Exchange–correlation effects in the monoclinic to tetragonal phase stabilization of Yttrium–doped $ZrO_2$: a first–principles approach

Davide Sangalli$^{1,2}$ and Alberto Debernardi$^1$

$^1$MDM Lab - IMM - CNR via C. Olivetti, 2 I-20864 Agrate Brianza (MB) Italy, European Union
$^2$Consorzio Nazionale Interuniversitario per le Scienze dei Materiali (CNISM)

(Dated: December 16, 2011)

Abstract

We describe, within an ab–initio approach, the stabilization of the tetragonal phase vs. the monoclinic one in Yttrium–doped Zirconia. The process is believed to be influenced from different mechanisms. Indeed we show that there is a delicate balance between the change in electrostatic and kinetic energy and exchange–correlation effects. In the tetragonal phase the perturbation induced by doping is better screened at the price of sacrificing correlation energy. Our work opens the opportunity to use the same approach to predict the tetragonal phase stabilization of materials like Zirconia or Hafnia, with different and less characterized dopants.

PACS numbers: 64.70.K-, 71.15.Mb, 81.05.Je, 81.30.Bx
I. INTRODUCTION

Zirconia (ZrO$_2$) is a hard and usually colorless material with a wide range of technological applications\textsuperscript{1}. Being corrosion resistant it is used as a dental material and because of its low cost, durability, and close visual likeness to diamond, it is widely used to synthesize artificial gems. It is used as a thermal barrier in coating engines due to its high resistance. The addition of cations (as for example $Y^{3+}$) induces the generation of oxygen vacancies for charge compensation which makes it useful as oxygen sensor. Moreover Zirconia is an high dielectric constant (high-$\kappa$) material with potential applications in the micro–electronics. Finally, very recently, transition–metals doped Zirconia has been predicted to be a dilute magnetic semiconductor (DMS) with high Curie Temperature $T_c$ with potential applications in the field of spintronics\textsuperscript{2}; while pure and doped ZrO$_2$ has been proposed as a candidate material for resistive switching memories devices\textsuperscript{3,4} (ReRAM) exploiting the high vacancies mobility of the system.

Pure ZrO$_2$ exhibits at ambient pressure three polymorphisms. The monoclinic (M) phase is stable at low temperature and is the less symmetric structure with the Zr$^{4+}$ ions exhibiting sevenfold coordination. Between 1440 and 2640 K the tetragonal (T) phase, with eightfold coordinated Zr$^{4+}$ ions is stable. Finally above 2640 K, till melting temperature ($\approx$ 3000K), the most symmetric cubic (C) phase is stabilized\textsuperscript{5,6}. The only difference between the (C) and the (T) phase is a distortion of the oxygen sub–lattice with a spontaneous symmetry breaking. The (T) and (C) phases are more used in technological applications and both the (T)/(C) structures and the (T) $\rightarrow$ (C) doping induced phase transition (DIPT) have been well characterized in the literature both form the theoretical and the experimental point of view\textsuperscript{5,7–16}, with $Y^{3+}$ ions the most used and studied dopants.

On the other hand the (M) $\rightarrow$ (T) DIPT is much less characterized, especially from the theoretical point of view. It presents a volume change of about 3 – 4% that causes extensive cracking in the material. This behavior destroys the mechanical properties of fabricated components and makes pure Zirconia useless for structural or mechanical application. Moreover the (T) phase, is metastable in pure and lightly doped ZrO$_2$ over a very long time; a growth sample of doped Zirconia must be annealed in order to check if the reached tetragonal phase is stable or meta–stable. Hence a better understanding of the (M) $\rightarrow$ (T) DIPT is desirable. In this paper we address the problem from a first principle perspective and,
Table I. Theoretical and experimental cell parameters of \(\text{ZrO}_2\) in the monoclinic and in the tetragonal phase.

