

Properties of Oxygen Vacancy and Hydrogen Interstitial Defects in Strontium Titanate: DFT + $U^{d,p}$ Calculations

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ABSTRACT: This work presents extensive theoretical studies focused on the mixed ion–electron transport in cubic strontium titanate (STO). A new approach to the description of this difficult system was developed within the framework of linear-scaling Kohn–Sham density functional theory, as realized in the ONETEP program. The description we present is free of any empirical parameters and relies on the Hubbard U and Hund’s J corrections applied to both Ti and O atoms. The proposed methodology was validated by considering perfect STO. Its calculated properties were found to be in close agreement with experiments and calculations at higher levels of theory. The validated approach was subsequently employed to study the oxygen vacancy (V_{O}) and the hydrogen interstitial (I_{H}), using very large supercells (625 ± 1 atoms). The relaxed configurations of defects were obtained through fastidious energy minimization and later analyzed from a number of perspectives. The calculated defect formation energies and charge transition levels (CTLs) were found to be in close agreement with the experiment. With the exception of the charge-neutral V_{O} , all considered defects were found to introduce shallow states, located down to 0.2 eV below the conduction band. Our calculations revealed a large 1 eV difference in the thermodynamic and optical CTLs of the neutral V_{O} , explaining the inconsistencies observed—till now—between conduction and spectroscopic measurements. The influence of defects on the bonding characteristics and the crystalline structure of STO was quantified, revealing that both V_{O} and I_{H} defects lead to a significant polarization and strong tilting of the TiO_6 octahedra.

Oxygen vacancy and hydrogen interstitial in SrTiO_3

Theoretical description	Modeling and calculations	Results
ONETEP DFT+ $U+J$ Ti 3d O 2p	$\hat{H}_{\text{eff}} = E_{\text{eff}}$ $E_{\text{IIC}} \Delta V_{\text{PA}}$	formation structure and bonding transitions
motivated and validated	physically sound and reliable	consistent with experiment

1. INTRODUCTION

1.1. Strontium Titanate. Strontium titanate (SrTiO_3 , STO) is a prototype perovskite material. At high temperatures, it crystallizes in a cubic structure (space group $Pm\bar{3}m$, see Figure 1), while at low temperatures, it adopts a tetragonal structure (space group $I4/mcm$) due to an antiferrodistortive phase transition which occurs at 105 K.^{1–3}

In the last few decades, significant effort has been dedicated to understanding the structural,^{4–9} mechanical,^{10–12} optical,^{13–24} and transport^{25–32} properties of STO. The influence of various dopants on these properties has also been extensively studied.^{28,33–35} This high interest in STO is driven by the variety of its potential applications, which span fields such as electronics, optoelectronics, and photocatalysis.³⁶

In recent years, STO has also become the subject of an increased interest in electrochemistry, as it has been demonstrated to be a good base material for the fabrication of the so-called triple-conducting oxides (TCOs).^{34,37,38} These materials are capable of conducting oxygen (O^{2-}) and hydrogen (H^+) ions, and electrons or holes, and are therefore considered promising for use as novel cathodes in proton-conducting fuel cells.^{39,40} One of the examples of promising STO-based TCO is $\text{SrTi}_x\text{Fe}_{1-x}\text{O}_{3-\delta}$, which was found to be

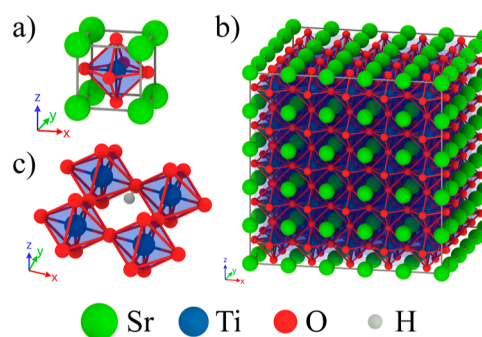
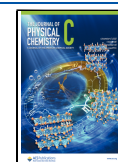


Figure 1. Unit cell of the cubic SrTiO_3 (panel a) and the $5 \times 5 \times 5$ supercell used in the calculations (panel b). Panel c presents the initial positioning of the hydrogen interstitial with respect to the closest TiO_6 octahedra. All visualizations presented in this work were prepared using OVITO.¹⁵⁶

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electrochemically stable, offering high ionic conductivity and good ion surface exchange rates.^{35,41–43}

In the last few years, significant advances were made in the experimental studies focusing on the synthesis, testing, and application of various TCOs.^{37–40,44} However, their better understanding still requires considerable theoretical work because even the base materials are still poorly understood in the context of mixed electron-ion transport. This knowledge gap mainly originates from the difficulties encountered in theoretical modeling studies.⁴⁵ The related issues can be well illustrated using the example of STO.

1.2. Difficulties in Theoretical Description. Understanding the mixed transport properties of STO requires a suitable description of all three constituent types of transport. However, even a description of the static properties of charge carriers constitutes a challenging task, as the theoretical methods used to date have been either incapable of describing the electronic structure correctly or have been too computationally expensive for studies concerning defects.

The local and semi-local density functional theory (DFT) methods that have been used typically in this context are known to poorly describe strong electron correlation effects, which leads to a poor description of the band structure with significantly underestimated band gaps. For STO, DFT calculations employing the local density approximation (LDA) and the generalized gradient approximation (GGA) for exchange–correlation (XC) interactions typically yield an indirect band gap of 1.6–2.0 eV,^{46–52} underestimating the experimental value (3.25 eV³¹) by as much as 40–50%.

Methods based on higher levels of theory (e.g., DFT using hybrid functionals^{46,47,49,53–57} or various GW approaches^{58–61}) perform considerably better in describing the band structure (band gaps very close to the experiment were reported in refs 49, 58, 59, and 61). However, because of their extremely high computational cost, they cannot, as of today, be fully applied to defects.

The investigation of defect properties needs to begin with structure relaxation, which is a highly demanding task, even when local and semi-local DFT methods are used. It cannot be performed with the higher level of theory applied throughout the entire minimization with the currently available computational resources. The apparent exceptions found in the literature⁶² report on the defect properties calculated at a higher level of theory but for structures optimized at a lower level.

One of the possible remedies to this accuracy-cost problem is the application of DFT + U approaches. These methods are characterized by a computational cost comparable to semi-local DFT, while providing significant improvement in the quality of the description, which is often comparable to DFT with hybrid functionals. The last two decades have witnessed considerable advances in the theory of DFT + U .^{63–65} As a consequence, DFT + U has gained significant popularity, becoming a standard tool for treating difficult (strongly correlated) systems containing transition-metal atoms.⁶⁶

Within DFT + U , the standard DFT energy functional is augmented by a Hubbard term, whereby additional Coulomb repulsion is applied between the strongly localized (typically d and f) electrons belonging to the same Hubbard atom. The magnitude of this interaction is proportional to the occupation numbers and is controlled by the Hubbard U parameter. In so doing, DFT + U addresses the self-interaction error, a well-known deficiency of standard DFT.

DFT + U can be extended to DFT + U + J by additionally correcting the interactions between unlike spins and adding an appropriate exchange interaction term in the corrective functional. The magnitude of thus introduced Hund's exchange term is controlled by another parameter, typically denoted with J . The + J correction has been demonstrated to be crucial in describing materials with strong spin–orbit coupling.^{63,64}

The application of DFT + U + J requires a prior determination of the correction parameters, U and J . Their choice can be motivated on theoretical grounds using the linear response approach,⁶⁷ its recent reformulations,^{68,69} or other approaches.^{70,71}

Although theoretical foundations for finding U and J exist, in many cases these parameters are determined empirically by fitting to experimental properties, like the band gap width.⁶⁶ Such an approach often leads to unphysical (i.e., very large) U values,^{72–76} often resulting in a poor description of the properties which were not taken into account when choosing U . More often than not, the values of U and J are transferred between various systems containing the same corrected element,⁷⁷ without regard for their dependence on the material. It should be emphasized that U and J are also sensitive to the methodology, in the sense that they depend on the details of the theoretical approach used, as the exact form of the corrective functional, the form of the projectors used to calculate the occupancies for the correction, and the XC functional used.

Because of the above, the (U , J) values used in the literature often differ significantly between different studies concerning the same material.⁶⁶ For example, two different (U , J) sets were used for correcting Ti 3d electrons in STO (all given in eV): (5.0, 0.64) proposed by Pavarini et al.⁷⁸ (used in refs 48, 77, 79–87) and (3.2, 0.90) proposed by Solovyev et al.⁸⁸ (used in refs 48, 80, and 89). We believe that neither of the above sets is, in fact, well justified, as they were transferred from studies concerning LaTiO₃ (Solovyev's set, Pavarini's set) and YTiO₃ (Pavarini's set). There are also other choices made by individual authors.^{62,75,90–92}

This inability to describe the underlying physics in a justified manner is reflected in the published results. The existing studies, which employed DFT + U (and + J) methods and focused on oxygen vacancies (V_{O}) in STO, have provided conflicting reports concerning the location and nature of defect levels introduced by the neutral V_{O} . DFT + U studies reported levels that are shallow or located within the conduction band (CB), while higher-level calculations typically predicted deeper locations (0.7 eV below the CB). This problem has been recently extensively addressed by Ricca et al.,⁹³ who questioned many of the previous results obtained for V_{O} in STO with DFT + U approaches. These failures can be attributed to the fact that standard DFT + U (which, when applied to STO, corrects only Ti 3d electrons through on-site terms only) is still unable to faithfully reproduce the band structure of perfect STO, offering only a minor improvement over semi-local DFT and yielding band gaps (typically 2.0–2.4 eV), which are still in poor agreement with the experiment, among other deficiencies.

The last few years have provided new developments in the corrective approaches. The proposed extensions (e.g., methods such as DFT + U + V ,⁹⁴ which also corrects off-site interactions) are now emerging as a valuable tool for studying STO and similar challenging systems. Ricca et al.⁹³ very

recently demonstrated that DFT + U + V is capable of reproducing STO's band gap width with an accuracy comparable to that of hybrid DFT calculations. Crucially, this was done using U and V parameters determined on theoretical grounds.

Recent work by O'Regan et al.⁹⁵ has shown that a similar improvement can also be achieved within the DFT + U + J framework. They demonstrated that the application of the + U + J corrections also to the 2p electrons of the oxygen atoms (in addition to Ti 3d electrons) considerably improves the predicted fundamental band gap of TiO₂, which constituted a long-standing challenge for DFT calculations. The band gaps calculated agreed almost perfectly with the experiment (differing by only 0.01 and 0.03 eV, for rutile and anatase, respectively). Reference 95 also achieved this using well-justified methods of choosing the corrective parameters. The U and J parameters were determined from the theory using the so-called minimum-tracking linear-response (MT-LR) method. This approach—also proposed recently by O'Regan and co-workers⁶⁹—can be considered as an analogue of the conventional linear response approach of Cococcioni and de Gironcoli,⁶⁷ with their procedure being adapted to the case of direct-minimization codes, like ONETEP^{96,97} used in refs 69 and 95.

1.3. Difficulties in Modeling of Defects. The accurate theoretical description is not the sole difficulty. By default, the calculations on point defects employ periodic boundary conditions (PBCs) and the so-called supercell approach. The use of PBCs means that the simulated defect is in fact a periodic array of defects. Particular care must be taken to guarantee that the calculated properties are not affected by spurious defect–defect interactions (mechanical, electrostatic, or magnetic).

This problem can be overcome—at least in principle—by extrapolating to infinity the results obtained for a series of finite systems, thus reaching the dilute defect limit. However, the available computational power constraints the system sizes tractable in practice. These typically do not exceed 320 atoms (corresponding to a supercell containing $4 \times 4 \times 4$ replicas of the five atom unit cell) for perovskite-type materials, for semi-local DFT methods or methods with a similar cost.

Additionally, not all system sizes are equivalent. A study carried out by Zhang et al.⁸⁶ has shown that there are better and worse system sizes $n \times n \times n$. This conclusion has been reached based on the calculated defect formation energies (V_O in STO was considered). Energies obtained for even n 's turned out to be systematically higher than those obtained for odd n 's. Reference 86 explained this effect referring to the relaxation of the crystal structure, which occurred after the introduction of the defect. The possible relaxation was limited in supercells with even values of n , which restricted the rotation of the TiO₆ octahedra.

Studies concerning defects often consider their various charge states in order to predict which state is energetically preferred at the given conditions specified by the Fermi level.^{98–103} By doing so, other quantities of interest—like thermodynamic and optical charge transition levels (CTLs)—can also be accessed.

In the above context, charge-neutral defects do not pose significant difficulty. In contrast, charged defects—which most often constitute the main interest—require additional treatment, that is, the calculation of an image interaction correction (IIC), with the aim of removing the spurious contributions to

the energy which originate from the electrostatic interactions of the defect charge density with its periodic images (an artifact of PBCs).

Another correction must be applied in calculations on charged defects performed in PBCs. In such calculations, a uniform neutralizing background charge, the so-called jellium, must be introduced to make the Ewald summation convergent. The introduction of jellium means that the electrostatic potentials of charged and neutral defects cannot be directly compared (the introduction of jellium corresponds to setting the average potential to zero). Therefore, before evaluating the formation energy of the charged defect, a so-called potential alignment (PA) must be performed to correctly account for the energy required to exchange the electrons in the process of defect charging.

A number of approaches to computing both corrections (IIC and PA) have been proposed over the last three decades.^{98,99,104–117} More often than not, these corrections are either omitted^{81,83} or are treated without the diligence they deserve (the underlying assumptions are not verified), even though they might give contributions to the energy which are significant in the context of mixed electron–ion transport. This requires verification.

Comparing theoretical calculations with the experiment also constitutes a challenge, and there are several reasons for that.

Theoretical calculations are typically carried out at 0 K. Therefore, comparing the properties measured in the experiment requires extrapolating results obtained for finite temperatures down to 0 K, which may introduce discrepancies. For example, the experimental determination of defect formation energies can be done by measuring the temperature dependence of conductivity and applying the methods of defect chemistry, which rely on a number of assumptions.²⁹

Differences between the conditions used in experiments and those assumed in calculations also manifest themselves through differences in the chemical potentials. Typically, in theoretical calculations, chemical potentials are computed, and as such they are known precisely. However, they are affected by the errors originating from the limitations of the chosen theoretical description. On the other hand, in the experiment chemical potentials are difficult to precisely control and measure. Many properties of defects, for example, their formation energies, strongly depend on chemical potentials, which, in fact, specify thermodynamic conditions. An appropriate comparison between calculations and experiments requires ensuring that the same conditions are considered, that is, that both use the same sets of chemical potentials.

Some possibilities of comparison are also provided—at least in principle—by the measurements of optical properties. The locations of defect levels, that is, the electron states introduced by defects are accessible from calculations. However, also here, considerable difficulties arise. Again, the related problems can be well illustrated on the example of STO.

With the exception of the charge-neutral V_O , the considered defects (oxygen vacancies in the 2+ and 1+ charge states, hydrogen interstitial) are problematic for optical observation because they do not introduce any states, or only introduce states which are very close to the CB (≈ 0.1 eV below). This proximity makes it challenging to experimentally obtain reliable information, which could be used to validate the theory.

The charge-neutral V_O introduces states, which might be seen as deep (≈ 1.1 eV below the CB^{14,15}). However, some

optical experiments measured shallow locations (≈ 0.1 eV below the CB¹⁶). The conduction measurements of the ionization energies also predicted a shallow location.^{25,26} This significant discrepancy is the reason why the charge-neutral V_O attracted considerable theoretical interest.^{53,56,57,62,77,81–83,86,87,93,118–132}

However, the calculations carried out to date could not explain the discrepancies seen in the experimental characterization of the charge-neutral V_O . Various theoretical studies gave locations of defect states which differed significantly and disagreed with spectroscopic and conduction measurements. This problem was discussed in detail in ref 93, to which we refer the interested reader. We will return to this problem later in this article, as it constitutes an essential aspect of this work. The methodology developed and applied in this work was partially validated by considering this problematic case of the charge-neutral V_O . We believe that a careful validation of the proposed approach should always constitute an indispensable test of its adequacy and quality.

Although there are many theoretical reports focusing on the oxygen vacancies in STO, the properties of hydrogen interstitials were not a subject of any broader study. The existing reports have serious limitations: using small supercells^{72,73,133–135} (typically $2 \times 2 \times 2$ or $3 \times 3 \times 3$) and/or employing inadequate descriptions (local and semi-local DFT, or DFT + U with unphysically large $U = 8.5$ eV,^{72,73,134,136} or even simpler methods^{137–139}). Furthermore, to the best of our knowledge, no single study focused on all three types of charge carriers (electrons, oxygen and hydrogen ions), studying them on the same theoretical footing within the context of mixed electron-ion transport. That hinders the assessment of their role in STO.

1.4. Aims and Outline of the Work. Building on the above observations and recent theoretical developments, we decided to fastidiously investigate the oxygen vacancy and hydrogen interstitial defects in STO. Our focus on these defects is motivated by the long-term scope of the research undertaken, a theoretical characterization of various STO-based (and similar) TCOs, with a particular interest in their mixed electron-ion transport. From this perspective, this work paves the way for future studies, constituting a proof of concept focused on developing and validating the underlying methodology (presented in Section 2).

