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Polaronic interactions between oxygen vacancies in rutile TiO₂

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Oxygen vacancy-vacancy interactions in rutile TiO_2 are studied in conjunction with polaron formation trends using density functional theory calculations. It is found that polarons strongly enhance the formation of oxygen vacancies in this material, and also mediate the interactions between existing vacancies. At distances below one nanometer, two isolated and charge-neutral vacancies exhibit attractive interactions with an equilibrium distance of about 4 Ångstroms. The attractive forces between vacancies partly arise from the polaronic transfer of excess electrons to reduce the potential energy. These discoveries provide microscopic explanations to the vacancy clustering phenomena, as well as a practical approach to stabilize the polarons at arbitrary Ti atoms in TiO₂.

I. INTRODUCTION

Recently, titanium-oxide nanomaterials have been widely investigated for a range of applications including photo-catalysis [1-5], solar energy conversion [6-7] and nano-electronic devices such as high-k gate dielectrics [8] and resistive random-access memories (RRAM) [9-12]. The conductivity of titanium oxide varies significantly in these applications, which is affected by factors such as non-stoichiometry, crystallinity and surface morphology. Especially, an increased level of bulk n-type conductivity is often observed without the intentional introduction of dopants during the fabrication process [13-15]. Therefore, the n-type semiconducting behaviors have traditionally been attributed to the presence of oxygen vacancies (V_0) which contribute excess electrons to the lattice [13-17]. However, the nature of these excess electrons still remains a matter of active debate due to the widely-observed electron self-trapping phenomena, i.e. the polaronic effects in titanium oxide. In pristine TiO₂, the small-polaron model has been proposed to describe the excess electrons localized around Ti atoms [18], supported by recent electron spin resonance (ESR) measurements [19]. Nevertheless, in the presence of oxygen vacancies (V_0), the picture becomes more complex and recent studies suggested the coexistence of both small polarons and delocalized electrons [17].

As the oxygen-vacancy concentration increases, titanium oxide eventually becomes metallic due to the clustering of these point defects. Long-range order of vacancies leads to crystallographic shear and transformation into the Magnéli phases (Ti_nO_{2n-1} , n>3 [20]). These substoichiometric phases have been widely observed in TiO₂ thin films [11], nanoparticles [21] and nanowires [22]. Due to the metallic conductivity, they have been widely considered as possible electrode materials in batteries and fuel cells [23-25]. The desirable chemical stability and abundant low-temperature behaviors of Magnéli phases also generated substantial research interests lately [26-30]. For example, various bipolaron models have been proposed to explain the two-step metal-insulator transitions of Ti_4O_7 between 125K and 150K [28-30].

In previous studies, the isolated and clustered oxygen vacancies in titanium oxide have been investigated separately and it is still unclear how a pair of oxygen vacancies will interact when they come close to each other. Although it has been suggested that clustered vacancies have lower energies compared to isolated vacancies in the neutral charge state [31-32], the effects of multiple polaronic states were not thoroughly considered. Moreover, during the approaching of two oxygen vacancies, the redistribution of charges coupled with the structural change has never been fully investigated. In this work, we report an in-depth study of the vacancy-vacancy interactions in rutile TiO_2 based on density functional theory (DFT) and an extensive search of potential polaronic states. The total energy of the system in each polaronic state has been calculated to identify the ground state at various vacancy-vacancy distances and to characterize the potential energy surface. Our results suggest that two isolated V_o in rutile TiO_2 exhibit attractive interactions, with an equilibrium distance of about 4 Å. The attractive forces between vacancies arise from the polaronic effects which redistribute the excess electrons to minimize their Coulomb repulsion. These insights provide an in-depth understanding of the vacancy clustering process from a microscopic point of view.

II. METHODS

Theoretical studies of defect states with density functional theory (DFT) are challenging since the standard local/semi-local approximations such as local density approximation (LDA) and generalized gradient approximation (GGA) severely underestimate the bandgap of TiO₂ and suppress the polaron formation [31]. In order to reproduce the correct polaronic behaviors, the self-interaction of electrons must be treated properly to balance the tendency of localization versus delocalization [33-44]. Hybrid functionals [33-40] and on-site Coulomb corrections [31, 32, 41-44] are two potential treatments. Using hybrid functionals, several groups of researchers have reported the co-existence of two distinct polaronic states [17, 35-38]. The first state is the widely-observed closed-shell solution, which has two electrons distributed in between three Ti neighbors [35-37]. The Kohn-Sham (KS) energy level of this defect state is relatively deep (0.4~1.0 eV below conduction band minimum [17, 33, 36, 40-45]), in contrast to the experimentally-observed shallow donor states [46-49]. Thus, they are unlikely to be the major contributor of n-type conductivity in rutile TiO₂. On the other hand, the second state reported has two excess electrons localized at two neighboring Ti sites, in the form of electron polarons [17, 36, 39]. This state, only discovered recently by hybrid functional calculations, was predicted to be more stable over the first one [17, 36].