|                  | Present work | Theory | Experiment |
|------------------|--------------|--------|------------|
| \(\text{(M) ZrO}_2\) |              |        |            |
| \(a\) (Å)        | 5.18         | 5.15   | 5.15       |
| \(b/a\)          | 1.011        | 1.012  | 1.012      |
| \(c/a\)          | 1.037        | 1.028  | 1.032      |
| \(\beta\)        | 99°10'       | 99°14' | 99°14'     |
| \(\text{(T) ZrO}_2\) |              |        |            |
| \(a\) (Å)        | 5.11         | 5.02   | 5.03       |
| \(c/a\)          | 1.030        | 1.014  | 1.017      |

in particular, we show that the DIPT is a balance of the mean field (MF) description (i.e. kinetic plus electrostatic: \(H = T + V_{\text{ext}} + V_{\text{H}[\rho]}\)) with exchange–correlation effects which cannot be captured by simplified models.

II. FIRST PRINCIPLES DESCRIPTION.

A. Computational details.

We work within density functional theory\textsuperscript{19,20} (DFT) in the generalized gradient approximation\textsuperscript{21} (GGA) with ultrasoft pseudopotentials\textsuperscript{22,23}. Both for Yttrium and Zirconium the pseudopotentials includes semicore electrons and non–local core correction. The Quantum Espresso\textsuperscript{24} package is used to solve the Kohn–Sham\textsuperscript{20} (KS) equations for a super–cell with 96 atoms (down to 92 when oxygen vacancies are considered). We checked that a cut–off of 35 Ry for the wave–functions and 400 Ry for the augmentation charge, a k-point grid \(2 \times 2 \times 2\) are needed in order to have the energy difference between the tetragonal and the monoclinic phase converged to up to \(5.0 \times 10^{-4}\) eV per \(\text{ZrO}_2\) molecular unit (m.u.), where the energy difference is of the order of 0.1 - 0.01 eV/m.u. .

As starting point we relaxed atomic positions and structure of pure \(\text{ZrO}_2\) with 12 atoms in the unit cell of the (T) and (M) phases. The (C) phase is unstable and is not considered.
Theoretical and experimental internal structural parameters of ZrO$_2$ in the Wyckoff notation. For the tetragonal structure the only free parameter is the z coordinate of the oxygen atoms (0.0, 0.5, 0.25 – $d_z$).

| Present work | Theory | Experiment |
|--------------|--------|------------|
| (M) ZrO$_2$  | Ref.[8] | Ref.[14]   |
| Zr           | (0.276,0.044,0.210) | (0.277,0.043,0.210) | (0.275,0.040,0.208) |
| O$_I$        | (0.065,0.327,0.350) | (0.064,0.324,0.352) | (0.070,0.332,0.345) |
| O$_{II}$     | (0.451,0.757,0.475) | (0.450,0.756,0.479) | (0.450,0.757,0.479) |

| (T) ZrO$_2$  | Ref.[13] | Ref.[26] |
| $d_z$        | 0.057    | 0.044    | 0.057    | 0.042    |

The results, reported in Tables II A-II A, are in good agreement with experimental data and previous theoretical works within the DFT error ($\approx 1 - 2\%$).

We correctly find that the (M) phase is favored with an energy difference $\Delta E_{M-T} = 0.109$ [eV/mol] in agreement with previous works (0.063 [eV/mol]$^{[8]}$, 0.144 [eV/mol]$^{[6]}$); the experimental estimation is 0.063 [eV/mol]$^{[27]}$.

B. Yttria–stabilized Zirconia

The (T) and the (C) phase of Zirconia can be stabilized with Yttria (Y$_2$O$_3$) doping, with a phase diagram for Yttria–stabilized Zirconia (YSZ) which has been extensively characterized experimentally. Also the (T) and (C) phases of Y–doped Zirconia have been studied within a theoretical approach in a number of works$^{[8,10]}$, while no systematic study of the Y–doped (M) phase or of the (M) $\rightarrow$ (T) DIPT has been reported in the literature.

Y–doping is known to induce oxygen vacancies. We show in Fig. 1 how the Kohn–Sham (KS) density of states changes if vacancies are present both in pure and doped ZrO$_2$ crystals at low doping$^{[28]}$.