In particular, we studied V_O and I_H defects in STO using the DFT + $U^{d,p}$ approach, recently proposed by O'Regan et al.⁹⁵ After validating the methodology (on perfect STO, Section 3), we considered a number of charge states of these defects, performed their geometry relaxation and characterized their various properties. Here, particular care was taken to correctly account for all essential theoretical and computational ingredients discussed in the above introduction. The results of this extensive study are presented in Section 4, where we report on formation energies, electronic structures, defect levels, CTLs, structural relaxations, and other properties. We conclude with a discussion and summary in Section 5, outlining the directions for future works.

2. METHODOLOGY

2.1. ONETEP Approach. All the calculations reported in this article were performed using ONETEP^{96,97} (order- N electronic total energy package), a linear-scaling approach for quantum-mechanical calculations based on DFT.

Within ONETEP, the cubic scaling limitation of conventional DFT is overcome by working directly with the density matrix

$$\rho(\mathbf{r}, \mathbf{r}') = \sum_{\alpha, \beta} \phi_{\alpha}(\mathbf{r}) K^{\alpha\beta} \phi_{\beta}^*(\mathbf{r}') \quad (1)$$

which is constructed from localized orbitals ϕ_{α} and ϕ_{β} centered on atoms. The orbitals are nonorthogonal-generalized Wannier functions (NGWFs), which can be chosen to be real-valued. They are expressed in an underlying basis of periodic cardinal sine (psinc) functions defined in real space on a regular Cartesian grid. The grid spacing d_{psinc} controls the quality of the basis set and can be related to the kinetic energy cutoff E_{cut} of traditional plane-wave (PW) DFT. Another parameter that affects the quality of the computations is the localization radius of the NGWFs (R_{ϕ}), beyond which they strictly vanish.

The linear-scaling property of the ONETEP approach is a consequence of the so-called nearsightedness of electronic matter.^{140,141} This phenomenon makes the density kernel $K^{\alpha\beta}$ short-ranged in systems with a large band gap. Therefore, it can be truncated for $|\mathbf{R}_{\alpha} - \mathbf{R}_{\beta}| > R_K$, leading to a highly sparse matrix representation, when large systems are studied. The parameter R_K is known as the density kernel cutoff.

The sparsity of $K^{\alpha\beta}$ and the localization of $\{\phi_{\alpha}\}$ are exploited to obtain a linear-scaling formalism. For this purpose, ONETEP utilizes sparse matrix algebra and the so-called FFT box technique.¹⁴² Here, as we only want to outline the most critical aspects of ONETEP approach briefly, we omit a detailed discussion of the underlying formalism, referring the interested reader to refs 96, 97, 143–147.

Within ONETEP, the ground state is found as in standard DFT, that is, through the variational principle. The minimization of the system's total energy is achieved using a two-level loop. The inner loop minimizes the energy with respect to the density kernel elements with the NGWFs fixed, while the outer loop optimizes the psinc coefficients of the NGWFs. The minimization continues until the convergence with respect to both quantities—that is, $K^{\alpha\beta}$ and $\{\phi_{\alpha}\}$ —is achieved.

In ONETEP—similarly to other approaches to DFT—the quality of the obtained results also depends on the basis set used to represent the orbitals. In this respect, the ONETEP approach is similar to the linear combination of atomic orbitals (LCAO) approach, as it also uses atom-centered basis functions, specifically the NGWFs. A different number of NGWFs can be assigned to different atoms. The type (i.e., symmetry) and the spatial extent of NGWFs can also be controlled, making it possible to work with basis sets of different qualities (like split-valence basis sets) and characteristics (like bases containing diffuse functions).

What strongly distinguishes the ONETEP approach from LCAO approaches is the fact that during the calculation the shapes of the NGWFs change, as they are optimized in situ. Consequently, the basis functions adapt to the local chemical environment experienced by the atom on which they are centered. This approach—as it facilitates expressing the sought Kohn–Sham eigenlevels—significantly improves the description of various systems, which within ONETEP can be accurately modeled with near-complete basis set accuracy even with minimal basis sets. The use of a minimal basis also reduces the computational cost, facilitating studying large

systems, which in the case of ONETEP may contain hundreds of thousands of atoms, depending on the system type and chosen methodology (see ref 96 and works cited therein).

2.2. Theoretical Description. In this work, we used the ONETEP approach to study defects (oxygen vacancy and hydrogen interstitial) in STO. We used the DFT method within the GGA with the Perdew–Burke–Ernzerhof (PBE)¹⁴⁸ XC functional, which was augmented with the Hubbard + U and Hund's + J corrections using the DFT + $U^{\text{d,p}}$ approach.⁹⁵ We motivate our choice of the PBE functional with the fact that in the calculations carried out, we also had to consider H₂ and O₂ molecules (used as the reference states in the calculation of defect formation energies). The versatility of PBE is the reason why we preferred it over GGA functionals that were specifically designed to model solids (like PBEsol¹⁴⁹).

The innermost core electrons of Sr, Ti, and O were treated with norm-conserving pseudopotentials obtained from the Rappe library.^{150,151} For Ti atoms, these sets do not include 3s or 3p semicore states, which were treated explicitly. Therefore, we used 13 NGWFs for each Ti atom, with the initial electron configuration being that of a neutral atom ($3s^2 3p^6 3d^2 4s^2 4p^0$). The same number of 13 NGWFs was used for Sr atoms (initial electron configuration $4s^2 4p^6 4d^0 5s^2 5p^0$). In both cases (Sr and Ti), the additional p-type NGWFs served as diffuse functions and were found by numerical experiments to be necessary for obtaining a well-converged ground state. For all O atoms, we used 4 NGWFs ($2s^2 2p^4$ initial electron configuration). The hydrogen atom was described with one NGWF function.

For all NGWFs, we used the same localization radius $R_\phi = 12 a_0$, where a_0 denotes the Bohr radius. The density kernel was not truncated. The cutoff energy was taken as $E_{\text{cut}} = 1090$ eV, meaning the NGWFs were expanded with a $d_{\text{psinc}} = 0.43 a_0$ spacing. The above values ensured well-converged results, as was concluded from the numerical tests performed for the perfect system.

The use of a relatively high energy cutoff is motivated by the fact that the performed calculations mainly comprised geometry optimizations, which require accurate forces for stable convergence. For the same reason, the stopping criterion for NGWF optimization was set to a more stringent value of $8 \times 10^{-7} E_h a_0^{-3/2}$ (E_h denotes the Hartree energy) and the density kernel was optimized on each update of the NGWFs and also in trial steps taken during the conjugate-gradient-based optimization of NGWFs. Denser than default working grids were also used for expressing the charge density and potential (with 340 grid points in each coordinate direction) to reduce numerical noise. Γ -Point sampling was used for reciprocal space integration. In the modeled supercells, this corresponds to sampling on a Γ -centered regular grid containing $5 \times 5 \times 5$ (or $7 \times 7 \times 7$, see later) points in the first Brillouin zone (FBZ).

In our studies, we also considered systems with an odd number of electrons (oxygen vacancy with a $q = 1 +$ formal charge and neutral hydrogen interstitial) and systems with locally broken spin symmetry (singlet and triplet states of the neutral oxygen vacancy and a negatively charged hydrogen interstitial). Therefore, all the calculations were performed in the spin-polarized scheme, treating both spin channels independently. We note that within ONETEP spin-polarization is introduced through the dependence of the density kernel $K^{\alpha\beta}$ on spin, but the basis functions (NGWFs) are spin-

independent. The employed spin-polarization was also a consequence of the fact that we applied + J corrections.

What distinguishes our approach from other DFT + U + J studies on STO is that we applied + U and + J corrections to both Ti (3d subspace) and O (2p subspace) atoms, following the guidelines provided in refs 95, 101, 152, and 153. Using the + U + J corrections requires choosing the so-called Hubbard projectors, which directly enter the definition of Hubbard and Hund's terms (therefore, affecting the effectiveness of the + U + J corrections), as they are used to calculate occupancies of the strongly localized orbitals being corrected. This is done by projecting the calculated Kohn–Sham states (molecular orbitals or bands) onto a basis of localized atomic-like states. In this work, we used Hubbard projectors corresponding to the atomic NGWFs (obtained from the pseudoatomic solver), according to a method proposed by O'Regan et al.^{144,145} and implemented in ONETEP. Here, we underline the particular importance of using localized Wannier-like projectors, which—according to many authors—is the most appropriate choice.^{69,95,154,155} Our procedure did not include projector self-consistency.¹⁴⁴

Although there are some sets of U and J parameters for correcting Ti in STO available in the literature, to the best of our knowledge, there are no parameters suitable for correcting oxygen 2p electrons in STO. Here, similarly to refs 64 and 69, we emphasize that the considered parameters are known to be case-specific, in the sense that they should not be transferred between different systems and, what is also important, between different methodologies. For these reasons, at the very beginning of our studies, we determined all the required parameters (i.e., U_{Ti} , J_{Ti} , U_{O} and J_{O}) from scratch for STO and the chosen methodology.

2.3. Calculations Performed. All the calculations were performed at zero temperature. A supercell consisting of $5 \times 5 \times 5$ repetitions of the 5-atom cubic unit cell (Figure 1) was used, with $N = 625$ atoms in the perfect system. Where clearly indicated, a $7 \times 7 \times 7$ supercell (with $N = 1715$ atoms) was used instead. PBCs were used in all directions. The lattice constant was set to the experimental value of $a = 3.91 \text{ \AA}$.³⁷

Our simulation protocol consisted of the following steps. At the very beginning, we determined the U and J parameters for Ti and O atoms. This was done by studying the response of the perfect system using the MT-LR method (see Supporting Information, Section S1).

The obtained parameters were subsequently validated by comparing the results obtained with the chosen methodology with the results obtained with different methodologies and with experimental data. Our validation included comparing measures describing the band structure (band gap width E_g , total and partial densities of states) and measures characterizing the structure and energetics of the perfect system. To this end, we have calculated the equation of state (EOS) and, based on it, determined the equilibrium lattice constant a , the bulk modulus B_0 , and its pressure derivative $B'_0 = \partial B_0 / \partial p$ (both for zero pressure).

With U and J parameters at hand, we moved to systems containing defects, studying the oxygen vacancy with a formal charge of $q = 0, 1+$, and $2+$ (denoted with V_{O}^0 , V_{O}^{1+} , and V_{O}^{2+}) and the hydrogen interstitial with a formal charge of $1-$, 0 and $1+$ (I_{H}^{1-} , I_{H}^0 , and I_{H}^{1+} , respectively). For the neutral oxygen vacancy, we considered two spin states, with two “excess” electrons (i.e., electrons remaining after the removal of the neutral O atom and located on the two neighboring Ti atoms)

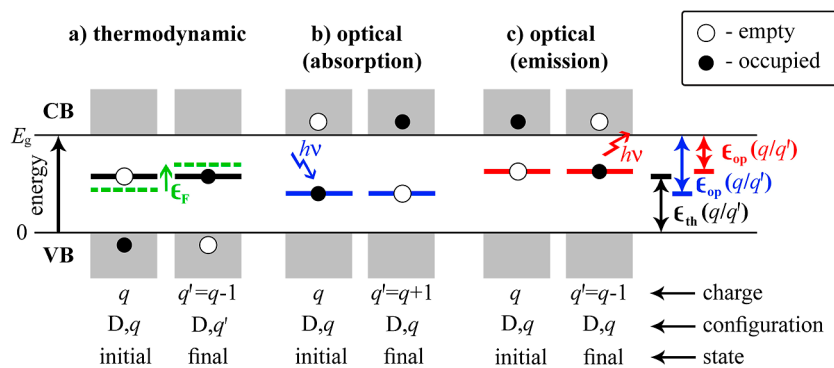


Figure 2. Thermodynamic (a) and optical (b,c) CTLs. In the thermodynamic transition, the increase of the Fermi level ϵ_F above the corresponding TTL ($\epsilon_{th}(q/q')$, see eq 3) causes a $q \rightarrow q' = q - 1$ charge transition. This process corresponds to a thermally induced excitation of an electron (obtained from the host material's valence band) to a state introduced by the defect. The opposite process may also occur during the lowering of ϵ_F . In the optical transition, the absorption (or emission) of a photon causes an excitation (or recombination) of the electron, which passes from the defect's level to the CB of the host material (or vice versa). Both optical processes result in a change of the defect's charge: $q \rightarrow q' = q + 1$ (for the absorption) or $q \rightarrow q' = q - 1$ (for the emission). The energy of the absorbed (or emitted) photon defines the location of the corresponding OTL ($\epsilon_{th}(q/q')$, see eq 4). In the thermodynamic transition, both states of the defect (initial and final) correspond to the relaxed configurations (D, q and D, q' , respectively), while in the optical transitions, only the initial state is relaxed (D, q).

having initially either antiparallel (singlet state, $V_O^{0,S=0}$) or parallel (triplet, $V_O^{0,S=2}$) spins. We also considered a non-magnetic singlet state, with two excess electrons located on the same Ti atom and having antiparallel spins. However, we found this state to be highly unpreferable. Singlet and triplet spin states were also considered for I_H^{1-} (denoted as $I_H^{1-,S=0}$ and $I_H^{1-,S=2}$).

For all considered defects, we performed geometry relaxation, which started from the perfect crystal with one O atom removed (V_O^q) or one H atom added (I_H^q). In the initial structure, the hydrogen atom was positioned on the TiO_2 plane, with the O–H bond length and the Ti–O–H valence angle taken as $d_{O-H} = 1 \text{ \AA}$ and $\theta_{Ti-O-H} = 90^\circ$ (Figure 1c).

To check the influence of the initial conditions (and numerical noise) on the obtained minima, we have also performed additional (and independent) geometry optimization, which started from structures with slightly distorted atomic positions (all atoms were displaced by 10^{-3} \AA in random directions). We have considered one additional structure for each charge/spin state of the V_O^q defect and three additional structures for each state of the I_H^q defect. In total, we have considered 4×2 (for V_O) + 4×4 (for I_H) different structures, performing 24 separate geometry optimizations. For this purpose, we employed a combination of two geometry optimization techniques, which were used alternately in a loop.

In the beginning, the optimization was carried out in the delocalized internal coordinates¹⁵⁷ (DI), according to a method proposed in ref 158. Through numerical experiments, we found this method to be more efficient in the early stages of geometry relaxation.

The structure obtained from the DI minimization was later subjected to further relaxation, carried out in Cartesian coordinates with the Broyden–Fletcher–Goldfarb–Shanno (BFGS) algorithm.^{159–161} In these two initial stages of relaxation, we used the following convergence criteria $\Delta E < 10^{-7} E_h/\text{ion}$ (change in the system's total energy between two successive iterations), $F_{\max} < 2 \times 10^{-4} E_h/a_0$ (the maximum force acting on any ion), and $\Delta R_{\max} < 5 \times 10^{-4} a_0$ (maximum displacement of any ion between two successive iterations), requiring all of them to be met simultaneously at convergence.

This two-stage process—consisting of repeated DI- and BFGS-based minimizations—was continued until no significant drop (less than 3 meV) in the system's total energy was observed throughout four consecutive minimizations (DI-, BFGS-, DI-, and BFGS-based). During this stage, more stringent (twice as strict as before) convergence criteria were used in the individual DI/BFGS minimizations. The criterion for NGWF convergence was also tightened to $2 \times 10^{-7} E_h a_0^{-3/2}$, to avoid “empty” iterations (corresponding to updates of the geometry with no significant update of the electronic structure).

Here, we note that the simulation cell was kept constant, as changing it would change the grid spacing d_{psinc} resulting in varying E_{cut} and, in turn, the basis set. However, we performed a complete relaxation of the atomic coordinates, that is, without constraining the positions of any (even distant) atoms. The performed complete relaxation allowed us to observe effects such as structural polarization and octahedral tiltings and to account for their influence on the defect's properties.

2.4. Formation Energies. The formation energies of defects were calculated using the well-known Zhang and Northup formula,¹⁶² according to which the formation energy of a defect D in charge state q is given as^{99,114}

$$\Delta H(q, \epsilon_F) = E_{D,q} - E_P - \sum_i n_i \mu_i + q(\epsilon_{VBM} + \epsilon_F) + E_{IIC} + E_{PA} \quad (2)$$

Here, $E_{D,q}$ and E_P represent the total energies of the defective and perfect systems, respectively. With n_i , we denote the number of removed ($n_i < 0$) or added ($n_i > 0$) i -type atoms, μ_i refers to their chemical potential in the chosen reference state, and the third term in eq 2 represents contributions to ΔH that are due to the exchange of the atoms with the environment. With ϵ_{VBM} and ϵ_F , we denote the energy level of the valence band maximum (VBM) of the perfect system and the location of the Fermi level (with respect to the VBM), respectively. The corresponding fourth term represents the energy change originating from the exchange of electrons (charging of a defect).

