO$_2$ and UO$_2$. In the presence of V$_O$ and ICe$_{U}$, the extra available electrons are localized on the $f$ orbitals of adjacent cations to form Ce$^{3+}$ and U$^{3+}$. In the presence of other neutral defects such as V$_{CeU}$, IO, and divacancy, there are no Ce$^{5+}$ ions in CeO$_2$ unlike in UO$_2$ in which U$^{5+}$ ions can be formed. The formation of Ce$^{5+}$ would require the removal of one electron from the fully occupied bands of Ce$^{4+}$ ions, which is very unfavorable. Indeed, the fifth ionization energy of Ce is high (it is estimated as 65.55 eV in the study of Reader and Epstein [70]) compared to fifth ionization energy of U (45.77 eV based on Pyper et al.’s study [71]) and also in comparison to the electron affinity of Ce$^{4+}$ to form Ce$^{3+}$, which is 36.76 eV [72].

We should note that our study of defects in CeO$_2$ and the comparison to UO$_2$ do not take into account the effect of the temperature. Temperature may have an influence on the defect formation energies due, in particular, to the temperature dependence of the oxygen molecule chemical potential, as studied by Mastrikov et al. [73]. However, our objective is to focus on the physics of point defects in CeO$_2$ and the comparison to UO$_2$, regardless of the conditions of use of these materials.

V. INVESTIGATION OF XE AND KR TRAPPING PROPERTIES IN CeO$_2$

In order to have an insight into the trapping behavior of the fission gases in CeO$_2$, the incorporation energies of Xe and Kr atoms were calculated as a function of the Fermi energy according to Eq. (14) in various defect sites: single vacancies, divacancies, and the bound Schottky defects. There is no dependence on the stoichiometry conditions because the chemical potentials $\mu_{CeO_2}$ and $\mu_{O}$ are canceled out in the calculation of the incorporation energy. $\Delta q = q' - q$, the change of the charge state of the defect site before $(q)$ and after $(q')$ the incorporation of the fission gas atom, is indicated on the corresponding curves of each defect site in Figs. 7 and 8.

A. Incorporation energies of Xe and Kr atoms

Figure 7 shows that the highest incorporation energy for the Xe atom is found when considering incorporation in the oxygen vacancy. For the Ce vacancy, the incorporation energy strongly depends on the Fermi energy in the region above the valence-band maximum from about $-3.7$ eV ($p$-doped ceria) to $+3.1$ eV in the middle of the band gap. For the divacancy $(V_{Ce}+V_{O})$, the incorporation energy of Xe for the Fermi energy in the middle of the band gap is $0.96$ eV, which is a lower energy than the one in the mono vacancy $(V_{Ce}$ or $V_{O})$. When considering the incorporation in the bound Schottky defects, we found that the BSD1 configuration with oxygen vacancies along the $(100)$ direction gives the lowest incorporation energy which is $0.15$ eV for the Fermi energy in the middle of the band gap. Regarding geometry of the three configurations of BSD in Fig. 5, the BSD1 is more compact than the BSD2 which is more compact than the BSD3. This explains that the BSD1 is the most preferable incorporation site for Xe atoms.

For the incorporation of Kr in CeO$_2$ (shown in Fig. 8), the preferred defect sites are found consistent with Xe. However, for Kr, the value of the Fermi energy has no effect as significant as for Xe on the incorporation energies. Table IV shows Xe and Kr incorporation energies in various defect sites for the Fermi energy in the middle of the band gap. The calculated incorporation energies of Kr are generally lower than the ones of Xe, which means that Kr is more easily incorporated in the defect sites than Xe in CeO$_2$. The BSD1 is still the favorable trapping site for Kr with an incorporation energy slightly lower than $0$ eV for the Fermi energy in the middle of the band gap. Considering the most stable charge states of the defect sites, we can see that for both Xe and Kr, $\Delta q$ is $0$ (except for the
incorporation of Xe in $V_{Ce}$ for the Fermi energy in the middle of band gap, which means that the most stable charge state remains unchanged after the incorporation of Xe and Kr in the defect sites. The incorporation energies for both Xe and Kr in the defect sites depend on the Fermi energy when $\Delta q$ is nonzero. The most significant variation of incorporation energies occurs for Xe incorporation into Ce vacancies and Fermi energies close to the valence-band maximum ($p$-doped case).