Moreover we compute the energy the system gains producing vacancies (see the caption of Fig. II for more details) for different doping concentrations. A correct modelling of the ma-
Figure 1. (colors online) Kohn–Sham density of states for (a) Yttrium doped zirconia ($\text{Zr}_1-x\text{O}_2\text{Y}_x$), (b) zirconia with oxygen vacancies ($\text{ZrO}_2-y\text{V}_y$) and (c) zirconia with one vacancy each two Yttrium atoms ($\text{Zr}_{1-x}\text{O}_{2-y}\text{Y}_x\text{V}_y$) in the tetragonal phase. The DOS are at $x = 0.25$ and $y = 0.125$ (red continuous line) and $x = 0.125$ and $y = 0.0625$ (orange dashed line); the DOS of pure $\text{ZrO}_2$ (thinner black dot–dashed line) is also shown. The system doped with Yttrium gains energy when vacancies are created, releasing oxygen molecules, (panel (d)). Here $E_{\text{gain}} = 1/y\{E(\text{Zr}_1-x\text{O}_2\text{Y}_x) - [E(\text{Zr}_{1-x}\text{O}_{2-y}\text{Y}_x\text{V}_y) + y/2\mu(\text{O}_2)]\}$, with $x = 2y$ for the “Y–doping” case (on the left) and $x = 0$ for pure $\text{ZrO}_2$ (on the right). The creation of vacancies is instead highly unfavored in the undoped system.

Material must consider the possible relative positions of the dopants (as substitutional defects) and the vacancies. We found out that the relative position of Y atoms among themselves plays a minor role, in agreement with the other works, and in our system we kept these as far as possible to mimic uniform doping. Instead the position of vacancies respect to Y atoms or to other vacancies influences significantly the total energy of the system. Stapper et al. reported that in the cubic system isolated vacancies tend to be next nearest neighbor (NNN)
to Y atoms with an energy gain of $\Delta E_c \approx 0.34$ eV per vacancy against the nearest neighbor (NN) configuration in a supercell with 1 Y atom and 1 vacancy. This result is confirmed also by Ostanin et al. in a supercell with 2 Y atoms and 1 vacancy with $\Delta E_c \approx 0.3$ eV between the two configurations $(O-Y_1, O-Y_2) = (NN, NNNN)$ and $(O-Y_1, O-Y_2) = (NNN, NNN)$ in favor of the second in the (C) phase and $\Delta E_t \approx 0.2$ eV in the (T) phase. Also in the present work we found $\Delta E_t \approx 0.12$ eV considering 1 vacancy and 2 Y atoms.

In the (M) phase two nonequivalent oxygen atoms exist, one with coordination 3 and another with coordination 4, and, accordingly there are four possible configurations. we found that the case $(NN, NNNN)x3$ favored of 0.21, 0.36 and 0.80 eV against $(NNN, NNN)x3$, $(NNN, NNN)x4$ and $(NN, NNNN)x4$ respectively; here the labels $x3$ and $x4$ refer to the coordination that would have the vacant oxygen. At the best of our knowledge, no results have been reported in the literature about the position of vacancies in the (M) phase.

The most favored configuration, for both the (T) and the (M) phase, is represented in Fig. 2 together with the forces acting on each atom in the initial geometry of pure ZrO$_2$. The Y ions are not attracted by the oxygen vacancies and indeed in the (M) phase they are pushed away from the vacancy. Hence both the analysis of the energetically favored configuration and the internal relaxation of the ions does not follow the intuitive scheme of charged defects with the vacancies a +2e and the Y ion a −1 charged sites, suggesting that exchange–correlation effects play a role in the stabilization mechanism.

Also the relative vacancy–vacancy orientation and position is in contrast with an electrostatic based model. In the (C)/(T) phase, while Stapper et al. based their calculation on the assumption that oxygen vacancies should remain as far as possible, Ostanin et al. results suggest that oxygen vacancies tend to couple along the [111] direction with a single Zr ion in between, thus at a relative distance of about $\approx 4.47$ Å. Also experimental X–ray data seems to support this idea. No data about the orientation of oxygen vacancies in the (M) phase is present in the literature.