Hybrid functional calculations usually require one order of magnitude more computational resources than standard DFT [50], limiting its application to systems with no more than a few hundred atoms. In this work, an alternative approach has been adopted to study the polaronic states in rutile TiO_2 , which is DFT with on-site Coulomb corrections (DFT+U). Our previous calculations confirmed its capability to reproduce the correct band-gap and lattice constants of TiO_2 [31]. Here the exact cancellation of self-interaction errors is further verified by checking the 'generalized Koopmans condition' [33-36]. The finely-tuned corrections were then applied to calculate the polaronic states in pristine and oxygen-deficient TiO_2 . Qualitative agreements were observed with the previously-reported hybrid-functional calculations, but obtained at much-reduced computational cost.

The predicted ground states of an isolated V_0 are found to be consistent with the recently-reported experimental results by electron spin resonance (ESR) [46-48]. By considering the polaronic effects, the defect levels of an isolated vacancy become significantly shallower (<= 0.1 eV), which provides a more consistent explanation for the n-type conductivity in TiO₂. With consistent description of an isolated V₀, the methodology was then applied to study the vacancy-vacancy interactions. By varying the distance between a pair of V₀, the change of thermodynamic stability and electronic structures in

various polaronic states are thoroughly discussed.

The DFT calculations were carried out using density functional theory as implemented in the Vienna ab initio simulation package (VASP [51]), in which the projector-augmented-wave pseudopotentials were employed [52]. For K-point sampling, 2x2x2 Monkhorst-Pack grid in the first Brillouin zone were applied. All calculations were based on a 3x3x4 supercell of rutile TiO₂ with a total of 216 atoms. The atomic relaxations were performed until the force on each atom is reduced to <0.005 eV/Å with a plane-wave cut-off energy of 353 eV. The exchange and correlation energy of electrons is described with LDA+U [53]. In order to overcome the underestimation of bandgap by LDA, on-site Coulomb corrections to both O 2p orbitals and Ti 3d orbitals were applied (LDA+ U^d+U^p). Here we use 8 eV for U^d , 6 eV for U^p and 0.6 eV for the exchange parameter J, which adequately reproduced the electronic structures and lattice constants of rutile TiO₂ [31]. These parameters are consistent with the values obtained by proper rescaling of U values of single ions (calculated by DFT) to fit the electronic bandgap of TiO₂ [43]. The importance of applying U to O-2p in addition to Ti-3d states was discussed elsewhere [31, 43, 54].

Compared to the widely-studied hybrid-functional method [17,35-38], DFT+U has significantly better computational efficiency, making it favorable for the studies of large systems and multiple polaronic states. On the other hand, DFT+U may be less accurate for the treatment of el-el correlation effects, and the choice of U values requires extra caution. In this work, the choice of U values followed our previous work which examined various U^d - U^p combinations and their effects on the geometry and band structures of rutile TiO₂ [31]. Here we further verify the capability of our model to properly describe the electron self-interaction, by checking the 'generalized Koopmans condition' (gKC) [33-36]. The gKC states that, in order to fully eliminate the self-interaction errors associated with the exchange-correlation functional, the K-S level of the defect state should be equal to the electron addition/removal energy of the system. When the gKC is satisfied, the linear behavior of total energy as a function of fractional occupation number is retained. In this work, we follow the procedure described by Deak et al. [35] to verify gKC. As shown in Table I, the fulfillment of gKC is confirmed to be within 0.1eV for both oxygen vacancy and Al dopant [55].

In constrast to the empirical determination of U, there are more rigorous ways to compute U fully *ab initio*, such as the constrained random phase approximation (cRPA) [56] or the linear response approach by Cococcioni et al. [57]. However, these methods often lead to lower U^d values and under-estimated bandgap. In order to achieve the correct bandgap for TiO₂, relatively high U^d value for Ti is needed, which enhances the tendency of electron localization and polaron formation at Ti sites. In general, it would be interesting to examine the polaronic behaviors of TiO₂ at various U values, which is beyond the scope of this study.

III. POLARONIC STATES OF AN ISOLATED OXYGEN VACANCY

Fig. 1(a) demonstrates the electron localization functions (ELF) indicating a small polaron localized at two different Ti sites in rutile TiO_2 . The ELF is a powerful tool to identify polaron formation, which has a value between 0 and 1 that characterizes the type of chemical bonding in the space [58]. The cross-shaped charge localization is a signature of electron polaron at the Ti atomic sites. As a

benchmark, we further simulated the migration of a small polaron by linear interpolation of the atomic positions [17,59-61]. The non-adiabatic polaron transfer energy in rutile [001] direction is calculated to be around 0.30 eV (Fig. 1(b)), consistent with previous GGA+U calculation [59].