The two remaining terms, E_{IIC} and E_{PA} , represent corrections. These corrections are applied a posteriori in

calculations on charged defects in PBCs to remove artificial contributions to $E_{D,q'}$ which arise from spurious interactions between the considered charged defect, its periodic images, and the neutralizing background charge. The first of these (E_{IIC}) is the so-called IIC, while the latter (E_{PA} , typically written as $q\Delta v_{\text{PA}}$) is the PA term. Both of these can be calculated using various approaches of different sophistication.^{98,99,104–117}

When determining the IIC and PA corrections, we used a very recent method proposed by Durrant et al.¹¹⁴ This method (for details, see [Supporting Information](#), Section S3) is in many aspects similar to the well-known Freysoldt–Neugebauer–Van de Walle (FNV) procedure,¹⁰⁵ as it also relies on the comparison of the charge densities (determination of IIC) and electrostatic potentials (determination of PA) of perfect and defected (both neutral and charged) supercells. However, it does not assume any functional form of the charge density associated with the defect's charge (like a Gaussian for the FNV procedure) and therefore, it is better suited to situations where the obtained defect is difficult to describe analytically (we found this to be the case in our calculations).

2.5. Charge Transition Levels. The calculated formation energies ΔH were used to determine CTLs (see [Figure 2](#) for their graphical interpretation). This was done based on the framework proposed by Lany and Zunger.⁹⁹

The thermodynamic transition levels (TTLs), $\epsilon_{\text{th}}(q/q')$, were calculated from the equation

$$\epsilon_{\text{th}}(q/q') = \frac{\Delta H(q', \epsilon_{\text{F}} = 0) - \Delta H(q, \epsilon_{\text{F}} = 0)}{q - q'} \quad (3)$$

The TTL $\epsilon_{\text{th}}(q/q')$, which in the above expression is calculated relative to VBM of the perfect system, describes a thermally induced charge transition $q \rightarrow q'$ (electron excitation or trapping). $\epsilon_{\text{th}}(q/q')$ corresponds to the Fermi level at which the formation energies of the defect in charge states q and q' become equal. Here, because thermal excitations are considered, the defect is allowed to relax in both charge states q and q' . Therefore, both formation energies from [eq 3](#) are evaluated for the relaxed configurations. Based on $\epsilon_{\text{th}}(q/q')$, the thermal ionization energy of the defect can be determined. For example, the distance of $\epsilon_{\text{th}}(1+/0)$ from CBM (and of $\epsilon_{\text{th}}(0/1-)$ from VBM) measures the thermal ionization energy of simple donors (and acceptors).

We have also calculated the optical transition levels (OTLs), $\epsilon_{\text{op}}(q/q')$, which measure the optical absorption (or emission) energy related to a $q \rightarrow q'$ excitation (or recombination). The $\epsilon_{\text{op}}(q/q')$ levels (relative to CBM, with $\epsilon_{\text{F}} = E_{\text{g}}$) were calculated from the equation

$$\epsilon_{\text{op}}(q/q') = \frac{\Delta H(q', \epsilon_{\text{F}} = E_{\text{g}})|_{D,q} - \Delta H(q, \epsilon_{\text{F}} = E_{\text{g}})}{q - q'} \quad (4)$$

Because optical (vertical) transitions are considered (which occur over extremely short time scales), both formation energies are evaluated for the atomic configuration corresponding to the initial charge state q , according to the Franck–Condon principle.

The expression for $\epsilon_{\text{op}}(q/q')$ can be written in the following form

$$\epsilon_{\text{op}}(q/q') = \frac{E_{D,q'}|_{D,q} - E_{D,q}}{q - q'} - \epsilon_{\text{CBM}} \quad (5)$$

Here, ϵ_{CBM} stands for the location of the conduction band minimum (CBM), while $E_{D,q'}|_{D,q}$ and $E_{D,q}$ represent the total energies of two supercells containing the defect with a charge state q' and q , respectively. In the interest of brevity, we omitted electrostatic corrections E_{IIC} and E_{PA} in [eq 5](#). They were, however, calculated (using Durrant's procedure, see [Supporting Information](#), Section S3) and added to the two total energies present in [eq 5](#).

Based on Janak's theorem,¹⁶³ the energy difference in [eq 5](#) can be approximated with:^{100,102,103}

$$E_{D,q'}|_{D,q} - E_{D,q} \approx \frac{1}{2}[\epsilon_{N+1}^{\text{KS}}(q', N+1)|_{D,q} + \epsilon_{N+1}^{\text{KS}}(q, N)] \quad (6)$$

which holds for $q' = q - 1$ (recombination), and with

$$E_{D,q'}|_{D,q} - E_{D,q} \approx -\frac{1}{2}[\epsilon_N^{\text{KS}}(q', N-1)|_{D,q} + \epsilon_N^{\text{KS}}(q, N)] \quad (7)$$

which is valid for $q' = q + 1$ (excitation). Here, $\epsilon_i^{\text{KS}}(q,N)$ denotes the i -th single-particle Kohn–Sham eigenvalue of an N -electron system with charge q . The notation $|_{D,q}$ serves as a reminder that the eigenvalue must be evaluated for the atomic configuration corresponding to charge state q . The eigenvalues, which appear in [eqs 6](#) and [7](#), correspond to the highest occupied ($\epsilon_{N+1}^{\text{KS}}(q', N+1)$ and $\epsilon_N^{\text{KS}}(q, N)$) and lowest unoccupied ($\epsilon_{N+1}^{\text{KS}}(q, N)$ and $\epsilon_N^{\text{KS}}(q', N-1)$) molecular orbitals. Similar—although more sophisticated—expressions can be derived for transitions (recombinations/excitations), which correspond to the removal/addition of two electrons (with $q' = q \pm 2$). The notation used in [eqs 6](#) and [7](#) does not distinguish spin channels. It is noted that the relevant (majority- or minority-spin) eigenvalues should be considered.

The approximations defined by [eqs 6](#) and [7](#) can be used to calculate $\epsilon_{\text{op}}(q/q')$ based on the Kohn–Sham eigenvalues ϵ_i^{KS} . It must be emphasized that—because of the spurious defect–defect interactions—the eigenvalues calculated for charged systems must be also corrected. This can be done using the formula^{103,111}

$$\Delta\epsilon_{\text{d}}^{\text{KS}} = -\frac{2}{q}(E_{\text{IIC}} + E_{\text{PA}}) \quad (8)$$

which defines the correction that must be added to the eigenvalue corresponding to the defect state $\epsilon_{\text{d}}^{\text{KS}}$. Here, the electrostatic corrections (E_{IIC} and E_{PA}) were also calculated using Durrant's procedure (see [Supporting Information](#), Section S3).

In the literature there are many reports (see e.g., refs [100](#) and [103](#)) which employed the approximations defined by [eqs 6](#) and [7](#) in the determination of OTLs $\epsilon_{\text{op}}(q/q')$. More often than not, the accuracy of these approximations was not tested (for a discussion of their correctness see ref [164](#)). In this work, we verified their validity practically by comparing the approximated OTLs with those calculated using the exact expression given by [eq 5](#) (for results of this comparison, see [Supporting Information](#), Section S4).

3. METHODOLOGY VALIDATION—PERFECT STO

3.1. Validation Outline. From the MT-LR method, we obtained the following values of the U and J parameters (for details, see [Supporting Information](#), Section S1): $U_{\text{Ti}} = 4.111$ eV, $J_{\text{Ti}} = 0.374$ eV, $U_{\text{O}} = 9.097$ eV, and $J_{\text{O}} = 0.935$ eV. To

Table 1. Properties of the Perfect STO as Predicted by Various Approaches Considered in This Work, Compared with Other Theoretical Calculations, and Experimental Data (for Room Temperature)^{a,b}

method	property				
	<i>a</i> (Å)	<i>B</i> ₀ (GPa)	<i>B</i> ' ₀	<i>E</i> _g ⁱ (eV)	<i>E</i> _g ^D (eV)
this work					
DFT (PBE)	3.966	161.9 (0.2)	4.37 (0.02)	1.66	1.96
DFT + <i>U</i> ^d	3.986	154.2 (0.3)	4.35 (0.02)	1.95	2.32
DFT + <i>U</i> ^{d,p}	3.957	172.1 (0.2)	4.25 (0.02)	2.92	3.20
other computational studies					
(a) DFT (all with PBE)					
GTO	3.93 ^a , 3.96 ^b	171 ^{a,b}		1.8 ^b , 1.99 ^a	2.1 ^b , 2.35 ^a
LAPW	3.94 ^{c,d} , 3.95 ^e	167 ^c , 169 ^c , 175 ^d	4.61 ^c	1.63 ^d , 1.97 ^c	
PW	3.94 ^{b,f,g} , 3.95 ^h	168 ^{f,r} , 169 ^b , 171 ^h		1.8 ^{b,f} , 1.85 ^h	2.1 ^b , 2.2 ^{f,r}
hybrid	3.94 ⁱ	172 ⁱ		1.74 ⁱ	
(b) DFT + <i>U</i> ^d (all with PBE and PW)					
<i>U</i> _{eff} = 4 eV	3.97 ^l	164 ^j	4.35 ^j	≈2.0 ^l	
<i>U</i> = 5 eV, <i>J</i> = 0.64 eV	3.90 ⁸ , 3.93 ^k , 3.98 ^l			2.39 ^j , 2.68 ^k	
(c) hybrid DFT					
B3LYP, GTO	3.94 ^a , 3.95 ^m	179 ^m , 180 ^a		3.57 ^a , 4.38 ^m	3.89 ^a
B3PW, GTO	3.89 ⁿ , 3.90 ^{o,p} , 3.92 ^b , 3.93 ^q	190 ^b		3.4 ^b , 3.6 ⁿ , 3.63 ^o , 3.64 ^p	3.7 ^b , 3.96 ^p
HSE06, PW	3.90 ^{f,r}	192 ^{f,r}		3.01 ^r , 3.07 ^{f,r}	3.47 ^{f,r}
PBE0, GTO	3.90 ^b	201 ^b		3.8 ^b	4.2 ^b
PBE0, PW	3.90 ^b	193 ^b		4.0 ^b	4.4 ^b
(d) various GW approaches (all based on LDA)					
QSGW				3.32 ^s	
<i>G</i> ₀ <i>W</i> ₀				3.40 ^t	3.76 ^t
QSGW80 + SO				3.56 ^u	
QSGW + LP + 0.8Σ				3.2 ^v	3.7 ^v
experimental data	3.905 ^{w,x}	178.8 (4.6) ^y	4.31 (0.1) ^y	3.25 ^z	3.75 ^z

^aValues given in brackets denote errors, which for the calculated lattice constants were lower than 10⁻³ Å. ^bReferences: a, ⁴⁶ b, ⁴⁹ c, ⁵¹ d, ⁵⁰ e, ¹⁶⁸ f, ⁴⁷ g, ⁸⁶ h, ⁴⁸ i, ⁵² j, ⁶² k, ⁸³ l, ⁸⁴ m, ⁵³ n, ⁵⁶ o, ⁵⁵ p, ⁵⁴ q, ¹²⁵ r, ⁵⁷ s, ⁵⁸ t, ⁵⁹ u, ⁶⁰ v, ⁶¹ w, ⁹ x, ³⁷ y, ¹² z. ³¹

validate these and the chosen approach, we calculated the electronic ground state of perfect STO at the experimental density (*a* = 3.91 Å). We also calculated its EOS by finding the ground state for eight other lattice constants. Here, to maintain the same accuracy, we had to restrict ourselves to system sizes that differ by an integer number of psinc grid spacings *d*_{psinc}.^{165–167}

To directly demonstrate how the chosen approach (employing +*U* +*J* corrections for both Ti and O atoms, denoted as DFT + *U*^{d,p}) improves the description of STO, similar calculations were also performed for two different methodologies. Within these, (i) the +*U* +*J* corrections were not applied (neither for Ti nor for O) or (ii) were applied only to Ti atoms (with the *U* and *J* parameters for Ti taken as in DFT + *U*^{d,p}). These two approaches will be denoted as DFT and DFT + *U*^d. The first one corresponds to a standard DFT calculation (GGA-based, without any corrections). The second one corresponds to a typical approach from the literature, in which only Ti atoms are corrected.^{48,62,72–77,79–87,89–92} In the DFT and DFT + *U*^d calculations, all other (i.e., not related to +*U* +*J* corrections) parameters were the same as in the DFT + *U*^{d,p} calculations to permit a direct comparison. The EOS calculations (Section 3.2) used a 5 × 5 × 5 supercell. The band structure calculations (Section 3.3) used a 7 × 7 × 7 supercell.

3.2. Equation of State. In Table 1, we present structural and mechanical properties of STO obtained with the three considered approaches, calculated by fitting the obtained *E*(*a*) dependencies (the total energy vs lattice constant, see Figure 3) with the Birch–Murnaghan EOS.

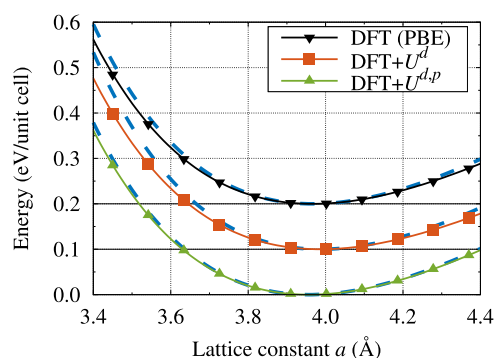


Figure 3. EOS of the perfect STO, calculated with the three considered approaches. Points correspond to the computational results, while solid lines represent the obtained fits (of the Birch–Murnaghan form). Blue dashed lines represent the adjusted experimental EOSs. To increase readability, the curves corresponding to DFT and DFT + *U*^d calculations were offset by +0.1 and +0.2 eV, respectively.

Standard DFT significantly underestimates *B*₀ by 9.4% when compared to the experimental result.¹² Interestingly, the application of the +*U* +*J* corrections for Ti atoms only (DFT + *U*^d) does not improve the bulk modulus and even makes it worse (the calculated *B*₀ is by 13.8% lower than the experimental value). In contrast to these two, the result obtained for DFT + *U*^{d,p} is lower by only 3.8%. For this approach, the calculated pressure derivative of the bulk modulus (*B*'₀) was also found as very close to the experiment being—in fact—within the experimental uncertainty (the same

holds for DFT and DFT + U^d calculations). The calculated lattice parameter a differs from the experimental result significantly (by 0.06–0.08 Å), but for DFT + $U^{d,p}$, the agreement is—still—the best.

The results obtained from the standard DFT calculations deviate slightly from analogous calculations reported in the literature but are comparable. The calculated lattice constant (3.97 Å) and bulk modulus (162 GPa) are higher by 0.02–0.03 Å and lower by 5–9 GPa than those obtained from PW calculations employing the same XC functional (PBE). Our DFT + U^d results ($a = 3.99$ Å, $B_0 = 154$ GPa, and $B'_0 = 4.35$) are similar to those presented in ref 62 (3.97 Å, 164 GPa, and 4.35, respectively), where a similar $U_{\text{eff}} = U - J = 4$ eV was employed (3.74 eV in our case). The results obtained from DFT + $U^{d,p}$ cannot be compared with the literature because no studies to date employed similar methodology.

In Figure 3, we present three calculated EOSs and compare them with the corresponding adjusted experimental EOSs, which were obtained by using (in the Birch–Murnaghan EOS) the experimental values of B_0 (178.8 GPa) and B'_0 (4.31). Their adjustment consisted in using the calculated (for the given approach) equilibrium lattice constant a .

Even a cursory glance reveals an excellent agreement obtained for DFT + $U^{d,p}$. The calculated EOS coincides with the (adjusted) experimental curve almost in the entire range (notable deviations are observed only at high compressions). For the two other approaches, the agreement is visibly much worse.

The presented comparison clearly demonstrates that the chosen methodology (DFT + $U^{d,p}$) significantly improves the description of energetics of STO, with good accuracy obtained in a wide range of strains. The chosen methodology correctly describes mechanical properties of STO both in the linear (reproduced B_0) and non-linear (correct B'_0) regimes. This gives us confidence that the lattice distortions, which appear around defects (and the related energetic effects) will also be correctly described by DFT + $U^{d,p}$.

3.3. Electronic Structure. The examination of the calculated band gaps (Table 1) clearly shows that the application of DFT + $U^{d,p}$ also significantly improves the description of the band structure around the Fermi level. Standard DFT calculations significantly underestimate the experimental indirect band gap (by more than 1.5 eV) and DFT + U^d improves the band gap only slightly (by 0.3 eV). The application of the $+U + J$ corrections also to O atoms yields a band gap, which agrees well with the experiment. Both calculated band gaps (indirect $R \rightarrow \Gamma$, $E_g^I = 2.92$ eV; direct $\Gamma \rightarrow \Gamma$, $E_g^D = 3.20$ eV) compare favorably with the experiment (3.25 and 3.75 eV,³¹ respectively) and with calculations based on a higher level of theory, like those employing hybrid functionals^{46,47,49,53–57,125} or various GW approaches^{58–61} (see Table 1). The calculated valence band width is also in excellent agreement with experimental results of 5.5–6.5 eV.^{169,170} 5.5 eV was obtained using DFT + $U^{d,p}$, while standard DFT and DFT + U^d calculations predicted 4.5 and 4.6 eV, respectively. DFT + $U^{d,p}$ also offers an improved description of the dispersion of the valence and conduction bands. To illustrate this, in Figure 4, we present the Kohn–Sham single-particle energies evaluated at high-symmetry points of the FBZ. Visual inspection reveals that the energies obtained from DFT + $U^{d,p}$ match quasi-particle energies obtained from G_0W_0 calculations.⁵⁹

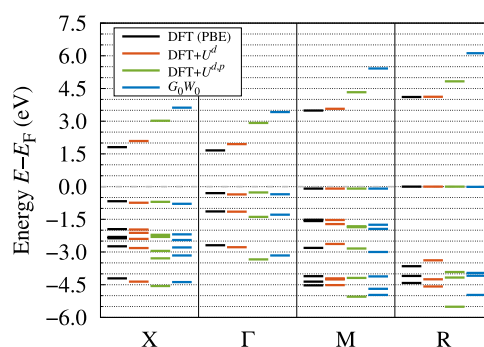


Figure 4. Electronic structure of the perfect STO. Single-particle energies of the nine highest valence bands and the lowest CB are represented with horizontal line segments. Four panels correspond to the X, Γ , M, and R points of the FBZ. Each panel presents results obtained with DFT (black, left), DFT + U^d (red), and DFT + $U^{d,p}$ (green), and also results of G_0W_0 calculations⁵⁹ (blue, right), which serve as a reference. The zero of energy (dashed gray line) was set to the Fermi energy E_F .