Changing the relative position of di–vacancy complex in our model (i.e. two vacancy at $x = 12.50\%$) we found differences in total energy of $\Delta E_{tot} \approx 1.1$ eV (per di–vacancy). The oscillations in the mean field (MF), i.e. the sum of the kinetic and electrostatic energy, and the exchange–correlation (xc) component of the energy are much more pronounced, with $\Delta E_{xc} \approx 8.2$ eV and $\Delta E_{MF} \approx 7.2$ eV. We will discuss this more in detail in the next section. Our results are, in some aspects, different from the ones suggested by the two cited
Figure 2. (colors online) Tetragonal (a) and monoclinic (b) structure of $Zr_{(1-2x)}Y_{2x}O_{(2-x)}V_x$, most stable configuration at $x = 0.03125$. The forces acting on the ions in the “starting geometry”, i.e. with the atoms placed at the coordinates of the relaxed $ZrO_2$ system, are represented with blue arrows. In the tetragonal structure the oxygen atoms ($O$, small red spheres) closer to the oxygen vacancy ($V$ represented as a small black sphere in the figure) move toward it of $\approx 0.3 - 0.5$ Å, while the Zirconium atoms ($Zr$, big dark/magenta sphere) next nearest neighbor move away from it of $\approx 0.1 - 0.2$ Å from the “starting geometry”. The Yttrium atom ($Y$, biggest light/yellow sphere) is almost fix. The behavior is similar in the monoclinic structure, though the vacancy has just 3 nearest neighbor here, one of which is an Yttrium atom. The Yttrium atom moves away from the vacancy suggesting that a charged–defects interaction model based on electrostatic is not correct and that exchange–correlation effects play a major role.

works. Indeed we found that, while vacancies does not repel, the configuration suggested as most favorable for the (T) structure by Ostanin et al.\textsuperscript{9}, is not the lowest in energy. Indeed, surprisingly, the latter comes out to be unstable, with one of the two vacancies which changes position if we let the system relax (Fig. 3). This last result suggests that, in presence of vacancies, oxygen atoms can move with (nearly) no potential barrier to overcome. This is a theoretical evidence that YSZ is a good ionic conductor. Indeed this property is important for many applications and has been investigated in other works\textsuperscript{30}.

We also considered some cases at atomic doping concentrations of $x = 18.75\%$ and one test case at $x = 25.00\%$. Both for $x = 12.50\%$ and $x = 18.75\%$ there is a huge number of possible relative position of the oxygens and we did not try to systematically explore which
Figure 3. (colors online) Starting (a) and final relaxed geometry (b) of the $\text{Zr}_{1-x}\text{O}_{2-x/2}\text{Y}_{x/2}$ system with $x = 12.5\%$. $O$ atoms are dark (red) small spheres, $\text{Zr}$ atoms are big dark spheres (magenta), and $\text{Y}$ atoms are biggest light spheres (Yellow). The oxygen which changes position is a small light (light red) sphere. Before relaxation the two vacancies are aligned along the [111] direction at a distance of 8.45 Bohr; they can be detected as in the figure the oxygens are aligned in groups of four atoms everywhere except where a vacancy exist. After the relaxation two vacancies are aligned along the [101] direction at a distance of $\approx 6.2$ Bohr (this is the distance that the missing oxygens would have at the geometry of pure $\text{ZrO}_2$).

is the best configuration. At $x = 12.25\%$ the best found configuration has two vacancies aligned in the $xy$ plane. A simple explanation of this fact could be provided considering the axial anisotropy of the dielectric constant ($\kappa$) of the (T) phase. The component $\kappa_{xx} = \kappa_{yy}$ are larger than the $\kappa_{zz}$, thus providing a more effective screening of charged oxygen vacancies placed along the $xy$ plane. However we did not find this to be the sole mechanism, as some of the other configurations with the vacancies aligned in the $xy$ plane have higher energy than some configurations with the vacancies aligned, for example, in the $xz$ plane.