With consistent results on a small polaron, we moved on to study the polaronic states of a single V_0 . With the creation of one V_0 , two excess electrons are contributed to the system. Fig. 2(a) depicts the atomic configuration of the single V_0 . Here we define the two nearest-neighboring Ti atoms in the diagonal direction as Ti(1) and Ti(2), and a non-neighboring Ti atom as Ti(3). We adopt the following criteria to distinguish the polaronic states:

- i. The charge state of the whole system: neutral, 1+, 2+;
- ii. The charge state of the V₀ [17], considering the number of electrons that occupy a defect state whose density is centered on the V₀: V_0^{0} , V_0^{1+} , V_0^{2+} ;
- iii. The number of polarons localized on Ti atoms: 0-polaron, 1-polaron, or 2-polaron;
- iv. The location of Ti sites at which each polaron is localized: Ti(1), Ti(2), or Ti(3);

In a previous study, we reported one defect state with both electrons localized in the vacancy without the formation of small polarons [31]. As shown in Fig. 2(b), this case is denoted as the 'neutral, V_0^0 , 0-polaron' state. In addition, we discovered that two other polaronic states can be stabilized after relaxation, by manually changing the initial atomic positions. As shown in Fig. 2(c-d), these two configurations correspond to the formation of one and two polarons, respectively. The calculated formation energies of V_0 (Table II) indicate that the 'neutral, V_0^{2+} , 2-polaron, Ti(1) and Ti(2)' state is the most stable one, while the 'neutral, V_0^0 , 0-polaron' and 'neutral, V_0^{1+} , 1-polaron, Ti(1)' states are metastable. Recently, ESR studies on the triplet ground state of neutral Vo in rutile TiO2 confirmed that the two electrons trapped by Vo are localized on two separate Ti neighbors, forming a pair of exchange-coupled Ti³⁺ ions aligned along the [001] direction [46]. Thus, the simulations indicated the correct polaronic ground state of neutral Vo among the three possible solutions, in perfect agreement with experimental results. Fig. 2(e) shows the metastable configuration with one polaron formed and migrated to a non-neighboring Ti atom, Ti(3). Compared to Fig. 2(c), the formation energy of the migrated-polaron configuration is higher, which can be attributed to the attractive Coulomb interactions between the electron polaron and the positively-charged (1+) Vo. These observations suggest that isolated oxygen vacancy tends to form bound polarons in neutral charge state [46].

In the 1+ charge state on the other hand, two possible polaronic states were found after atomic relaxations, as shown in Fig. 2(f-g). Following similar notations they are denoted as '1+, V_0^{1+} , 0-polaron' and '1+, V_0^{2+} , 1-polaron, Ti(1)', respectively. The '1+, V_0^{2+} , 1-polaron, Ti(1)' state has a lower formation energy, which is consistent with the ESR results of the singlet ground state of positively-charged oxygen vacancy [49]. Furthermore, the electron densities of this singlet state is roughly half of the electron densities of the triplet ground state in neutral charge state, which explains why the hyperfine matrix **A** of the triplet state is approximately twice that of the singlet state, as found by ESR [46,49].

Fig. 3(a-d) show the electron density of states (DOS) for the configurations in Fig. 2(b-e). The 'neutral, V_0^0 , 0-polaron' configuration has a defect state at 0.4 eV below the bandgap, consistent with Ref. 31. The 'neutral, V_0^{1+} , 1-polaron, Ti(1)' configuration has the defect state split into two levels that are 0.36

eV apart, with the upper level at 0.21 eV below the conduction band minimum (CBM). The 'neutral, $V_0^{2^+}$, 2-polaron, Ti(1) and Ti(2)' configuration restores the single defect level again, which is now only 0.1 eV below CBM. When one polaron is formed and migrated away from the vacancy as shown in Fig. 3(d), the defect state is split further by 0.48 eV, while the upper level begins to merge with CBM. To further demonstrate the nature of the two defect levels in Fig. 3(b) and 3(d), the peak-resolved charge isosurfaces are plotted in Fig. 3(e-f) for 'neutral, $V_0^{1^+}$, 1-polaron, Ti(1)'. It is verified that the deeper defect level (peak-A) corresponds to charge densities centered on the V_0 , while the shallower defect level (peak-B) corresponds to the polaron localized on Ti(1). It is worth noting that the defect level of the small polaron is much shallower compared to that calculated by hybrid functional [17]. This discrepancy could potentially be attributed to the local nature of U. It is also worth mentioning that the Fermi-levels of these systems are elevated compared to pristine TiO₂, generating larger electron concentration near CBM. These results further support the bound-polaron picture of the shallow donor states induced by oxygen vacancy, and also explains the significant enhancement of UV absorption by reduced TiO₂ in photocatalysis [62-63].