To explain this success of DFT + $U^{d,p}$, we use Figure 5 to present the local density of states (l-DOS) calculated for all three considered approaches. A comparison of characteristics obtained for DFT and DFT + U^d shows that the valence band edge essentially remains unaffected when correcting the Ti 3d subshell only. The additional application of corrections to the O 2p subshell lowers its energy. This re-establishes the possibility of hybridization with Ti 3d (see Figure 5), resulting in a significant band gap opening with DFT + $U^{d,p}$. Similar observations were made in ref 95.

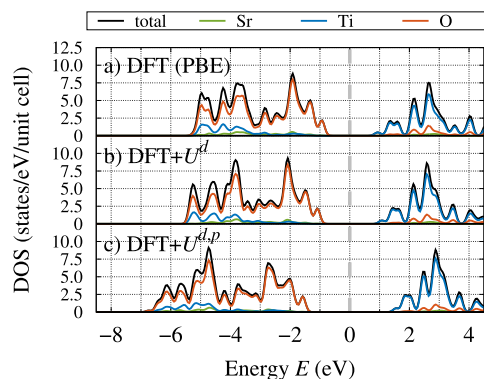


Figure 5. Total and l-DOSs of perfect STO as predicted by various approaches considered in this work. To allow direct comparison, all panels use the mid-gap energy of the DFT calculation for 0 eV (dashed gray line). Gaussian broadening of 0.1 eV was used.

Figure 6 presents the total DOS calculated with DFT + $U^{d,p}$ and its decomposition into local angular-momentum projected densities (lp-DOS).¹⁷¹ The low-lying three narrow maxima (located at -55.7 , -32.6 , and -29.6 eV) correspond to Ti 3s, Ti 3p, and Sr 4s semicore states. Two broad groups of features located at energies between -16 and -12 eV correspond to bands formed from O 2s and Sr 4s levels. The valence band, which starts at -5.5 eV, is mainly composed of O 2p states, with a small contribution of Ti 3d states. The first two groups of conduction states (located between 3 and 8 eV) have a Ti 3d- t_{2g} and Ti 3d- e_g character. Another group of features—originating from the Sr 4d levels—is found above 8 eV.

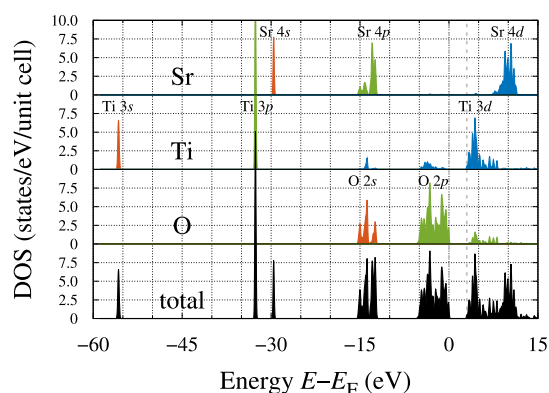


Figure 6. Total DOS of perfect STO and its decomposition into local angular momentum-projected densities (red—s, green—p, and blue—d). The zero of the energy corresponds to the Fermi energy E_F . Two vertical dashed lines indicate the VBM and CBM.

The above results are consistent with G_0W_0 calculations,⁵⁹ which—similarly—located the Ti 3p state at -32.5 eV below E_F and yielded a similar position of the O 2p valence band center. Some quantitative differences are—however—visible in the locations of other bands, which in our picture are positioned at higher energies (see Table 2). The same

Table 2. Electronic Structure of Perfect STO^a

	Ti 3s	Ti 3p	Sr 4s	O 2s	Sr 4p	O 2p
DFT + $U^{d,p}$	-23.1	0	3.0	18.8	19.4	30.0
G_0W_0	0	0	0	16.0	18.1	29.8
XPS	-24.8	0		15.7	18.3	≈31.7

^aFor DFT + $U^{d,p}$ (this work) and G_0W_0 (ref 59), weighted centers of various features are presented (calculated from the lp-DOS). All values are given in eV and are measured relative to the location of Ti 3p state (the sharpest feature) to allow comparison with the XPS experiment of ref 172.

conclusions can be drawn from a comparison with recent X-ray photoelectron spectroscopy (XPS) data,¹⁷² which also provided locations similar to ours. Interestingly, while the position of the Sr 4s state was not identified in ref 172, our DFT + $U^{d,p}$ results suggest that this level may lie slightly (3 eV) above the Ti 3p level. This could explain the asymmetry of

the peak assigned to Ti 3p in the referenced photoelectron spectroscopy experiment.

The presented validation clearly shows that DFT + $U^{d,p}$ significantly improves the description of STO. The calculated measures characterizing its mechanical properties and the electronic structure agree with recent experiments. They are also consistent with recent theoretical calculations based on a higher level of theory, and this can be obtained within DFT + $U^{d,p}$ for a fraction of their cost. The well-reproduced energetics and good reproduction of valence band-related features (character, width, dispersion, forbidden gaps) give us confidence that the defect-related properties (such as formation energies, transition levels, location of defect levels) will also be correctly described by DFT + $U^{d,p}$.

One more issue deserves comment here, namely our choice not to apply the + U + J corrections to Sr 4d states. These states are located high in energy, mostly from the upper CBs, and contribute only marginally to the VB and lower CBs (see Figures 5 and 6). Therefore, it can be expected that the Sr 4d states (and—consequently—their correcting) will have negligible influence on the investigated ground-state properties. We verified this by performing DFT + $U^{d,p}$ calculation in which the Sr 4d states were also corrected (we used $U_{Sr} = 0.934$ eV and $J_{Sr} = 0.152$ eV, which were obtained from the MT-LR method). These calculations showed that the additional correcting of the Sr atoms does not appreciably influence the results. For example, the calculated direct band gap width increased by only 4 meV.

4. RESULTS—DEFECTIVE SYSTEMS

4.1. Structural Relaxation. We begin by discussing the structural relaxations, which proved to be computationally very demanding. In the most difficult cases, we found it necessary to perform up to 15 cycles of DI + BFGS optimizations, with up to 1000–1600 updates of the geometry required to reach the assumed convergence criterion (energy change below 3 meV/2 cycles). One relaxation took between 6 and 9 months of continuous calculations on 16 nodes (equipped with two 14 core processors each). The total wall time of all carried relaxations exceeded 70 M CPU-hours.

The difficulty originates from the very high dimensionality of the considered problem (1869 ± 3 degrees of freedom). However, to a large degree, it is also a consequence of the problem's specificity, that is, the fact that the introduction of a

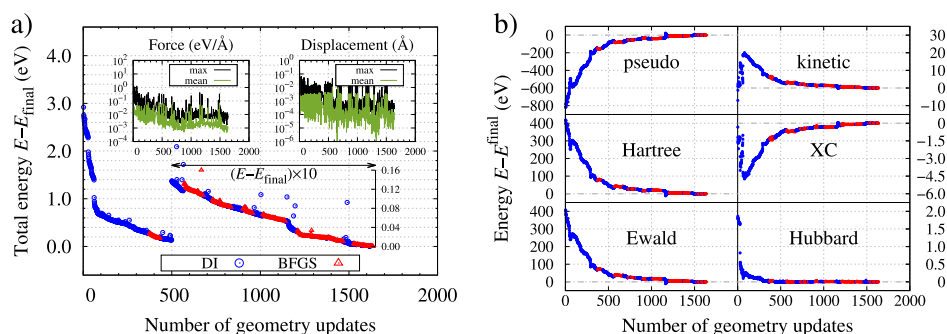


Figure 7. Structural relaxation of the system containing a neutral oxygen vacancy $V_O^{0,S=0}$. Panel (a) presents the evolution of the total energy (relative to the converged value). Note that in the right part (i.e., for energy evaluations >500), the vertical axis was adjusted to better show detail. Two insets present the evolution of forces and displacements (the latter relative to the previous iteration). Panel (b) presents the evolution of the components of the total energy. These are also measured with respect to their converged values. Blue and red identify DI- and BFGS-based minimizations.

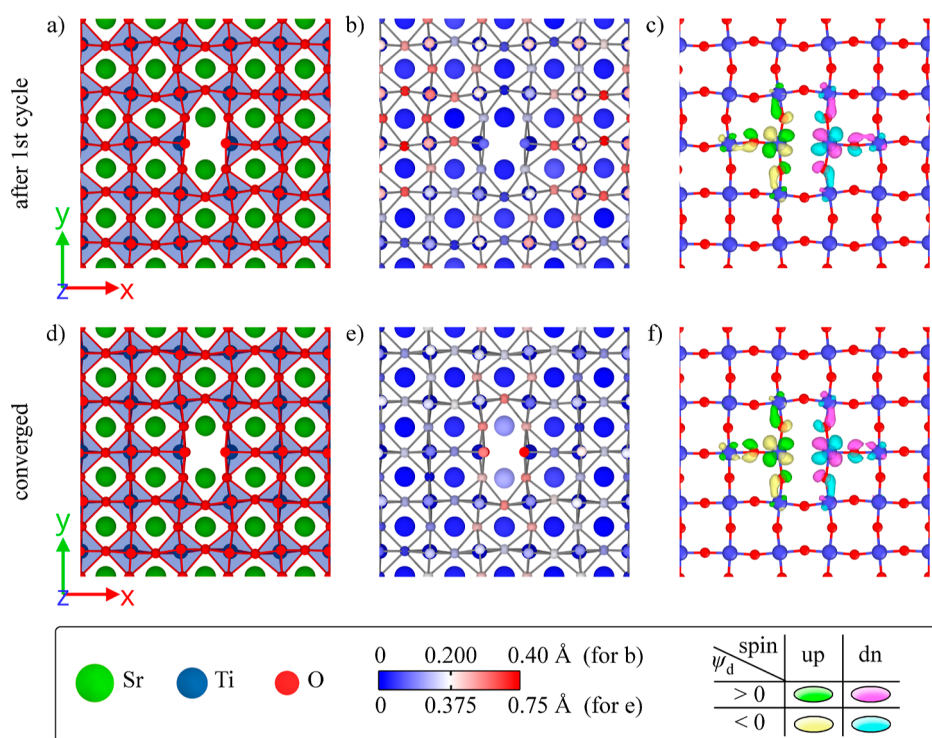


Figure 8. Structural relaxation of the neutral oxygen vacancy $V_{\text{O}}^{0,S=0}$ (cont'd). Configurations obtained after the first minimization cycle (a–c) and at convergence (d–f). All panels present a cross-sectional view through the supercell: the central TiO_2 (001) plane containing the defect (located in the center) is shown, and (for a,b and d,e) SrO (001) plane located above. Panels (b,e) present atomic displacements, measured relative to the converged structure (b) or the initial structure (e). Panels (c,f) visualize two electronic states introduced by the defect, using their single-particle wavefunctions ψ_d^{KS} . Here, isosurfaces $|\psi_d^{\text{KS}}|^2 = 10^{-2} \text{ e}/\text{\AA}^3$ are shown.

Table 3. Minima Identified from the Relaxations Carried out^a

defect state		total energy (eV)	defect level position (eV)	integral of $ \rho^{\uparrow} - \rho^{\downarrow} $ (e)	distances (Å)	
/init.	config.				$\Delta d_{\text{Ti-Ti}}$	$\Delta d_{\text{Sr-Sr}}$
V_{O}^{2+}	d	+404.640	n/a	0	+0.221	+0.144, +0.155
	p	+0.039	n/a	0	+0.240	+0.132, +0.170
V_{O}^{1+}	p	+415.791	-0.099	1.88	+0.209	+0.145, +0.155
	d	+0.155	-0.098	1.88	+0.212	+0.143, +0.160
$V_{\text{O}}^{0,S=0}$	p	+427.045	-0.716, -0.716	2.39	+0.046	+0.134, +0.208
	d	+0.046	-0.719, -0.718	2.45	+0.045	+0.167, +0.186
$V_{\text{O}}^{0,S=2}$	d	+427.044	-0.724, -0.721	2.47	+0.046	+0.165, +0.186
	p	+0.015	-0.719, -0.714	2.45	+0.046	+0.134, +0.208

^aData for various charge/spin states (given in the first column) of the V_{O} defect is presented. The second column indicates if the relaxation started from perfect (p) or distorted (d) positions. The total energies of the identified lowest energy minima (first rows) are measured relative to the perfect system ($E_{\text{D,q}} - E_{\text{p}}$ is given). For other—higher-lying—minima, the energy difference to the corresponding lowest energy minimum is given. Positions of the defect levels are given as $\epsilon_{\text{d}}^{\text{KS}} - \epsilon_{\text{CBM}}$, that is, they are measured with respect to the CBM of the perfect system. The fifth column gives the integral of $|\rho^{\uparrow}(\mathbf{r}) - \rho^{\downarrow}(\mathbf{r})|$ (spin up and spin down densities) over the entire supercell. The last two columns give the changes in the Ti–Ti and Sr–Sr distances for Ti and Sr atoms closest to the defect. For $\Delta d_{\text{Sr-Sr}}$ the minimum and maximum is given.

defect leads to significant rotations and deformations of TiO_6 octahedra, and to notable displacements of the nearest Sr ions. These effects feed back into one another, propagating through the crystal lattice, and significantly affecting the system's energy.

To better illustrate the above problem, we use Figures 7 and 8 to present the evolution of various measures during the relaxation of a $V_{\text{O}}^{0,S=0}$ defect. The data corresponds to the structure whose minimization started from perfect (undistorted) positions. For $V_{\text{O}}^{0,S=0}$, this initial structure led to a lower final energy (however, this is not always the case, see later discussion).

The analysis of the total energy, forces, and displacements (Figure 7a) shows that the employed strategy overcame many local barriers. As expected, the energy drop was the highest in the first minimization cycle (2.8 eV). However, the subsequent minimizations reduced the energy further (by approx. 0.2 eV), yielding a more suitable configuration for calculating CTLs or studying ion transport. For such transport, the energy barriers are of the order of tenths of eV and—therefore—very fine minima are required to obtain reliable (not underestimated) barrier heights.

The validity of the carried out relaxation can also be supported with an analysis of the total energy decomposition (Figure 7b), which shows that all contributions to the total

Table 4. Minima Identified from the Relaxations Carried Out (cont'd)^a

defect state		total energy (eV)	defect level position (eV)	integral of $ \rho^{\uparrow} - \rho^{\downarrow} $ (e)	distances (Å)			angles (degrees)	
/init.	config.				$\Delta d_{\text{Ti-Ti}}$	$\Delta d_{\text{Sr-Sr}}$	$d_{\text{O-H}}$	$\theta_{\text{Ti-O-H}}$	$\theta_{\text{Sr-O-H}}$
I_{H}^{1+}	d	-27.737	n/a	0	+0.283	+0.323	0.956	86.5	44.4
	d	+0.071	n/a	0	+0.298	+0.308	0.957	85.1	43.9
	p	+0.123	n/a	0	+0.300	+0.306	0.957	85.5	43.8
	d	+0.148	n/a	0	+0.301	+0.303	0.957	85.4	43.8
I_{H}^0	d	-16.468	-0.092	1.88	+0.283	+0.317	0.956	86.3	44.3
	p	+0.119	-0.089	1.88	+0.295	+0.301	0.957	85.5	43.8
	d	+0.129	-0.092	1.88	+0.290	+0.308	0.956	85.8	44.0
	d	+0.148	-0.090	1.88	+0.296	+0.300	0.957	85.4	43.7
$\text{I}_{\text{H}}^{1-,S=0}$	d	-5.288	-0.115, -0.115	3.67	+0.282	+0.334	0.963	77.6	46.3
	d	+0.014	-0.108, -0.104	3.66	+0.279	+0.325	0.962	77.9	46.2
	d	+0.092	-0.115, -0.115	3.66	+0.288	+0.326	0.957	86.2	44.3
	p	+0.105	-0.107, -0.105	3.65	+0.280	+0.323	0.956	86.5	44.4
$\text{I}_{\text{H}}^{1-,S=2}$	d	-5.298	-0.116, -0.110	3.75	+0.282	+0.334	0.963	77.5	46.2
	p	+0.009	-0.107, -0.099	3.75	+0.280	+0.324	0.961	78.5	45.9
	d	+0.021	-0.106, -0.099	3.75	+0.279	+0.326	0.962	78.0	46.2
	d	+0.045	-0.096, -0.089	3.73	+0.282	+0.331	0.962	78.1	46.0

^aData for the I_{H} defect. Columns 1–6 have the same meaning as in Table 3. Column 7 presents the change in the distance between two Sr atoms closest to the H atom. The last three columns describe the position of the H atom.

energy converged to their final values before the relaxation was complete.