B. Density of states of CeO$_2$ with Xe and Kr incorporated in the BSD1

In order to have an insight into the modification of the electronic structure of CeO$_2$ induced by fission gas atoms, we plotted in Fig. 9 the density of states of CeO$_2$ with Xe and Kr incorporated in their most favorable trapping site, that is the neutral bound Schottky defect (BSD1). The density of states for CeO$_2$ with the neutral BSD1 only and for perfect CeO$_2$ are presented as well for comparison.

We see that the densities of states of CeO$_2$ with Xe and with Kr in a BSD1 Schottky defect are very similar to each other and to the one of CeO$_2$ with the BSD1 Schottky defect only. However, in comparison to the density of states of perfect CeO$_2$, we find that the band gaps slightly decrease: when Xe is trapped in BSD1, the conduction band is shifted by 0.5 eV towards the valence band. And when Kr is trapped in BSD1, the conduction band is shifted by 0.4 eV towards the valence band.

C. Comparison with UO$_2$

Our results of the Kr and Xe incorporation in CeO$_2$ are compared with the ones obtained in UO$_2$ by Andersson et al. [74], Vathonne et al. [51], and Dorado et al. [52]. As already mentioned, the difference between LDA+$U$ and GGA+$U$ is not essential for that comparison. The same OMC scheme and the supercell-size corrections to formation and incorporation

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**TABLE IV.** Incorporation energies of Xe and Kr atoms in various defect sites for the Fermi energy in the middle of the band gap in CeO$_2$ using LDA+$U$ and comparison with the corresponding values for UO$_2$ taken from the literature (GGA+$U$ in Refs. [51,52], LDA+$U$ in Ref. [74]) where the incorporation energies were calculated from their data using Eq. (12).

<table>
<thead>
<tr>
<th>Defect sites</th>
<th>Xe atom</th>
<th>Kr atom</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CeO$_2$</td>
<td>UO$_2$</td>
</tr>
<tr>
<td>$V_O$ (eV)</td>
<td>7.32</td>
<td>6.89$^{b,c}$</td>
</tr>
<tr>
<td>$V_{Ge}+V_O$ (eV)</td>
<td>2.68</td>
<td>2.29$^{a,3,84}$$^{b,c}$</td>
</tr>
<tr>
<td>BSD1 (eV)</td>
<td>0.15</td>
<td>0.18$^{a,1.18}$$^{b,c}$</td>
</tr>
<tr>
<td>BSD2 (eV)</td>
<td>1.18</td>
<td>1.82$^{b,c}$</td>
</tr>
<tr>
<td>BSD3 (eV)</td>
<td>2.10</td>
<td>2.25$^{b,c}$</td>
</tr>
</tbody>
</table>

$^a$Reference [74].

$^b$Reference [51].

$^c$Reference [52].

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energies were used in the studies of UO$_2$ and CeO$_2$. Table IV shows the comparison of Xe and Kr incorporation energies between CeO$_2$ and UO$_2$. Common features in the trapping properties of Xe and Kr between UO$_2$ and CeO$_2$ are found: the calculated incorporation energies of Xe and Kr in the same defect sites are in good agreement between UO$_2$ and CeO$_2$ for the Fermi level in the middle of the band gap; the BSD1 has the lowest incorporation energy of both Xe and Kr for UO$_2$ and CeO$_2$, which means that the bound Schottky defect within this configuration is the most energetically preferable site to trap either the Xe or the Kr atoms in both compounds. Furthermore, there is a correlation between the fission gas incorporation and the vacancy size: the larger the vacancy size is, the more favorable the fission gas atoms are trapped in.