Since a polaron is essentially a localized electronic state bound by the electron-phonon interactions, it is expected that multiple metastable states exist, especially in the presence of oxygen vacancies which disrupt the local symmetry. In this study, the atomic coordinates in all polaronic states of one oxygen vacancy are summarized in Table III. When no polaron is formed, the surrounding Ti-Ti distances are slightly larger compared to pristine TiO₂. When one electron polaron is formed at Ti(1), the Ti-O distances around this Ti atom are stretched due to enhanced Coulomb repulsion, breaking also the symmetry between Ti(1) and Ti(2) and causing the split of DOS in Fig. 3(b). At the same time, the Ti-Ti distances around the V_O become larger due to the decrease of negative charges at the vacancy site which mitigates the Coulomb repulsion among Ti cations. Similarly, when a second polaron is formed, the Ti-Ti distances around the V_O are further stretched. The energy required to achieve these lattice distortions is compensated by the reduced Coulomb energy through the spatial and orbital redistributions (e_g to t_{2g}) of electrons.

In ab initio studies of polaronic states, different strategies can be adopted to trap the polarons at specific sites, such as manual structural modifications [64-65], using vacancy substitutions [66], or setting different values for the on-site U [67]. The information from Table III can act as a guideline for manual structural modification to stabilize arbitrary polaron states. A similar but less systematic approach has been explored recently to stabilize a bipolaron state on the surface of rutile TiO₂ [65].

IV. VACANCY-VACANCY INTERACTIONS

As two V_0 approaches each other, the excess electrons and corresponding lattice distortions will start to interact. An extreme case is when the two V_0 migrate into the same TiO₆ octahedron and share one Ti atom as their nearest neighbor. In this work, we studied all the five non-equivalent configurations that belong to this category (labeled as C1-C5, as shown in Fig. 5(a)). The TiO₆ octahedron is the basic building block of various crystal structures of TiO₂, including rutile, anatase and brookite. In addition, four other double-vacancy configurations with the two V_0 located in separate TiO₆ octahedra were studied (labeled as C6-C9, see Fig.5(b)). The supercell with only one V_0 is referred to as the non-interactive case, i.e. when the V_0 - V_0 distance is very large. Based on the atomic relaxation data from Table III, all the double-vacancy configurations were positioned into 9 different polaronic states (3 polaronic states for each V_0). These initial configurations were then fully relaxed to search for the ground states.

To determine the most stable configurations, the interaction energy (E_{int}) is calculated with respect to the non-interactive reference, using the following formula:

$$E_{\rm int} = E_{2V_{\rm O}} + E_{\rm TiO_2} - 2E_{\rm IV_{\rm O}} \tag{1}$$

Where E_{TiO2} is the total energy of the pristine supercell and $E_{1\text{VO}}$, $E_{2\text{VO}}$ are the total energies of the supercell including one or two vacancies respectively. The polaronic states with the lowest interaction energies are identified as the ground state of each configuration and Fig. 5(c) shows the calculated E_{int} of each configuration's ground state, plotted as a function of the V_O-V_O distances.

According to Fig. 5(c), the interaction energies of all configurations are calculated to be negative. This suggests that two isolated V₀ exhibit attractive interaction when they approach each other. A simplified way to understand this attractive interaction is to view the oxygen vacancy and its excess electrons as a dipole. When this dipole approaches an ion or another dipole, its direction and amplitude can be adjusted through the redistribution of excess electrons. The distance dependence of such charge-dipole and dipole-dipole interactions can be estimated as $\sim 1/r^2$ and $\sim 1/r^6$, respectively [68].

When the distance between two V_0 becomes very small, they can no longer be treated as two free-rotating dipoles. Instead, the space between them is too small for all the excess electrons to freely redistribute, as in the case of C1-C5. Thus, the Coulomb repulsion between two positively-charged vacancies should become stronger as they cannot be fully mediated by excess electrons. This argument is confirmed by the calculation results of C1-C5, as E_{int} increases with decreasing Vo-Vo distance. Among them, the most stable configurations are C4 and C5. The two V_0 in these cases are located in the opposite directions of the Ti atom, with V_0 -Ti- V_0 angles of 180 degrees. On the other hand, C1-C3 have V_0 -Ti- V_0 angles of approximately 90 degrees. Among them, C3 has the largest Vo-Vo distance and also lowest interaction energy. The distance dependence of unscreened Coulomb repulsion is $\sim 1/r$. Thus, the interaction energies are fitted with the following equation to verify the proposed physical mechanisms:

$$E_{int}(r) = A/r - B/r^2 - C/r^6$$
⁽²⁾

As shown in Fig. 5(c), satisfactory fitting result is obtained, with A = 0.753 eV/Å, $B = 11.83 \text{ eV/Å}^2$ and $C = -383.45 \text{ eV/Å}^6$. Now, this set of parameters can be applied to evaluate Vo-Vo interactions in TiO₂ beyond *ab initio* calculations.