We note that the main features of the electronic states introduced by the defects, such as their spatial character (described by the corresponding single-particle wavefunction, $\psi_{\text{d}}^{\text{KS}}$) and position within the gap (given by the corresponding single-particle Kohn–Sham eigenvalue, $\epsilon_{\text{d}}^{\text{KS}}$), are already well reproduced after the first minimization cycle. For example, at this point, the two $\psi_{\text{d}}^{\text{KS}}$ states of $\text{V}_{\text{O}}^{0,S=0}$ were located $\epsilon_{\text{CBM}} - \epsilon_{\text{d}}^{\text{KS}} = 0.714$ eV below the CB edge, which is very close to the converged result (difference of <2 meV). After the first DI + BFGS cycle, the single-particle wavefunctions $\psi_{\text{d}}^{\text{KS}}$ were also very similar to those at convergence (see Figure 8).

However, a single-cycle relaxation does not provide an accurate picture of the defect's influence on the crystal structure, with the atomic positions undergoing significant changes during later cycles. For example, for $\text{V}_{\text{O}}^{0,S=0}$, the positions obtained after the first cycle differed by as much as 0.4 Å (and by 0.12 Å on average) from those at convergence (see Figure 8). This clearly shows that multiple DI + BFGS cycles are required to obtain a reliable picture of the defect.

The relaxation proved to be comparatively difficult for hydrogen interstitials, although the opposite could be expected due to the smaller size of I_{H} . This can be explained by the fact that in the considered range of distances (limited by half of the supercell side, ≈ 10 Å), the lattice distortions introduced by I_{H} are comparable to those associated with the presence of V_{O} .

We found the equilibrium structure obtained from the relaxation to depend—to some extent—on the initial configuration. However, as it concerns the defects' static (i.e., not related to their migration) properties, the differences between the minima corresponding to the same defect state were negligible. Tables 3 (data for V_{O}) and 4 (I_{H}) summarize all the relaxations carried out. The differences in the calculated total energies of the corresponding minima are small (below 0.05–0.15 eV). The locations of the defect levels are also comparable (differences in $\epsilon_{\text{d}}^{\text{KS}}$ are below 0.01 eV). The similarity of the spin density distributions shows that the determined ground states are also similar.

For the oxygen vacancy, the measures which characterize lattice distortions that occurred in the very vicinity of the defect (the outward displacements of two closest Ti atoms and four closest Sr atoms) are also comparable between the corresponding minima. The positions of the H atoms are also similar for all minima corresponding to I_{H}^0 , as well as for I_{H}^{1+} .

For $\text{I}_{\text{H}}^{1-,S=0}$, considerable differences are, however, seen in the calculated $\theta_{\text{Ti-O-H}}$ valence angle, which is either ≈ 78 or $\approx 86^\circ$, for the two lowest and two highest energy minima, respectively. Because of this difference, these two pairs of minima can be classified as two distinct motifs. They also differ (by 0.005–0.007 Å) in the calculated $d_{\text{O-H}}$ bond length. Similar motifs (with $\theta_{\text{Ti-O-H}} = 90$ and 76°) were identified by T-Thienprasert et al.¹³⁶ for the I_{H}^{1+} defect.

The absence of the low $\theta_{\text{Ti-O-H}}$ angle motif for the $q = 0$ and $1+$ charge states of I_{H} does not originate from differences in the used initial configurations. The relaxations performed for each charge state of I_{H} used the same initial atomic positions. The underlying reasons are difficult to identify.

In general, we could not ascertain any pattern that would make one initial structure more suitable for the relaxation of V_{O} . For $\text{V}_{\text{O}}^{0,S=0}$ and V_{O}^{1+} , relaxations which started from perfect (undistorted) positions led to a lower final energy. For the remaining states introducing initial distortions proved to be beneficial. This was also the case for I_{H} , for which all identified minimum energy structures corresponded to distorted initial guesses.

4.2. Formation Energies. Table 5 presents the calculated formation energies and the obtained IIC and PA corrections, which were calculated^{108,173–175} using Durrant's procedure¹¹⁴ (for details, see Supporting Information, Section S3).

The IIC corrections proved to be negligible: for both of V_{O} and I_{H} defects, they are in the order of 2–10 meV. The main reason for that is the very high dielectric constant of STO (we assumed 330^{27}). A similar observation—about the practical insignificance of IICs for STO—was made in ref 93.

The obtained PA corrections were proved to be larger than IICs. However, they are still small (typically below ± 0.1 eV), and their neglect would not change the obtained picture of the defects' relative stability. The negligibility of the PA corrections

Table 5. Calculated Formation Energies ΔH (Uncertainty in Brackets) and the Contributions of the IIC and PA Corrections^a

defect state	formation energy		corrections	
	ΔH (eV)	E_{IIC} (eV)	E_{PA} (eV)	
V_{O}^{2+}	+1.45 (0.12)	[0, 0.009]	[-0.062, +0.139]	
V_{O}^{1+}	+4.36 (0.11)	[0, 0.002]	[-0.058, +0.017]	
$V_{\text{O}}^{0,S=0}$	+7.32 (0.02)	0	0	
$V_{\text{O}}^{0,S=2}$	+7.31 (0.01)	0	0	
I_{H}^{1+}	-3.56 (0.13)	[0, 0.002]	[-0.050, +0.063]	
I_{H}^0	-0.54 (0.07)	0	0	
$I_{\text{H}}^{1-,S=0}$	+2.40 (0.11)	[0, 0.002]	[-0.093, +0.068]	
$I_{\text{H}}^{1-,S=2}$	+2.34 (0.07)	[0, 0.002]	[-0.085, +0.058]	

^aHere, we do not distinguish different minima of the same defect. Instead, the results obtained for the various minima were used to determine the presented uncertainties. These uncertainties also account for the uncertainties of the E_{IIC} and E_{PA} corrections, for which bounds are given. Note that both corrections are zero by definition for the neutral defects.

can be explained by referring to the details of the calculations we carried out. The large size of the supercells causes the jellium to have a low charge density and, therefore, it does not appreciably shift the average electrostatic potential.

The formation energies presented in Table 5 correspond to the Fermi level ϵ_{F} set to the VBM, and, crucially, oxygen- and hydrogen-rich limits (the H_2 and O_2 molecules were taken as the reference states, for more details, see Supporting Information, Section S2).

Our DFT + $U^{\text{d,p}}$ calculations reveal that the oxygen vacancy prefers the $q = 2+$ charge state, which has the lowest formation energy (1.45 eV for the oxygen-rich conditions). This result is in line with experimental observations. The formation energies of V_{O}^{1+} and V_{O}^0 are higher by ca. 2.9 and 5.9 eV, respectively, than the formation energy of V_{O}^{2+} . Keeping the uncertainties in mind, the singlet and triplet states of the neutral V_{O} are characterized by the same formation energy, which equals 7.3 eV.

The hydrogen interstitial strongly prefers the 1+ charge state, which has the lowest formation energy (≈ -3.6 eV). Its negative value informs that under the considered conditions (hydrogen-rich conditions, Fermi level set to the VBM), hydrogen will be naturally incorporated into STO in the form of protons, and this also agrees with experimental observations. Although the formation energy of I_{H}^0 was also negative (-0.5

eV), its significantly higher value (by ≈ 3 eV) suggests that in STO, the formation of neutral hydrogen defects will not be as favorable. The formation of negatively charged hydrogen defects is strongly unfavorable (their formation energy is higher by 6 eV than that of I_{H}^{1+}).

We now turn to comparing our results against those of other computational studies. There are many literature reports, which considered the formation of V_{O} using various theoretical approaches. LDA-DFT calculations¹²⁰ yielded significantly smaller formation energies: 4.5 eV (V_{O}^0), 2.4 eV (V_{O}^{1+}), and 0.3 eV (V_{O}^{2+}). Results in line with our calculations were obtained by Carrasco et al.,¹²² with $\Delta H = 7.65$ eV for V_{O}^0 based on GGA-DFT calculations with a PW91 functional. DFT + U calculations predicted a lower formation energy of V_{O}^0 (5.3,⁶² 6.1,⁸⁶ 6.7,⁸³ and 6.8 eV⁷⁷), but a similar—to ours—formation energy of V_{O}^{2+} (1.2 eV⁸³). Formation energies obtained from DFT calculations with hybrid functionals are also lower than those reported here. Reference 53 reported 6.8 eV for V_{O}^0 (obtained with B3LYP), while Mitra et al.⁵⁷ obtained 5.9 eV (V_{O}^0), 3.2 eV (V_{O}^{1+}), and 0.7 eV (V_{O}^{2+}) using the HSE functional. A comparison with other existing studies^{56,123,125} is difficult, as they used a different reference chemical potential μ_{O} .

The above comparison highlights significant discrepancies among the existing reports and our work. However, we found a reasonable agreement with a very recent work of Ricca et al.,⁹³ who used a similar method (DFT + U + V , with self-consistently adjusted U and V parameters). Reference 93 reported $\Delta H = 6.4$ eV for V_{O}^0 (7.3 eV in this work), 3.4 eV for V_{O}^{1+} (here, 4.4 eV), and 0.6 eV for V_{O}^{2+} (1.4 eV in the present study). While the corresponding formation energies differ (in this study, they are higher by ≈ 1 eV), the differences between the formation energies of various charge states are similar. We found that the formation energy of a neutral oxygen vacancy is by 2.9 and 5.9 eV higher than that of a single and double charged vacancy, respectively. Reference 93 reported similar differences: 3.0 and 5.8 eV, respectively. This similarity suggests that the apparent shift of 1 eV in the formation energies may originate from the choice of the reference chemical potential μ_{O} . Reference 93 also used $\mu_{\text{O}} = E_{\text{O}_2}/2$; however, this energy was obtained directly from DFT calculations and not from an adjustment procedure, as in the present work (see Supporting Information, Section S2). This may introduce a difference in the chemical potential μ_{O} and, consequently, a uniform shift in the calculated formation energies. We note that ref 93 found that the same formation

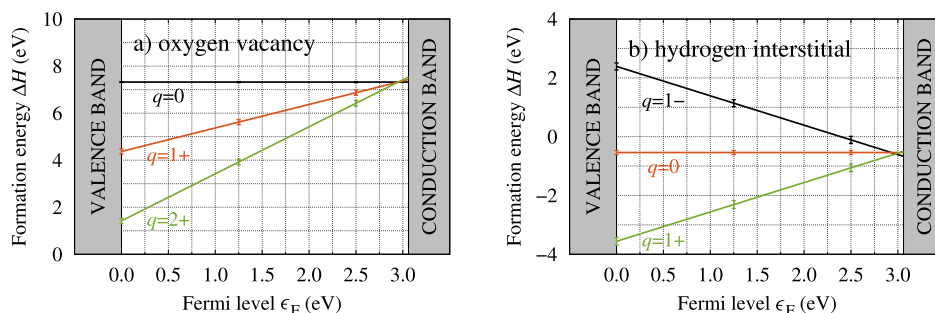


Figure 9. Dependence of the formation energy ΔH on the Fermi level ϵ_{F} . Two panels present the result obtained for the oxygen vacancy (a) and hydrogen interstitial (b). Individual lines correspond to various considered charge states q . The error bars drawn at $\epsilon_{\text{F}} = 0, 1.25,$ and 2.5 eV depict the uncertainties of ΔH . Here, no distinction is made between singlet and triplet states of V_{O}^0 , which gave similar results. The presented data encompasses both cases. The same holds for I_{H}^{1+} .

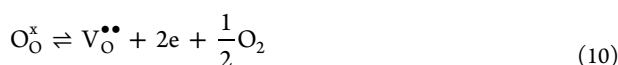
energy characterizes the singlet and triplet states of the neutral oxygen vacancy, and this agrees with our result.

The formation energy of V_{O}^{2+} can be compared with experimental data. ΔH (see eq 2) is defined through the following reaction (written using Kröger–Vink notation¹⁷⁶)



that is, a process in which the formation of a doubly ionized oxygen vacancy (denoted with $V_{\text{O}}^{\bullet\bullet}$, in our notation V_{O}^{2+}) and gaseous oxygen reduces the hole (h) concentration. This process occurs in the region of p-type conductivity (ϵ_{F} close to VBM). Its activation energy (the so-called oxidation enthalpy) was measured by Moos and Härdtl²⁹ to be $\Delta H_{\text{Ox}} \approx 1$ eV (see Section 4.7 therein). A similar value was measured by Choi and Tuller,¹⁷⁷ who obtained $\Delta H_{\text{Ox}} = 1.29$ eV. Our result, $\Delta H(q = 2+, \epsilon_{\text{F}} = 0) = (1.45 \pm 0.12)$ eV, agrees very well with both experimental measurements.

Similarly, the calculated $\Delta H(q = 2+, \epsilon_{\text{F}} \approx E_{\text{g}})$ can also be compared with the experiment, as it corresponds to the activation energy (the so-called reduction enthalpy, ΔH_{Red}) of the following reaction



This reaction, in which two excess electrons are emitted into the CB of the host material, is seen in the region of n-type conductivity (for high ϵ_{F}). Moos and Härdtl²⁹ measured $\Delta H_{\text{Red}} = 6.1$ eV, which compares well with our calculated $\Delta H(q = 2+, \epsilon_{\text{F}} \approx E_{\text{g}}) \approx 7.3$ eV (see Figure 9 for ΔH vs ϵ_{F} dependencies). The observed discrepancy of ≈ 1 eV might originate from the difference in the Fermi level ϵ_{F} (its exact value was not specified in ref 29).

The fact that the calculated formation energies ΔH are higher than the experimental results should be expected. This can be explained by accounting for the limitations of the modeling technique used. The application of the electrostatic corrections means the obtained ΔH is unaffected by spurious electrostatic interactions. However, the calculated total energies still suffer from other finite-size effects. The application of the supercell approach limits the possible long-range relaxation of the structure around the defect, making the obtained total energies higher than those which would be obtained for larger systems (for demonstration, see, e.g., refs 86 and 120) and, consequently, in the dilute defect limit. Thus, we expect the formation energies calculated for finite systems to be higher than those measured in the experiment, as we found in this work for V_{O}^{2+} .

Many of the previous theoretical studies obtained formation energies that were lower than those measured experimentally. In the context of the above discussion, this suggests that the theoretical approach used in these studies must suffer from one or more deficiencies, as it would give even more underestimated formation energies in the dilute defect limit.

Our results obtained for the hydrogen interstitial disagree with those presented by Varley et al.,¹⁷⁸ who obtained $\Delta H = -2.00$ eV (for I_{H}^{1+}), 1.59 eV (for I_{H}^0), and 5.55 eV (for I_{H}^{1-}), based on DFT calculations with hybrid functionals. Their formation energies are higher by ≈ 1.6 eV (I_{H}^{1+}), 2.1 eV (I_{H}^0), and 3.2 eV (I_{H}^{1-}) than the formation energies obtained in this work. This discrepancy can be explained by the observation that ref 178 used a smaller supercell (135 atoms), which strongly limits relaxation. A comparison with other studies

which considered $I_{\text{H}}^{72,73,134,136}$ is difficult, as the formation energies were either not calculated or their presentation omits important details (like the chosen reference state).

4.3. Charge Transition Levels. Figure 9 presents the dependence of the formation energy ΔH on the position of the Fermi level ϵ_{F} . For V_{O} , the $q = 2+$ charge state is preferred in almost the entire range of ϵ_{F} . V_{O}^{2+} thermodynamically transitions to states with lower nominal charge q ($1+$ and 0) when the Fermi level is ≈ 0.1 eV below the CBM (location of the $(2+/1+)$ and $(2+/0)$ TTLs). Keeping in mind the uncertainties, we cannot say if V_{O}^{2+} transitions directly into V_{O}^0 (this would suggest a negative- U behavior, a similar observation was made in ref 120), or through a V_{O}^{1+} charge state. All three ΔH versus ϵ_{F} lines presented in Figure 9 cross at $\epsilon_{\text{F}} = 2.95$ eV.

For the Fermi level close to the CBM, the formation energies of all three charge states of V_{O} are very similar. This suggests that neutral, single-charged, and double-charged oxygen vacancies may coexist under the corresponding conditions.

The band gap $E_{\text{g}} = 3.06$ eV, as shown in Figure 9, corresponds to the result obtained for a $5 \times 5 \times 5$ supercell (the same system size was used for defect calculations). Calculations performed for a larger $7 \times 7 \times 7$ supercell gave a slightly lower band gap $E_{\text{g}} = 2.92$ eV (see Table 1, the main reason for this is the better sampling of reciprocal space), which is below the location of the transition levels observed for V_{O} . This would mean that for V_{O} in cubic STO, there are no TTLs in the band gap. A similar observation was made by Choi et al.⁸² based on DFT + U calculations.