Comparing the incorporation energy graphs between CeO$_2$ and UO$_2$ (Ref. [51, Fig. 4.1]), we notice that the incorporation energy of Kr in the defect sites of UO$_2$ remains almost constant ($\Delta q = 0$) as a function of the Fermi energy. However, for CeO$_2$, we can see that the incorporation energy of Kr (but also of Xe) in V$_{\text{Ce}}$ changes with the Fermi energy in the region between the valence-band maximum and the middle of the band gap [shown in Fig. 8(a)]. We should note that the charge states of cation vacancies are not formal for the Fermi energy in this region. The differences between UO$_2$ and CeO$_2$ arise since there may be U$^{5+}$ ions created in UO$_2$, yet no formation of Ce$^{5+}$ ions in CeO$_2$ is possible. Instead of the formation of +5 valence state for Ce, the incorporation of the Kr atom in V$_{\text{Ce}}$ results in the charge loss on the 4$p$ orbitals of the Kr atom and the 2$p$ orbitals of the eight neighboring O atoms in CeO$_2$. Bader charge analysis indeed shows that after the incorporation of one Kr atom in the cation vacancy under nonformal charge states, the Kr atom loses some charge in CeO$_2$ but not in UO$_2$.

For instance, for nonformal charge states of V$_{\text{Ce}}$, the Kr atom loses charge up to 0.8 $|e|$ from Bader charge analysis. This charge loss behavior has also been found in our study of the incorporation of Xe into the Ce vacancy, which agrees well with Xiao et al.’s study showing the same Bader charge loss of 1.4 $|e|$ for a Xe atom in a neutral V$_{\text{Ce}}$ [38].

This charge loss behavior mainly changes the difference between UO$_2$ and CeO$_2$ for the incorporation of Xe and Kr in defect sites. However, it does not appear for a Fermi energy close to the middle of the band gap where the most stable trap sites are under their formal charge state. In addition, considering the significant common features found in this section between these two compounds, we can conclude that the incorporation properties of the fission gases Xe and Kr are similar in CeO$_2$ and in UO$_2$.

VI. CONCLUSIONS

In this study, we investigated the formation of various charged defects and the incorporation of the fission gas atoms Xe and Kr in CeO$_2$. The electron potential (Fermi energy) is an important factor that influences the formation and incorporation energies of small and large charged defects. By calculating the formation energy of defects under various charge states in CeO$_2$, great similarity was found between CeO$_2$ and UO$_2$. For the incorporation of fission gas atoms, CeO$_2$ and UO$_2$ have similar trapping properties of Xe and Kr atoms. The bound Schottky defect BSD1 is the most energetically preferable site to host fission gas atoms in both compounds. So, we can conclude that CeO$_2$ could largely simulate UO$_2$ under stoichiometric and nonstoichiometric conditions. That is true for nearly all positions of the Fermi energy within the band gap besides the heavily p-doped case (Fermi level close to the valence-band maximum). Let us note that a certain difference between UO$_2$ and CeO$_2$ exists for the incorporation of Xe and Kr atoms in the cation sites under nonformal charge states: Xe and Kr atoms lose charge in CeO$_2$ but not in UO$_2$, leading to differences in the incorporation energies. The reason is the absence of the +5 valence state for Ce in contrast to its existence for U. However, this difference between CeO$_2$ and UO$_2$ is not found for the defect sites under their formal charge state, which is their favorable configuration for the Fermi level near the middle of the band gap. Therefore it does not influence the good agreement of the incorporation properties of Xe and Kr between CeO$_2$ and UO$_2$.

Based on these common features on point defects and fission gas atoms that we have obtained by DFT+$U$ calculations for CeO$_2$ and UO$_2$, we could conclude that our study contributes to justify the following assumption: CeO$_2$ might be considered as a surrogate of UO$_2$ to study experimentally the radiation damage. However, our study is only one of the first steps to ascertain the similarities of UO$_2$ and CeO$_2$ in terms of radiation damage. More investigations could be conducted to compare the atomic transport properties of defects and fission gases, grain boundary behavior and fission gas release in UO$_2$ and CeO$_2$ using the relevant modeling techniques.

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