In order to reveal the microscopic origin of the interactions between vacancies, we further analyzed the electronic wavefunctions by plotting the ELF. Fig. 6 compares the ELF of the ground states and the second-most stable polaronic states of C1-C3 and C6-C9. As for C4 and C5, only the ground state is obtained from relaxations. In these cases, the ground state of each configuration always has more small polarons formed at nearby Ti atoms compared to the second-most stable state.

At longer V_0 - V_0 distances (C5-C9), the ELF exhibits similar patterns as two isolated 'neutral, 2-polaron' states. In these cases, the lowering of interaction energies can be attributed to the reduction of Coulomb repulsion through charge redistribution. On the other hand, at smaller V_0 - V_0 distances (C1-C3) the ground state no longer resembles two 'neutral, V_0^{2+} , 2-polaron' states, since the two V_0 share the same Ti atom as their nearest neighbor. Now the two vacancies must eject at least one

electron polaron to a distant Ti atom to reach the ground state. In these cases, the lowering of interaction energies is partly due to the reduction of Coulomb repulsion, and partly due to the sharing of lattice distortion between two V_0 .

It is worth noting that this work is primarily focused on systems with one or two polarons. The ELF suggests that the ground state usually has the polarons localized in the close vicinity of the V_0 . In previous studies, it has been shown that polarons can be rather mobile in rutile TiO₂ [56,69] and not necessarily bound to V_0 . For future studies, it would be interesting to check if this is still the case for systems with a larger V_0 concentrations (e.g. the Magnéli phases).

Finally, we would like to discuss the effects of polaronic interactions on the conductivity of TiO₂. Fig. 7 shows the spin-polarized electron density of states (DOS) of configurations C1, C4, C5, C8 and C9, chosen to represent various V_0 - V_0 distances. The positions of defect levels are quite different in these cases: (1) C1, C4 and C5 (Fig. 7(a-c)) exhibit very strong V_0 - V_0 interactions. Defect states are considerably split and the highest defect levels merge with the conduction band minimum (CBM). (2) C8 (Fig. 7(d)) has two oxygen vacancies that are farther apart (~6Å). The splitting of defect states is still observed, but to a lesser extent compared to C1, C4 and C5. The highest defect level is located at 0.1eV below CBM, similar to the 'neutral, V_0^{2+} , 2-polaron' state of a single V_0 . (3) C9 (Fig. 7(e)) has two well-separated V_0 (~9Å), and its DOS looks similar to the 'neutral, V_0^{2+} , 2-polaron' state of a shallower defect states. Since shallower defect states have lower donor-activation energies, we may conclude that the clustering of oxygen vacancies will increase the conductivity of rutile TiO₂.

V. CONCLUSIONS

In this work, we report an in-depth study of the vacancy-vacancy interactions in rutile TiO_2 using density functional theory with on-site Coulomb corrections (DFT+U), and extensive search of potential polaronic states. The stability of the di-vacancy systems in each polaronic state is determined, and the ground state at each vacancy-vacancy distance is predicted.

Our results suggest that polaron formation plays a crucial role in the formation and interactions of oxygen vacancies in rutile TiO_2 . Two isolated vacancies exhibit attractive interactions, with an equilibrium distance at about 4Å. The attractive forces between vacancies arise from the polaronic effects which redistribute the excess electrons to minimize their Coulomb repulsion. These new discoveries provide better understanding to the vacancy clustering phenomena from a microscopic point of view.

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REFERENCES

[1] A. L. Linsebigler, G. Lu and J. T. Yates Jr., Chem. Rev. 95, 735 (1995).

[2] M. Ni, M. K. H. Leung, D. Y. C. Leung and K. A. Sumathy, Renewable and Sustainable Energy Reviews 11, 401 (2007).

[3] R. Asahi, T. Morikawa, T. Ohwaki, K. Aoki and Y. Taga, Science 293, 269 (2001).

[4] S. U. M. Khan, M. Al-Shahry Jr and W. B. Ingler, Science 297, 2243 (2002).

[5] S. G. Kumar and L. G. Devi, Journal of Physical Chemistry A, 115, 13211 (2011).

[6] G. K. Mor, K. Shankar, M. Paulose, O. K. Varghese and C. A. Grimes, Nano Letters 6, 215 (2006).

[7] K. Zhu, N. R. Neale, A. Miedaner and A. J. Frank, Nano Letters 7, 69 (2007).

[8] M. Kadoshima, M. Hiratani, Y. Shimamoto, K. Torii, H. Miki, S. Kimura and T. Nabatame, Thin Solid Films 424, 224 (2003).

[9] I.G. Baek, M.S. Lee, S. Seo, M.J. Lee, D.H. Seo, D.-S. Suh, J.C. Park, S.O. Park, H.S. Kim, I.K. Yoo, U.-I. Chung and J.T. Moon, in IEDM Tech. Dig. 2004, p. 587.