Figure 9b shows that the hydrogen interstitial prefers a $q = 1+$ charge state over almost the entire range of ϵ_{F} . At $\epsilon_{\text{F}} = 2.97$ eV, it thermodynamically transitions to I_{H}^{1-} . In this case, it also cannot be concluded if a direct transition takes place, or if the intermediate $q = 0$ state is involved.

Similarly to V_{O} , all hydrogen interstitials have comparable formation energies for $\epsilon_{\text{F}} \approx E_{\text{g}}$. This suggests that they might coexist under the corresponding conditions.

Table 6 presents the calculated TTLs and OTLs (see eqs 3 and 4 for their definition), which are also schematically

Table 6. Thermodynamic and Optical Transition Levels^a

transition type (q/q')	charge transition level (eV)		
	thermodynamic $\epsilon_{\text{th}}(q/q')$	absorption $\epsilon_{\text{op}}(q'/q)$	emission $\epsilon_{\text{op}}(q/q')$
oxygen vacancy			
(2+/1+)	-0.11 (0.20)	-0.08 (0.08)	-0.09 (0.10)
(1+/0, $S = 0$)	-0.11 (0.13)	-0.98 (0.06)	-0.02 (0.04)
(1+/0, $S = 2$)	-0.12 (0.12)	-0.98 (0.06)	-0.02 (0.04)
(2+/0, $S = 0$)	-0.11 (0.05)	-1.09 (0.09)	-0.07 (0.09)
(2+/0, $S = 2$)	-0.12 (0.05)	-1.09 (0.09)	-0.08 (0.08)
hydrogen interstitial			
(1+/0)	-0.04 (0.20)	-0.11 (0.06)	-0.04 (0.06)
(0/1-, $S = 0$)	-0.12 (0.18)	-0.07 (0.08)	-0.02 (0.07)
(0/1-, $S = 2$)	-0.18 (0.14)	-0.08 (0.07)	-0.03 (0.06)
(1+/1-, $S = 0$)	-0.08 (0.12)	-0.08 (0.07)	-0.03 (0.06)
(1+/1-, $S = 2$)	-0.11 (0.10)	-0.07 (0.07)	-0.04 (0.06)

^aLocations of all levels are measured with respect to CBM. A negative value informs that the level is located below CB. The value given in brackets is the uncertainty. Note that the OTL presented in column 3 corresponds to the opposite transition $q' \rightarrow q$.

illustrated in Figure 10. V_O introduces both shallow and deep OTLs, which lie 0.08 eV ((1+/2+) level) and 1.0–1.1 eV ((0/

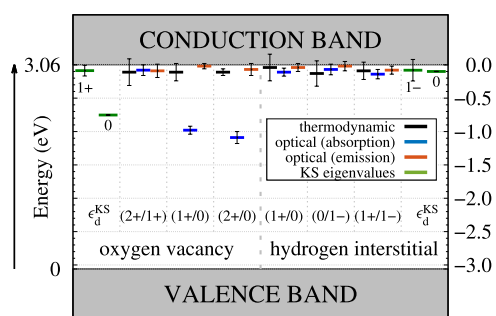


Figure 10. Thermodynamic $\epsilon_{th}(q/q')$ and optical $\epsilon_{op}(q/q')$ CTLs. The (q/q') labels describe the transition type. For absorption, the levels correspond to the opposite transition (q'/q) . The Kohn–Sham eigenvalues $\epsilon_d^{KS} + \Delta\epsilon_d^{KS}$ (corrected values) are also presented. Black error bars represent uncertainties. For V_O^0 and I_H^- no distinction is made between singlet and triplet states, for which results were similar.

1+) and (0/2+) levels) below the CBM when the excitations are considered. The OTLs related to the excitation of the hydrogen interstitial are shallow, located ≈ 0.1 eV below the CB edge.

For the neutral oxygen vacancy, significant differences are seen between OTLs related to excitation and those corresponding to recombination. While the (0/1+) and (0/2+) excitations of V_O^0 produce deep features (1.0–1.1 eV below CBM), the recombination of electrons from the CB to the states introduced by V_O^0 results in features that are very close to the CB edge (less than 0.1 eV below CBM, for all

recombinations to V_O^0). This observation explains the significant differences in the experimental studies concerning oxygen vacancies in STO.

Absorption techniques reported deep locations of defect levels associated with V_O . Ultraviolet photoemission spectroscopy measurements identified a feature located between 1.0¹⁴ and 1.2 eV¹⁵ below the CB, which agrees almost perfectly with our calculated (0/1+) and (0/2+) transition levels (1.0–1.1 eV below CBM). On the other hand, spectroscopy based on emission analysis reported shallow locations. For example, Kan et al.,¹⁶ based on photoluminescence (PL) measurements, observed a state which is located slightly (0.1 eV) below the CB. This agrees very well with our calculated (2+/1+), (1+/0), and (2+/0) OTLs. We note that recent PL studies also reported another level (located ≈ 0.4 eV below the CB). However, this feature was attributed^{16,18} to exciton recombination not considered in this work.

The calculated TTLs also agree well with conduction measurements of donor ionization energies, which all predicted a very shallow location (down to 0.16 eV below the CBM^{25,26}) of the TTLs of V_O .

Based on the above comparison, we conclude that the DFT + $U^{d,p}$ results obtained for V_O are in very close agreement with the experiment. A lack of experimental data prevents comparing the OTLs obtained for the hydrogen interstitials. However, the calculated TTLs are qualitatively consistent with recent experimental studies,^{32,34} which observed hydride conduction in STO subjected to reducing conditions and high temperatures. These experiments suggested that the (1+/1-) TTL of I_H is located very close to the CB edge, which agrees very well with our result.

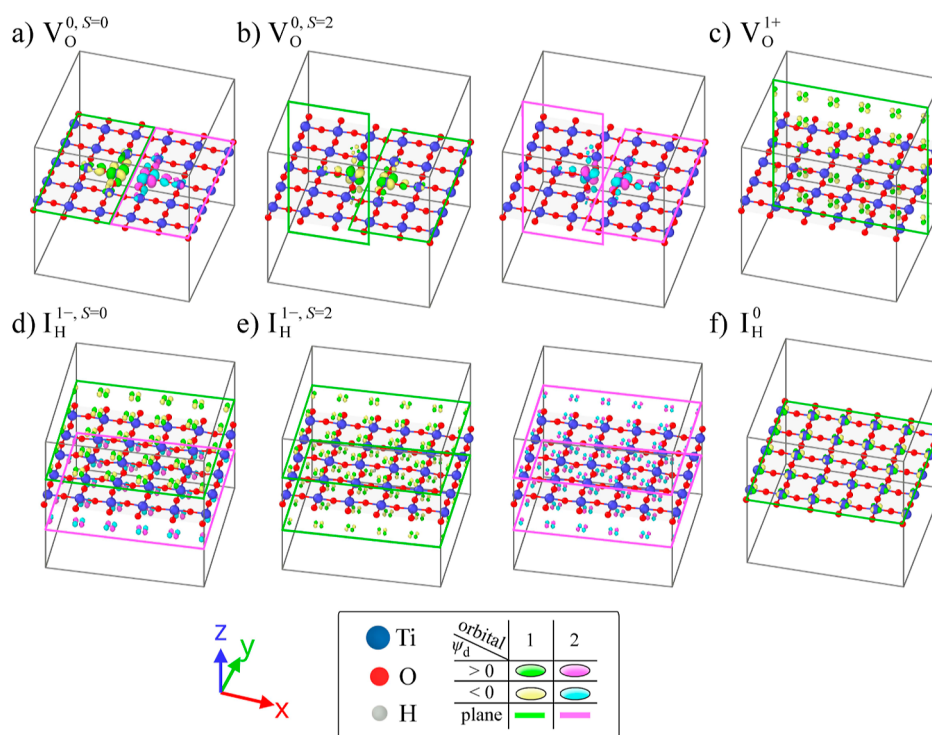


Figure 11. Levels introduced in the forbidden gap by the V_O (a–c) and I_H (d–f) defects. All panels show a cross-sectional view through the supercell (the central TiO_2 plane is shown, the defect is located in the center) and visualize isosurfaces corresponding to an electronic density of $|\psi_d^{KS}|^2 = 10^{-2} e/\text{\AA}^3$. In the interest of readability, planes containing lobes of ψ_d^{KS} were framed with color borders. For the triplet states ($V_O^{0,S=2}$ and $I_H^{1-,S=2}$), the two spin majority levels are presented separately.



For I_H , the obtained values of ϵ_d^{KS} (≈ 0.1 eV below CBM) correctly reflect the positioning of the TTLs and OTLs. For V_O^{1+} , the calculated ϵ_d^{KS} also agrees reasonably with the obtained TTLs and OTLs. However, the ϵ_d^{KS} eigenvalues obtained for V_O^0 (≈ 0.75 eV below CBM) differ strongly from the calculated TTLs (0.1 eV below CBM) and also do not reflect the positioning of the corresponding OTLs (1.0–1.1 eV below CBM for excitations of V_O^0 and 0.1 eV for recombinations to V_O^0). The obtained results show that—in general—the ϵ_d^{KS} Kohn–Sham eigenvalues are not accurate measures of the real locations of the defects' levels. Their correct estimation requires the calculation of the CTLs.

4.4. Defect Levels and Electronic Structure. The following analysis will be based on Kohn–Sham single-particle levels (ψ_i^{KS} wavefunctions) and the corresponding single-particle energies (ϵ_i^{KS} eigenvalues). Although both measures—as argued in the previous section—have some limitations, they are still helpful in the qualitative description of how the considered defects influence the electronic structure of STO. The presented analysis will also show why the Kohn–Sham eigenvalues ϵ_d^{KS} failed in describing the V_O^0 defect.

Figure 11a visualizes the two levels introduced in the forbidden gap by $V_O^{0,S=0}$. Each of these is localized on one neighboring Ti atom (Ti1 or Ti2), being mainly composed of its $3d-t_{2g}$ level (with a small addition of the O 2p levels of the nearby O atoms, see the DOS in Figure 12f). Because of this

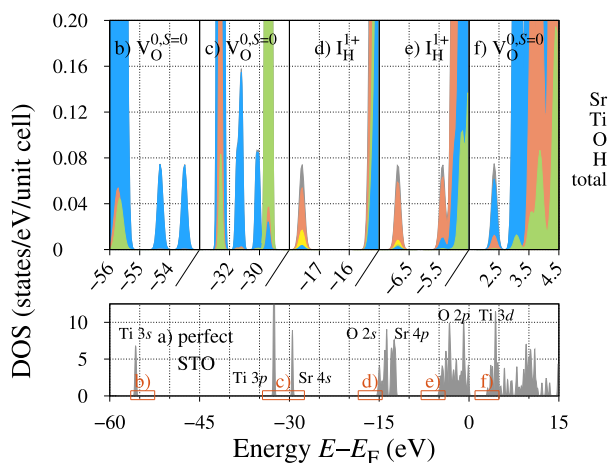


Figure 12. Influence of the V_O and I_H defects on the electronic structure of STO. Panel (a) (bottom) presents the total DOS of perfect STO and serves as a guide. It indicates the position of regions shown in panels (b–f), which present features introduced by $V_O^{0,S=0}$ (b,c,f) and I_H^{1+} (d,e) defects in the low-energy parts (b–e) and within the forbidden gap (f). Panels (b–f) present the local (atom-decomposed) DOS. All panels use the same reference energy (E_F of perfect STO).

trapping, the Ti1 and Ti2 atoms adopt a 3+ oxidation state. This change is visible in the DOS and manifests itself in a 1–2.5 eV increase of the Kohn–Sham eigenvalues corresponding to the 3s and 3p levels of the Ti1 and Ti2 atoms (see Figure 12, panels b and c).

Similar electronic behavior is observed for $V_O^{0,S=2}$, which also introduces two levels with the t_{2g} character in the forbidden gap. In this case, the decrease of the oxidation state is also visible in the lower energy parts of the DOS (not shown here). It also manifests itself in the outward displacements of the Ti1

and Ti2 atoms, which for $V_O^{0,S=2}$ (and also $V_O^{0,S=0}$) are by ≈ 0.2 Å lower than for the other charge states of V_O (see Table 3).

The levels introduced by $V_O^{0,S=2}$ (see Figure 11b) differ in symmetry from those introduced by $V_O^{0,S=0}$. For the triplet state, each level is composed of two orthogonal t_{2g} levels of the Ti1 and Ti2 atoms, such that the lobes of each level are located around two perpendicular planes. Two levels of $V_O^{0,S=2}$ differ in the sign of their single-particle wavefunctions.

The above analysis explains why the Kohn–Sham eigenvalues ϵ_d^{KS} are inadequate for predicting the CTLs of V_O^0 . During the charge transition from (and to) the neutral V_O , a change in the oxidation state of the Ti1 and Ti2 atoms manifests. This change is reflected in a considerable change of the Kohn–Sham eigenvalues of the deep levels (Ti 3s and 3p). Because the ϵ_d^{KS} eigenvalues obtained for V_O^0 do not account for this change in the low-energy eigenvalues, they cannot correctly describe the total energy change in the transitions to (and from) V_O^0 . From the above analysis, a general conclusion can be drawn: single (i.e., corresponding to one charge state) Kohn–Sham eigenvalues ϵ_d^{KS} are insufficient for predicting the locations of CTLs, which involve significant, that is, chemical, changes in the electronic structure.

In contrast to V_O^0 , V_O^{1+} introduces a shallow state, which is delocalized. This level is formed from the $3d-t_{2g}$ levels of multiple Ti atoms belonging to the same TiO_2 (010) plane (see Figure 11c). The participation of the 2p levels of the O atoms belonging to the same TiO_2 plane is also visible. For V_O^{1+} , there is no change in the oxidation state of the Ti1 and Ti2 atoms, as evidenced from the DOS (not shown here) and an analysis of the Ti1–Ti2 distance.

Similarly to V_O^{1+} , the levels introduced by I_H^0 and I_H^{1-} are shallow, delocalized, and also formed from the $3d-t_{2g}$ and 2p levels of the Ti and O atoms belonging to the same TiO_2 plane (see Figure 11, panels d–f).

V_O^{2+} does not introduce any levels in the forbidden gap and no other new features in the DOS. A similar situation (no levels within the forbidden gap) is observed for I_H^{1+} , which, however, manifests its presence in the lower energy parts of the DOS.

For all I_H defects, the addition of the H atom leads to new features appearing in the DOS below the O 2s and 2p valence bands (see Figure 12, panels d and e). These features are also composed of the H 1s level and should be attributed to the formation of an OH bond (and accompanying energy shifts). These new levels lie 2.31 eV below the O 2s band and 1.88 and 0.33 eV below the O 2p band. These locations are—within the obtained uncertainty ± 0.04 eV—independent of the charge state q . The given locations correspond to the Kohn–Sham eigenvalues and therefore should be considered only as estimates. A more accurate evaluation of the real positioning of these levels (energies which might be observed in the optical spectroscopy and which correspond to excitations from the identified OH levels) is not possible within the applied approach, requiring methods suitable for studying excited states.

The analysis of these (O 2s band- and O 2p band-derived, OH bond-related) states shows that they are predominantly formed by the states provided by the bonded O and H atoms. However, to a large degree they are also composed of the states provided by the nearest Ti atoms (see Figure 13). This reveals the importance of the Ti sublattices for protonic transport. With this in mind, future research could focus on verifying how the OH bonding could be affected by doping STO with other



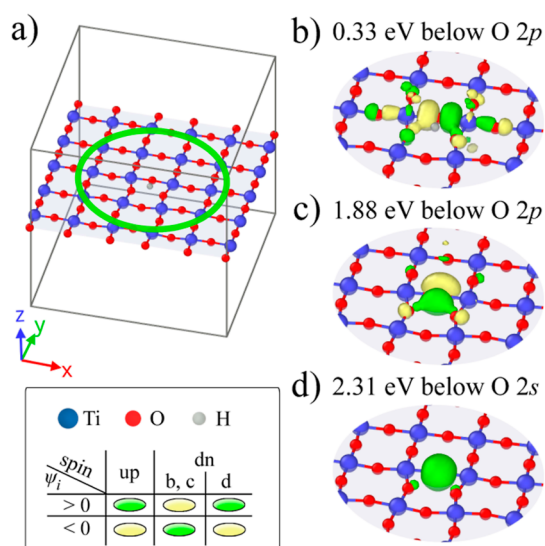


Figure 13. Influence of the I_H^+ defect on the electronic structure of STO. All panels present a cross-sectional view through the supercell, visualizing atoms from the central TiO_2 plane (with the H atom in the center). Panel (a) serves as a guide, showing the location of the region visualized in panels (b–d). These panels present the Kohn–Sham single-particle wavefunctions ψ_i^{KS} ($|\psi_i^{KS}|^2 = 10^{-2} \text{ e}/\text{\AA}^3$ isosurfaces are shown) of deep lying states, found below O 2p (b,c) and O 2s (d) bands. Only spin up ψ_i^{KS} are presented.

cations. Possibilities include enhancing proton transport by weakening the OH bond via electron drain to a more electronegative dopant.