[10] D. B. Strukov, G. S. Snider, D. R. Stewart and R. Stanley Williams, Nature 453, 80 (2008).

[11] D.-H. Kwon, K. M. Kim, J. H. Jang, J. M. Jeon, M. H. Lee, G. H. Kim, X.-S. Li, G.-S. Park, B. Lee, S. Han, M. Kim and C. S. Hwang, Nature Nanotech. 5, 148 (2010).

[12] H.-S. P. Wong, H.-Y. Lee, S. Yu, Y.-S. Chen, Y. Wu, P.-S. Chen, B. Lee, F. T. Chen and M.-J. Tsai, Proc. of the IEEE 100, 1951 (2012).

[13] E. Yagi, R.R. Haiguti and M. Aono, Phys. Rev. B 54, 7945 (1996).

[14] L. Forro, O. Chauvet, D. Emin, L. Zuppiroli, H. Berger, and F. Levy, J. Appl. Phys. 75, 633 (1994).

[15] M. K. Nowotny, T. Bak, and J. Nowotny, J. Phys. Chem. B 110, 16270 (2006).

[16] N. A. Deskins, R. Rousseau and M. Dupuis, J. Phys. Chem. C 114, 5891 (2010).

[17] A. Janotti, C. Franchini, J. B. Varley, G. Kresse and C. G. Van de Walle, Phys. Status Solidi RRL 7, 199 (2013).

[18] I. G. Austin, N. F. Mott, Advances in Physics 18, 41 (1969).

[19] S. Yang, A. T. Brant, N. C. Giles and L. E. Halliburton, Phys. Rev. B 87, 125201 (2013).

[20] S. Andersson, B. Collen, U. Kuylenstierna and A. Magnéli, Acta Chemica Scandinavica, 11, 1641 (1957).

[21] J. Y. Shin, J. H. Joo, D. Samuelis and J. Maier, Chemistry of Materials, 24, 543 (2012).

[22] W.-Q. Han and Y. Zhang, Appl. Phys. Lett. 92, 203117 (2008).

[23] X. Tao, J. Wang, Z. Ying, Q. Cai, G. Zheng, Y. Gan and Y. Cui, Nano letters 14, 5288 (2014).

[24] J. R. Smith, F. C. Walsh and R. L. Clarke, Journal of applied electrochemistry 28, 1021 (1998).

[25] T. Ioroi, Z. Siroma, N. Fujiwara, S. I. Yamazaki and K. Yasuda, Electrochemistry Communications, 7, 183 (2005).

[26] R. F. Bartholomew and D. R. Frankl, Phys. Rev. 187, 828 (1969).

[27] I. Leonov, A. N. Yaresko, V. N. Antonov, U. Schwingenschlögl, V. Eyert, and V. I. Anisimov, Journal of Physics: Condensed Matter 18, 10955 (2006).

[28] L. Liborio and N. Harrison, Physical Review B 77, 104104 (2008).

[29] L. Liborio, G. Mallia and N. Harrison, Physical Review B 79, 245133 (2009).

[30] M. Weissmann and R. Weht, Physical Review B 84, 144419(2011).

[31] S. G. Park, B. Magyari-Köpe and Y. Nishi, Phys. Rev. B 82, 115109 (2010).

[32] L. Zhao, S. G. Park, B. Magyari-Köpe and Y. Nishi, Mathematical and Computer Modelling, 58, 275 (2013).

[33] S. Lany and A. Zunger, Phys. Rev. B 80, 085202 (2009).

[34] S. Lany, Phys. Status Solidi B 248, 1052 (2011).

[35] P. Deak, B. Aradi and T. Frauenheim, Phys. Rev. B 83, 155207 (2011).

[36] P. Deak, B. Aradi and T. Frauenheim, Phys. Rev. B 86, 195206 (2012).

[37] A. Janotti, J. B. Varley, P. Rinke, N. Umezawa, G. Kresse, and C. G. Van de Walle., Phys. Rev. B 81, 085212 (2010).

[38] H.-Y. Lee, S. J. Clark, and J. Robertson, Phys. Rev. B 86, 075209 (2012).

[39] J. Robertson and R. Gillen, Microelectronic Eng. 109, 208 (2013).

[40] E. Finazzi, C. Di Valentin, G. Pacchioni and A. Selloni, J. Chem. Phys. 129, 154113 (2008).

[41] B. J. Morgan and G. W. Watson, Phys. Rev. B 80, 233102 (2009).

[42] H. Kamisaka, T. Suenaga, H. Nakamura and K. Yamashita, J. Phys. Chem. C 114, 12777 (2010).

[43] N. Umezawa and J. Ye, Phys. Chem. Chem. Phys. 14, 5924 (2012).

[44] T. S. Bjørheim, A. Kuwabara, and T. Norby, J. Phys. Chem. C 117, 5919 (2013).

[45] A. M. Czoska, S. Livraghi, M. Chiesa, E. Giamello, S. Agnoli, G. Granozzi, E. Finazzi, C. Di Valentin and G. Pacchioni, J. Phys. Chem. C 112, 8951 (2008).

[46] A. T. Brant, E. M. Golden, N. C. Giles, S. Yang, M. A. R. Sarker, S. Watauchi, M. Nagao, I. Tanaka, D. A. Tryk, A. Manivannan and L. E. Halliburton, Phys. Rev. B 89, 115206 (2014).

[47] S. Yang, L. E. Halliburton, A. Manivannan, P. H. Bunton, D. B. Baker, M. Klemm, S. Horn, and A. Fujishima, Appl. Phys. Lett. 94,162114 (2009).

[48] F. D. Brandao, M. V. B. Pinheiro, G. M. Ribeiro, G. Medeiros-Ribeiro, and K. Krambrock, Phys. Rev. B 80, 235204 (2009).

[49] A. T. Brant, N. C. Giles, S. Yang, M. A. R. Sarker, S. Watauchi, M. Nagao, I. Tanaka, D. A. Tryk, A. Manivannan and L. E. Halliburton, J. Appl. Phys. 114, 113702 (2013).

[50] E. Kioupakis, P. Rinke, A. Janotti, Q. Yan and C. G. Van de Walle, Chapter 8: Energy Conversion: Solid-State Lighting, in Computational Approaches to Energy Materials (John Wiley & Sons Ltd, Oxford, UK, 2013).

[51] G. Kresse and J. Hafner, Phys. Rev. B 47, 558 (1993); G. Kresse and J. Hafner, Phys. Rev. B 49, 14251 (1994).

[52] P. E. Blochl, Phys. Rev. B 50, 17953 (1994).

[53] V. I. Anisimov and O. Gunnarsson, Phys. Rev. B 43, 7570 (1991).

[54] I. A. Nekrasov, M. A. Korotin and V. I. Anisimov, arXiv:cond-mat/0009107.

[55] L. Zhao, S. G. Park, B. Magyari-Köpe and Y. Nishi, Applied Physics Letters 102, 083506 (2013).

[56] M. Setvin, C. Franchini, X. Hao, M. Schmid, A. Janotti, M. Kaltak, C. G. Van de Walle, G. Kresse and U. Diebold, Phys. Rev. Lett. 113, 086402 (2014).

[57] M. Cococcioni and S. Gironcoli, Phys. Rev. B 71, 035105 (2005).

[58] A. D. Becke, J. Chem. Phys. 92: 5397 (1990).

[59] N. A. Deskins and M. Dupuis, Phys. Rev. B 75, 195212 (2007).

[60] S. P. Ong, V. L. Chevrier, and G. Ceder, Phys. Rev. B 83, 075112 (2011).

[61] H. Ding, H. Lin, B. Sadigh, F. Zhou, V. Ozoliņš and M. Asta, J. Phys. Chem. C 118, 15565 (2014).

[62] G. Wang, H. Wang, Y. Ling, Y. Tang, X. Yang, R. C. Fitzmorris, C. Wang, J. Z. Zhang and Y. Li, Nano Letters 11, 3026 (2011).

[63] F. Zuo, L. Wang, T. Wu, Z. Zhang, D. Borchardt and P. Feng, J. Am. Chem. Soc. 132, 11856 (2010).

[64] T. Shibuya, K. Yasuoka, S. Mirbt and B. Sanyal, J. Phys.: Condens. Matter 24, 435504 (2012).

[65] T. Shibuya, K. Yasuoka, S. Mirbt and B. Sanyal, J. Phys. Chem. C 118, 9429 (2014).

[66] N. A. Deskins, R. Rousseau and M. Dupuis, J. Phys. Chem. C 115, 7562 (2011).

[67] X. Hao, Z. Wang, M. Schmid, U. Diebold and C. Franchini, Phys. Rev. B 91, 085204 (2015).

[68] P. W. Atkins, and J. De Paula, Physical Chemistry: Thermodynamics, Structure, and Change, 2014.

[69] P. M. Kowalski, M. F. Camellone, N. N. Nair, B. Meyer and D. Marx, Phys. Rev. Lett. 105, 146405 (2010).