The picture obtained in this work for V_O^0 disagrees (in many aspects) with previous theoretical studies. Their predictions are

schematically summarized in Figure 14 (scenarios a–d), which also presents the findings of this work (Figure 14e).

Standard DFT calculations and DFT + U studies, which used low U values,^{80,119–121} predicted that V_O^0 introduces two delocalized t_{2g} levels, which are positioned close to or in the CB (scenario a). This would mean that the $q = 0$ charge state is unstable. This failure can be attributed to a significant underestimation of the STO band gap. Hybrid DFT calculations^{53,56,57,122,123,125,126} predicted deep levels for V_O^0 but found that they have a Ti 3d– e_g character. DFT + U studies that used larger U values (yielding a reasonable band gap) predicted deep localized levels with an e_g character^{57,80–82,130} (scenario c, obtained for the singlet state) or positioned one of the defect levels in the CB^{81,82} (scenario d, obtained for the triplet state). Our results also disagree with those obtained recently based on DFT + U + V calculations by Ricca et al.⁹³ They also found that $V_O^{0,S=0}$ introduces two deep localized levels but with an e_g character (scenario c). For $V_O^{0,S=2}$, two distinct levels were identified in ref 93: one deep, localized level (with an e_g character) and one delocalized level (with t_{2g} character) located within the CB (scenario d). This also disagrees with our result.

To the best of our knowledge, none of the theoretical studies carried out to date reported defect levels with characteristics similar to that found in this work. We do not set out to identify the reason or reasons for the observed discrepancies (like differences in the supercell size, accounting for structural relaxation, the type and the quality of the basis set used, the XC functional chosen, the details related to the Hubbard + U and Hund's + J corrections, and others), only noting that some efforts toward establishing these were undertaken in ref 93. Instead, we emphasize that the results obtained in this work are supported by the agreement of the calculated CTLs with those

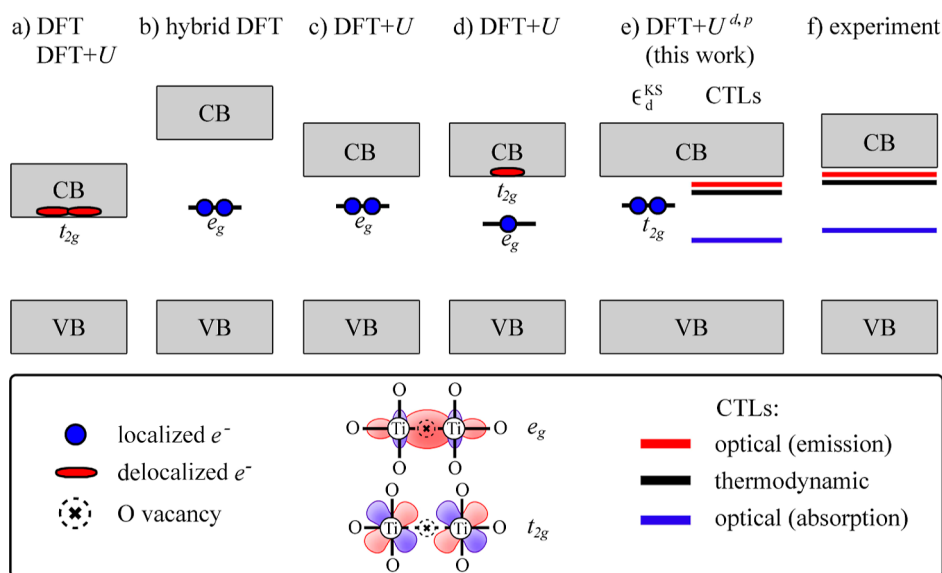


Figure 14. Defect levels introduced by the neutral oxygen vacancy V_O^0 in the forbidden gap, as predicted in the literature (a–d). Scenario (e) corresponds to the picture obtained in this work, in which the Kohn–Sham eigenvalues (ϵ_d^{KS}) still—as in previous works—disagree with the experiment (f). However, the calculated CTLs agree well with the available experimental data and explain the differences between the spectroscopic and conduction measurements done for V_O^0 . The presented schematic was partly (scenarios a–d) redrawn after ref 93 (Figure 2 therein): C. Ricca, I. Timrov, M. Cococcioni, N. Marzari, U. Aschauer, Self-consistent DFT + U + V study of oxygen vacancies in $SrTiO_3$, Phys. Rev. Res. 2, 023313, 2020, DOI: <https://doi.org/10.1103/PhysRevResearch.2.023313>; licensed under the Creative Commons Attribution 4.0 International license. (a) References 80 and 119–121, (b) refs 53, 56, 57, 122, 123, 125, and 126, (c) refs 57, 80–82, 93, and 130, (d) refs 81, 82, and 93, (f) refs 14–16, 25, and 26.

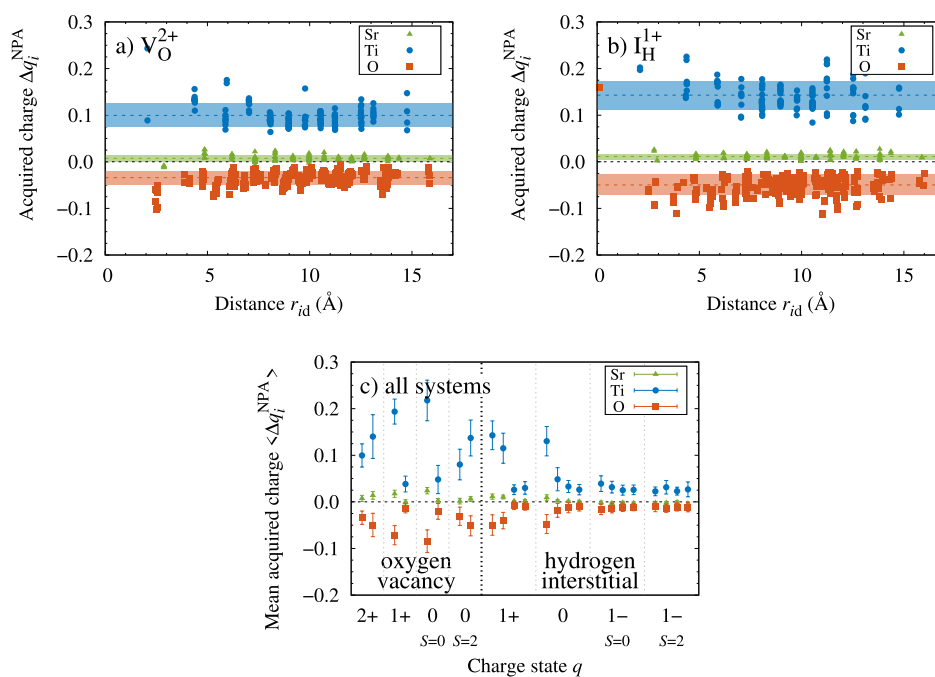


Figure 15. NPA carried out for the V_O^{2+} (a) and I_H^{1+} (b) defects, and all considered systems (c). Panels (a,b) show how the charges acquired by individual atoms (Δq_i^{NPA}) depend on their separation from the defect (r_{id}). Here, each point represents a single atom, dashed lines represent mean acquired charges of Sr, Ti, and O atoms, and the shaded regions depict the corresponding standard deviations. Panel (c) summarizes results obtained for all identified minima (the ordering of minima is the same as in Tables 3 and 4 if read from left to right). Here, the points represent the mean acquired charges of Sr, Ti, and O atoms, and the error bars depict standard deviations [corresponding to the regions highlighted in panels (a,b)].

measured experimentally^{14–16,25,26} (compare scenario e—our work—with scenario f—experiment).

4.5. Atomic Charges. We assessed the influence of the V_O and I_H defects on the bonding of STO using natural population analysis (NPA).^{179,180} The NPA atomic charges calculated for the perfect system are +1.75 for Sr, +2.12 for Ti, and −1.31 and −1.25 for equatorial and apical O atoms, respectively. The atomic charge of Sr is very close to its oxidation state (2+), showing the ionic character of its bonding. In contrast to this, the atomic charges on Ti and O differ strongly from their oxidation states (4+ and 2−, respectively), revealing a degree of covalency in their bonding.

The atomic charges obtained for the perfect system were used as reference charges in the analysis of charge redistribution in the defective systems. For this purpose, we calculated the acquired charges, that is, the differences

$$\Delta q_i^{NPA} = q_i^{NPA}(D, q) - q_i^{NPA}(P) \quad (11)$$

of the NPA atomic charges corresponding to the defective (D, q) and perfect (P) systems, with i indexing atoms. All the studied defective systems were analyzed in this manner.

Figure 15 shows the dependence of the acquired charge Δq_i^{NPA} on the distance from the defect $r_{id} = |\mathbf{R}_i - \mathbf{r}_d|$ (\mathbf{R}_i denotes the position of atom i and \mathbf{r}_d represents the position of the defect). Results for two example systems are presented: the lowest energy structures of V_O^{2+} and I_H^{1+} . Polarization of the crystal structure is apparent. The introduction of either defect causes the Ti and O atoms to acquire additional charge—mean values (averaged over all Ti/O atoms) are given: +0.10 for Ti and −0.034 for O (for V_O^{2+}) and +0.14 for Ti and −0.050 for O (for I_H^{1+}). The Sr atoms, in contrast, acquire only negligible charge.

Although the observed polarization is modest (for Ti and O atoms the mean acquired charges constitute 5–7% and 2–4% of their reference charges, respectively), it extends over the entire system and—in the studied systems—does not diminish appreciably even at the largest considered distances r_{id} (limited by the size of the employed supercell). Our calculations do not set out to determine how localized this effect is.

The $\Delta q_i^{NPA}(r_{id})$ characteristics obtained for the other considered systems (other charge states and other minima) are qualitatively similar to those presented. However, some quantitative differences were identified. These are shown in Figure 15c, which summarizes all the population analyses we carried out.

The polarization identified above occurs for all systems and is manifested through the charging of the Ti and O atoms but not Sr atoms. The fact that the polarization is observed for the completely ionized (V_O^{2+} and I_H^{1+}) defects means that it is not related to the redistribution of electrons from the defect levels. It is also not a consequence of a non-zero defect charge because it is also observed for V_O^0 and I_H^0 . These two observations lead us to attributing the observed polarization to the removal/addition of the defect's atom.

As seen from Figure 15c, the magnitude of the polarization varies between systems and can even be different for systems containing the same type of defect (compare results obtained for various minima of V_O^{1+} , $V_O^{0,S=0}$, I_H^{1+} , and I_H^0). This is illustrated in Tables 7 and 8, where we present the atomic charges of the atoms close to the defect. These are the two nearby Ti atoms (Ti1 and Ti2), the four closest Sr atoms, and the 10 closest O atoms (neighbors of Ti1 and Ti2), as well as (only for I_H defects) the O and H atoms forming the OH group.

Table 7. Atomic Charges for Atoms Close to the V_O Defect. Data Corresponding to Various Charge/Spin States (First Column) and Various Identified Minima (Consecutive Rows) Is Presented^a

defect state	acquired charge Δq_i^{NPA}			
	Ti1	Ti2	O (10 atoms)	Sr (4 atoms)
V_O^{2+}	+0.089	+0.243	-0.069 (0.024)	-0.010 (0.002)
	+0.092	+0.253	-0.067 (0.032)	-0.003 (0.002)
V_O^{1+}	+0.163	+0.288	-0.088 (0.029)	-0.005 (0.004)
	+0.012	+0.132	-0.034 (0.023)	-0.021 (0.002)
$V_O^{0,S=0}$	-0.077	+0.001	-0.145 (0.012)	+0.012 (0.001)
	-0.187	-0.065	-0.099 (0.026)	-0.008 (0.002)
$V_O^{0,S=2}$	-0.125	-0.056	-0.115 (0.022)	-0.015 (0.002)
	-0.130	-0.023	-0.127 (0.018)	-0.008 (0.002)

^aFor the O and Sr atoms, the mean value (averaged over the specified number of atoms) and its standard deviation (in brackets) are given. They specify the charge acquired by a single Sr or O atom. The ordering of the minima is the same as in Table 3.

The most significant differences are observed for V_O^{1+} , $V_O^{0,S=0}$, and I_H^{1+} defects, where the charge acquired by Ti1 and Ti2 atoms differs (between the identified minima) by as much as 0.15 (V_O^{1+}), 0.11 ($V_O^{0,S=0}$), and 0.14 (I_H^{1+}). They are also visible in the charges acquired by the nearest O atoms, where the total (summed) atomic charge may differ even by 0.54 (V_O^{1+}), 0.46 ($V_O^{0,S=0}$), and 0.51 (I_H^{1+}). The strongly differing minima identified here correspond to those which visibly differ in the mean acquired charges (cf. Figure 15c). This reveals the connection between the polarization observed in the very vicinity of the defect and the entire structure.

The charges of the H and O atoms from the OH group (see Table 8) are also—to some degree—sensitive to the atomic positions (they differ slightly between the identified minima). By itself, the hydrogen atom carries a charge, which ranges from +0.34 to +0.44. The remaining charge of the I_H defect can be either localized or delocalized.

The localized charge is also accommodated on the nearby O atom (from the OH group), whose atomic charge increases by

0.08–0.16 after introducing I_H . The delocalized charge is distributed in the entire supercell, as either a polarization charge (see Figure 15) or in the form of delocalized defect levels (for I_H^0 and I_H^{1-} , see Figure 11).

Population analysis reveals that the I_H defect carries a localized +0.5 e charge (here, Δq_i^{NPA} of the O atom and q_i^{NPA} of the H atom were summed), which seems to be independent of the formal charge q of the I_H defect. A similar conclusion was made by Bork et al.,¹³⁴ who, by combining DFT calculations with Bader charge analysis, found the same (+0.5) charge for I_H^{1+} and I_H^0 .

4.6. Crystalline Structure. The influence of the defects on the crystalline structure was quantified in terms of interatomic distances, with the aid of the pair correlation function (PCF), defined as

$$g(r) = \frac{N(r)}{4\pi r^2 \Delta r} \frac{V}{N} \quad (12)$$

Here, $N(r)$ represents the number of pairs of atoms (of any type) that are located within the distance $[r, r + \Delta r)$, while V and N stand for the volume and the total number of atoms in the system, respectively. The PCF relates the local atomic number density at a distance r from any atom to the average atomic number density of the entire system, N/V .

In Figure 16, we present the total PCFs of all considered defects and decompositions of PCFs obtained for V_O^{2+} and I_H^{1+} into partial functions, corresponding to various pairs of atom types.

The similarity of all the presented PCFs shows that in the considered short and medium ranges, the V_O and I_H defects have a similar effect on the crystalline structure of STO. The degree of this influence is comparable among various charge/spin states.

As already mentioned in Section 4.1 (see also Figure 8), the introduction of defects mainly leads to rotations and distortions of the TiO_6 octahedra. Changes of another character, like the displacements of the Sr and Ti ions, are only visible in the very vicinity of the defect. This picture allows explaining the obtained PCFs.

Table 8. Atomic Charges for Atoms Close to the I_H Defect^a

defect state	acquired charge Δq_i^{NPA}				atomic charge q_i^{NPA}	
	Ti1	Ti2	O (10 atoms)	Sr (4 atoms)	O	H
I_H^{1+}	+0.197	+0.202	-0.059 (0.018)	+0.013 (0.011)	-1.152	+0.340
	+0.142	+0.141	-0.044 (0.014)	+0.006 (0.012)	-1.207	+0.403
	+0.061	+0.056	-0.008 (0.015)	+0.006 (0.012)	-1.209	+0.427
	+0.175	+0.181	-0.050 (0.016)	+0.019 (0.009)	-1.218	+0.408
I_H^0	+0.071	+0.066	-0.008 (0.016)	+0.002 (0.006)	-1.197	+0.419
	+0.057	+0.075	-0.014 (0.017)	+0.004 (0.008)	-1.230	+0.437
	+0.037	+0.036	-0.010 (0.019)	+0.002 (0.005)	-1.207	+0.429
	+0.053	+0.058	-0.009 (0.017)	-0.001 (0.013)	-1.199	+0.416
$I_H^{1-,S=0}$	+0.076	+0.126	-0.020 (0.011)	-0.005 (0.010)	-1.132	+0.343
	+0.083	+0.053	-0.014 (0.009)	-0.008 (0.011)	-1.140	+0.362
	+0.015	+0.015	+0.001 (0.019)	-0.004 (0.009)	-1.197	+0.417
	+0.012	+0.027	-0.002 (0.015)	-0.006 (0.009)	-1.217	+0.437
$I_H^{1-,S=2}$	+0.016	+0.029	-0.002 (0.017)	-0.007 (0.010)	-1.132	+0.366
	+0.048	+0.084	-0.016 (0.013)	-0.005 (0.010)	-1.192	+0.393
	+0.003	+0.008	+0.001 (0.017)	-0.006 (0.009)	-1.188	+0.409
	+0.070	+0.104	-0.020 (0.013)	-0.006 (0.010)	-1.155	+0.375

^aThe meaning of columns 1–5 is the same as in Table 7. The last two columns present the atomic charges of the O and H atoms from the OH group. The ordering of the minima is the same as in Table 4.