Defect Type	$e_i(N)$ (eV)	<i>E(N)-E(N-1)</i> (eV)
Rutile: V ₀	2.097	2.193
Rutile: Al _{Ti}	0.036	0.022

TABLE I. Fullfillment of the generalized Koopmans condition by $LDA+U^d+U^p$ approach (Energies referred to the valence band maximum)

Charge state	Polaronic state	Formation energy (eV)				
	V_0^0 , 0-polaron	3.203				
Neutral	V ₀ ¹⁺ , 1-polaron, Ti(1)	2.768				
	V ₀ ²⁺ , 2-polaron, Ti(1) and Ti(2)	2.490				
	V_0^{1+} , 1-polaron, Ti(3)	2.978				
1+	V_0^{1+} , 0-polaron	3.542				
	V_0^{2+} , 1-polaron, Ti(1)	3.177				

TABLE II. Formation Energies of the Isolated $\mathbf{V}_{\mathbf{O}}$ in Various Polaronic States

		Ti-Ti distances around V ₀ (Å)			Ti-O distances around Ti(1) (Å)				Ti-O distances around Ti(2) (Å)					
Charge state	Polaronic Atomic state pair #	1	2	3	4	5	6	7	8	9	10	11	12	13
Neutral	Pristine TiO ₂	3.54	3.54	2.99	1.95	1.95	1.95	1.95	1.95	1.95	1.95	1.95	1.95	1.95
	V_0^{0} , 0-polaron	3.64	3.64	3.01	1.99	1.96	1.96	1.95	1.95	1.99	1.96	1.96	1.95	1.95
	V_0^{1+} , 1-polaron, Ti(1)	3.73	3.78	3.11	2.03	1.98	2.01	1.98	1.98	1.97	1.93	1.89	1.95	1.95
	$V_0^{2^+}$, 2-polaron, Ti(1) and Ti(2)	3.94	3.94	3.23	2.03	1.95	1.96	1.98	1.98	2.03	1.95	1.96	1.98	1.98
	V ₀ ¹⁺ , Ti(3)	3.78	3.81	3.15	1.94	1.9	1.92	1.94	1.94	1.9	1.84	1.94	1.95	1.95
1+	V_0^{1+} , 0-polaron	3.79	3.79	3.15	1.94	1.89	1.92	1.94	1.94	1.94	1.89	1.92	1.94	1.94
	V_0^{2+} , 1-polaron, Ti(3)	3.92	4.02	3.27	2.03	1.95	2.00	1.97	1.97	1.93	1.81	1.88	1.93	1.93

TABLE III. The Atomic Distances around One Single Oxygen Vacancy in Rutile TiO2 with Various Polaronic States*

* The number of atomic pairs are labeled in Fig. 4.



FIG. 1. (a) Electron localization functions (ELF) which demonstrate small polarons formed at two different Ti atomic sites. (b) Energies of the polaron migration pathway between A and B, calculated by linear interpolation of atomic positions. Insets: the isosurface plot of partial charge densities (isosurface density is 0.1 electron/Å³), representing the small-polaron localization at different Ti sites.



FIG. 2. (a) The atomic configuration of a single V_0 in rutile TiO₂. (b-g) The isosurface plots of partial charge densities (top row) and the electron localization functions of the spin-up states (bottom row) of various configurations: (b) neutral, V_0^{0} , 0-polaron; (c) neutral, V_0^{1+} , 1-polaron, Ti(1); (d) neutral, $V_0^{2^+}$, 2-polaron, Ti(1) and Ti(2); (e) neutral, V_0^{1+} , 1-polaron, Ti(3); (f) 1+, V_0^{1+} , 0-polaron; (g) 1+, $V_0^{2^+}$, 1-polaron, Ti(1).



FIG. 3. (a-d) Electron spin density of states (DOS) of the following configurations: (a) neutral, V_0^{0} , 0-polaron; (b) neutral, V_0^{1+} , 1-polaron, Ti(1); (c) neutral, V_0^{2+} , 2-polaron, Ti(1) and Ti(2); (d) neutral, V_0^{1+} , 1-polaron, Ti(3). The Fermi level is represented by red dashed lines. (e-f) Charge-density isosurface plots of (e) peak-A and (f) peak-B of the 'neutral, V_0^{1+} , 1-polaron, Ti(1)' state.



FIG. 4. Numbered atomic pairs around the single V_0 . The corresponding atomic distances are summarized in the Table III.



FIG. 5. (a) Schematics of the five configurations of which the two V_0 are located in the same TiO₆ octahedron (C1-C5). (b) Schematics of the four configurations of which the two V_0 are located in different TiO₆ octahedra (C6-C9). The oxygen vacancies are represented by black circles. (c) The interaction energies between two oxygen vacancies in each of these configurations, with respect to the non-interactive case.



FIG. 6. The electron localization functions (ELF) of the ground states of C1-C9 and the second-most stable polaronic states of C1-C3 and C6-C9 (excluding C7). The results for C7 is not included because it looked almost exactly the same as C6. The black circles represent the locations of oxygen vacancies, and the golden circles represent small polarons localized at Ti atoms.



FIG. 7. Spin-dependent electron density of states (DOS) of the ground state of (a) C1, (b) C4, (c) C5, (d) C8 and (e) C9. The Fermi level is represented by red dashed lines. (f) shows the DOS of the 'neutral, $V_0^{2^+}$, 2-polaron, Ti(1) and Ti(2)' state as a comparison.