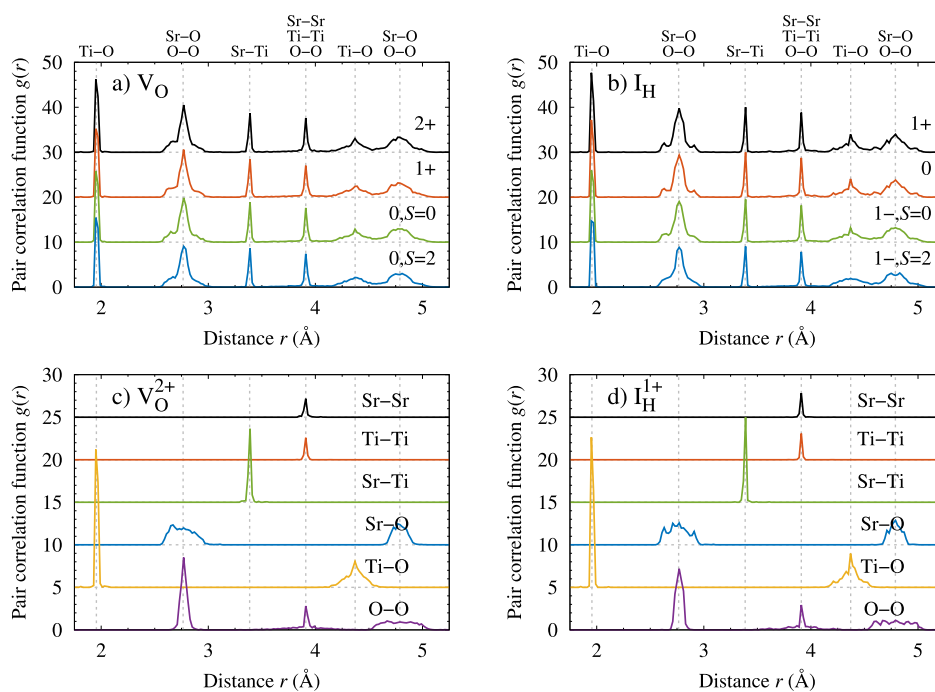


Figure 16. Influence of the V_O and I_H defects on the short- and medium-range order of STO. Panels (a,b) compare the total PCFs $g(r)$ obtained for various charge/spin states of V_O (a) and I_H (b). Panels (c,d) present the decompositions of the total $g(r)$ obtained for V_O^{2+} and I_H^{1+} into partial PCFs, corresponding to various pairs of atom types. The vertical dashed lines correspond to distances in the perfect system (described at the top). Note that all panels use vertical offsets. Their values can be directly read from the labeled tick-marks.

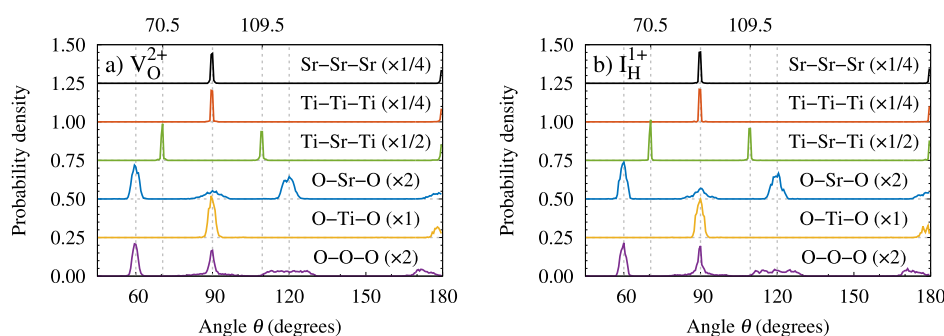


Figure 17. Influence of the V_O and I_H defects on the short- and medium-range order of STO (cont'd). The two panels present distributions of various valence angles (described within the plots) for V_O^{2+} (a) and I_H^{1+} (b) defects. Distributions were normalized to values given in the round brackets. The vertical dashed lines correspond to angles in the perfect system. Note that all panels use vertical offsets. Their values can be directly read from the labeled tick-marks.

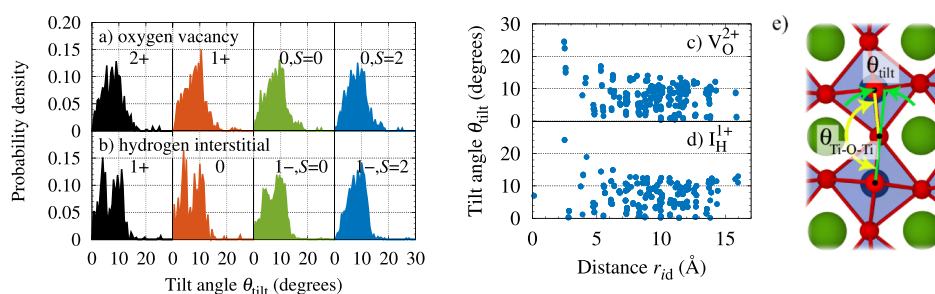


Figure 18. Defect-induced octahedral tilting in STO. Panels (a,b) present distributions of the tilt angle θ_{tilt} calculated for all considered defects (labeled in the plot). Panels (c,d) show the dependence of the tilt angle on the distance from the defect for two exemplary systems (minimum energy structures of V_O^{2+} and I_H^{1+}). Here, each point represents a single O atom, presenting its distance from the defect and the value of the tilt angle θ_{tilt} centered on it. Panel (e) illustrates the meaning of θ_{tilt} .

The rotations of the TiO_6 octahedra cause a significant change in the O–O, Ti–O, and Sr–O distances. These changes are well visible in all PCFs and manifest through the

broadening of the corresponding peaks: second (O–O and Sr–O distances), fourth (O–O distances), fifth (Ti–O distances), and sixth (O–O and Sr–O distances). In contrast,

no significant broadening (or shift) of the first peak is observed. This shows that the shortest (intra-octahedral) Ti–O distance remained constant, on average, in defect-containing systems.

The sharp fourth peak (corresponding to the shortest Sr–Sr and Ti–Ti distances) indicates no significant changes in either the Sr or the Ti sublattice. Their relative displacement is also not observed (sharp third peak, Sr–Ti distances).

Figure 17 reveals that in the defect containing systems the valence angles $\theta_{\text{Sr-Sr-Sr}}$ and $\theta_{\text{Ti-Ti-Ti}}$ (formed by a triple of the nearest Sr- or Ti-atoms) are close to 90 and 180°. This shows that the Sr and Ti sublattices remained regular. The $\theta_{\text{Ti-Sr-Ti}}$ angles also are similar to those seen in the perfect system, demonstrating no relative displacement of the Sr and Ti sublattices (except for the very vicinity of the defect).

The distributions of all valence angles incorporating O atoms display significant broadening, up to $\pm 10^\circ$. This, again, shows the rotations (changes visible in the $\theta_{\text{O-Sr-O}}$ angles) and distortions ($\theta_{\text{O-Ti-O}}$ and $\theta_{\text{O-O-O}}$ angles) of the TiO_6 octahedra.

There is one more valence angle, which is especially important in the context of proton conductivity, namely, the $\theta_{\text{Ti-O-Ti}}$ angle. As shown in Figure 18e, this angle (spanned by two Ti atoms belonging to two neighboring octahedra, and the O atom being their common neighbor) can be used to measure the relative tilting of the TiO_6 octahedra, with the tilt angle defined as

$$\theta_{\text{tilt}} = 180^\circ - \theta_{\text{Ti-O-Ti}} \quad (13)$$

Its importance originates from the fact that the octahedral tilting is suspected to reduce the activation energy for proton migration, enhancing proton conductivity.^{181,182}

Figure 18 demonstrates that the introduction of V_O and I_H defects alike results in a well pronounced octahedral tilting of $\approx 7\text{--}8^\circ$ on average (a similar mean was found for all systems), which may reach even $20\text{--}30^\circ$ close to the defect. This tilting effect does not decay with the separation from the defect. However, it can be suspected that the limited size of the simulated systems (≈ 20 Å supercells), as well as the use of PBCs, may have some influence on this non-vanishing behavior. The similarity of the $\theta_{\text{tilt}}(r_{\text{id}})$ and $\Delta q_i^{\text{NPA}}(r_{\text{id}})$ (see Figure 15) dependencies shows that the octahedral tilting and the observed polarization of the TiO_6 octahedra are related.

5. CONCLUSIONS

5.1. Summary. This work investigated oxygen vacancy and hydrogen interstitial defects in cubic STO. A new approach to describing this difficult system was developed within the framework of linear-scaling Kohn–Sham DFT. The band gap problem was overcome by applying Hubbard + U and Hund's + J corrections to both Ti (3d subshell) and O (2p subshell) atoms, following the DFT + U^{dp} method recently proposed by O'Regan et al.⁹⁵

The U and J parameters were determined on theoretical grounds, using the MT-LR approach⁶⁹ and later validated for perfect STO. DFT + U^{dp} significantly improved the description of STO's electronic structure close to the Fermi energy, as well as its mechanical properties, both of which are essential for the accurate modeling of defects.

The validated methodology was applied to the study of oxygen vacancy (V_O) and hydrogen interstitial (I_H) defects. Various charge and spin states of these defects were considered

and modeled using large supercells, containing $N = 625 \pm 1$ atoms. The relaxed configurations of defects were obtained through fastidious energy minimization, which consisted of a series of continued minimizations, carried out alternately in the delocalized internal and Cartesian coordinates.

Reproducibility tests showed that the potential energy surface of STO contains multiple local minima, which differ in the atomic positions and the observed internal polarization, but have comparable energies and correspond to similar static (i.e., not related to the ion transport) properties of defects.

For charged defects, the calculated total energies were corrected using a method proposed recently by Durrant et al.¹¹⁴ It was found that for STO (which has a high dielectric constant) and for large supercells, the image interaction and PA corrections are small, not exceeding 0.01 and 0.15 eV, respectively.

The obtained relaxed configurations of defects were analyzed from a number of perspectives. The calculated formation energies showed that the oxygen vacancy and hydrogen interstitial strongly prefer a fully ionized state, with a 2+ and 1+ formal charge, respectively. The predicted formation energy of V_O^{2+} was in close agreement with experimental oxidation and reduction enthalpies.

The locations of defect levels were determined using the CTL formalism.⁹⁹ With the exception of the neutral V_O , all considered defects introduced states that would be seen as shallow in both conductivity and optical measurements being located down to 0.2 eV below the CB. The neutral oxygen vacancy introduced shallow (thermodynamic, optical for the emission) and deep (optical for the absorption) CTLs. This observation explains the inconsistencies seen in experimental characterizations of V_O .

It was revealed that, in general, the locations of defect levels cannot be reliably predicted based on the Kohn–Sham eigenvalues corresponding to a single charge state, as these were unable to reflect the positioning of the defect levels of V_O . The underlying reason was the change in the oxidation state of two nearby Ti atoms.

The V_O^{1+} , I_H^0 , and I_H^{1-} defects introduced shallow and delocalized states in the forbidden gap, which were composed of the 3d- t_{2g} and 2p states of the Ti and O atoms belonging to the same TiO_2 plane. The deep-lying states of the V_O^0 defect were composed of the 3d- t_{2g} states of two nearby Ti atoms, which is in contrast to previous theoretical reports that predicted a Ti 3d- e_g character.

Both V_O and I_H defects manifested their presence also in the lower parts of the DOS. The neutral oxygen vacancy considerably (by 1.0–2.5 eV) shifted the 3s and 3p levels of two nearby Ti atoms. All hydrogen interstitials introduced states located below the O 2s and O 2p bands, which can be attributed to the formation of the OH bond.

Analysis revealed that V_O and I_H defects have a similar influence on the bonding and crystalline structure of STO, causing polarization and tilting of the TiO_6 octahedra. Both effects occurred due to the defect's atom removal/addition. Our calculations did not answer how localized these effects are. The Sr sublattice was found to be insensitive to the presence of the V_O and I_H defects.

5.2. Discussion and Future Work. This work demonstrated the DFT + U^{dp} approach to be successful for describing STO, continuing along the direction set by O'Regan et al.⁹⁵ (and also other authors^{101,152,153}), who demonstrated its

adequacy for describing other difficult transition-metal oxides, like TiO_2 , CuO , and CeO_2 .

The calculations we carried out yield a picture that is highly consistent with the experiment. We attribute this success to: (i) an adequate formal description (theoretically motivated, free of empirical parameters, and validated for the perfect system), (ii) high quality of the calculations (near-complete basis set accuracy, accounting for the semi-core states, and a tailored procedure for carrying out geometry relaxations), and (iii) physically sound calculations and analysis (large super-cells, careful accounting for structural relaxations, diligent treatment of electrostatic corrections, and explicit consideration of various charge states and calculation of the CTLs).

The adopted methodology can be applied almost straightforwardly to other perovskite materials. Here, we underline the importance of the MT-LR method, which provides a unified framework for finding the essential (correction defining) U and J parameters for various elements, even those that display magnetization.⁶⁹

The MT-LR method relies on the perturbation of individual sites and, therefore, allows determining the U and J parameters individually for each atom chosen for correcting. This opens the possibility of studying dopants (like Fe substitutes at the Ti sites in $\text{SrTi}_{1-x}\text{Fe}_x\text{O}_{3-\delta}$) and better accounting for chemical changes, which may occur due to the introduction of defects or their migration. The MT-LR approach also enables the application of the $+U$ and $+J$ corrections in a self-consistent fashion (similarly as it was done in ref 93). We did not account for the potential site dependence of the U and J parameters in the present study. Therefore, in future work, it would be prudent to check, for example, how the predicted CTLs of V_O^0 change if the changes in the oxidation state of two nearby Ti atoms are explicitly taken into account (e.g., by reevaluating their U and J parameters).

This work used the ONETEP program, which provides a formalism for linear-scaling DFT calculations. However, linear scaling was not exploited here, as we did not (for now) truncate the density kernel as we wanted to ensure that the results are not affected by additional approximations. In the next step, it will be essential to verify if the quality of the results can be maintained in the presence of truncation (the large band gap of STO suggests such possibility), also checking how other numerical parameters (like the localization radius of the NGWFs) influence the theoretical predictions. In this respect, the obtained results serve as a trustful checkpoint, allowing to test what further approximations can be made to reduce the computational cost. Its significant reduction will enable studying larger systems and efficiently investigating the ionic transport through transition state searches and even Born–Oppenheimer molecular dynamics simulations.

The obtained results suggest that defected STO (and, perhaps, other perovskites) might be characterized by a potential energy surface with multiple local minima, all with a similar total energy but with markedly different spatial distributions of the charge. These differences—which manifest in the observed polarization—may prove to be significant in the context of ion transport. It should be expected that the paths for O and H ion migration (and the corresponding energy barriers) depend on the atomic charges seen in the immediate vicinity of the defect and are strongly felt by the migrating ion. Our results show that these charges may be different for the minima of the same defect type. This reveals the necessity to verify how reproducible the minimum energy

paths are (and the corresponding barrier heights) calculated based on the theory using, for example, the nudged elastic band method (see e.g., refs 72, 73, 134, 183, and 184).

The obtained picture of how the defects influence the structure and bonding of STO also requires a discussion within the context of protonic transport. According to the commonly accepted view, in perovskites, it occurs through the so-called Grotthuss mechanism,^{185–187} which consists of two basic events: proton rotations (around the O atoms that they are bonded to) and proton hopping (to another—neighboring—O atom). It is also believed that proton rotations are energetically less expensive than proton hopping.

Our calculations suggest that the I_H defect causes a notable polarization and tilting of the TiO_6 octahedra. Let us—within this context—consider a scenario, in which the migrating proton moves from one equilibrium position to another (the exact type of the migration event is not relevant here). The associated polarization and octahedral tilting can be expected not to follow the proton immediately. Thus, after the migration event (rotation or hopping), the crystal lattice has locally higher energy, temporarily displaying non-equilibrium polarization and tilting. We note that the above reasoning can be motivated based on purely inertial and configurational grounds: rotations and distortions of many octahedra consisting of much heavier (than the proton) O ions are required for the crystal lattice to reach its new equilibrium state.

With the above in mind, in perovskite materials, protonic transport might be characterized by more than two energy scales (barrier heights for the rotation and jump) because, in the discussed picture, the knowledge of the lattice excitation energies is also required. This suggests that theoretical characterizations of the ion transport, which considered only the migration between two minima of the same energy, might be incomplete.

In this work, we do not attempt to answer the questions which naturally arise from the above discussion, like: “what are the heights of the energy barriers and energies of the related lattice excitations?”, “is the proton rotation/jump to its previous position more likely?”, “what are the related probabilities?”, “how long does it take for the crystal lattice to reach new equilibrium after the proton migration event?”, “what influence does the temperature have on all of these?”. Here, we only draw the scientific community’s attention, noting that the related problems are the topic of our ongoing research.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.jpcc.2c04681>.

Detailed information related to the choice of U and J parameters, choice of the reference chemical potentials, calculation of the electrostatic corrections, and determination of the optical transition levels (PDF)

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Notes

The authors declare no competing financial interest.

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