To conclude this section, the behavior of the oxygen vacancies present some differences, and some similarities, between the two phases of Zirconia. A key important difference however is that the formation of vacancies is more favored in the (T) phase than in the (M) phase as one can see from Fig. 1(d). This fact indeed implies that $\text{Y}_2\text{O}_3$ doping tends to stabilize the (T) phase.
Figure 4. (colors online) The difference in total energy per molecular unit (up panel) between the monoclinic and the tetragonal phase decrease increasing the $Y_2O_3$ doping (black circles). The result is not simply the sum of the effect induced by $Y$ doping without vacancies (violet triangles, dot dashed line for the linear fit) and oxygen vacancies without Yttrium (green squares, dashed line for the linear fit). When no Yttrium is present the value of the $x$ coordinate is chosen so that the number of vacancies for the $Y_2O_3$ doping case is equal to the number of vacancies for the oxygen vacancies only case. In the bottom panels the variations in the total energy as a function of the configurations is compared with the variations in the mean field and exchange–correlation energy. The pink shadowed area are a guide for the eyes, to highlight the trend of the two energy components.

C. Theoretical stabilization: $xc$–energy and vacancies.

The configuration explored allow to model the (M)$\rightarrow$(T) DIPT as a function of the Yttria doping. In Fig. 4 we see that the energy difference between the (T) and the (M) phase decrease increasing the doping concentration. This is a clear signature that we are correctly describing the DIPT. However the stabilization of the (T) phase happens, after a linear fit of our data, at higher doping concentration, $\approx 14\%$, than the experimental value $\approx 7\%^{15}$. The main error is due to an overestimation of the zero–doping energy difference of 0.056.
eV/m.u. ($E_{\text{exp}} \approx 0.063$ eV/m.u., while $E_{\text{DFT}} \approx 0.109$ eV/m.u.). Indeed it is reasonable to assume that the trend of the energy difference is better computed than its absolute value and accordingly assuming a constant “zero–doping error” for every $Y$ concentration we can subtract it (the zero energy in Fig. 4 shifts from the black line to the blue line) obtaining $x_{\text{DIP T}} \approx 7.5\%$, which is in excellent agreement with the experimental value. Similar results are obtained if one consider, instead of the linear fit, the energy difference between the two best configurations (orange dashed line in Fig. 4).

For the pure $\text{ZrO}_2$ we have also computed the contribution to the Helmholtz free energy due to thermal excitations of phonons. We found that the difference in the thermal contribution is negligible at room temperature, while the difference of the zero point phonon energy between the two phases is $\approx 0.005$ eV/m.u., with a small effect on the value of $x_{\text{DIP T}}$ (In Fig. 4 this amounts to the shift of the zero from the dashed to the continuous line).

It is interesting to compare how the mean field energy, $E_{\text{MF}}[\rho] = T[\rho] + V^{\text{ext}} + V_H[\rho]$, and the exchange–correlation energy, $E_{\text{xc}}[\rho]$, change considering different configurations. In Fig. 4 bottom panels, we see that the two component oscillate about 10 times more than the total energy as a function of the chosen atomic configuration. The mean field energy, in pure $\text{ZrO}_2$, is lower in the (T) phase (Fig. 4 central bottom panel); that is, if the xc–energy is neglected, the (T) phase is thermodynamically the most stable. The inclusion of $Y$ doping and oxygen vacancies enhance this aspect as, on average $d(\Delta E_{\text{MF}})/dx > 0$, with $x$ the atomic doping content and $\Delta E_{\text{MF}} = E^{(\text{M})}_{\text{MF}} - E^{(\text{T})}_{\text{MF}}$. This is likely because the tetragonal phase, which has an higher dielectric constant, better screens the electrostatic perturbation $V^{\text{ext}}_{\text{doping}}$ induced by the doping.

In order to have a picture of the screening effect we have defined (see Fig. 5 and caption) the polarization charge, that is the difference in the electronic density between the pure and the doped system. In Fig. 5 we see that the main perturbation is induced by the oxygen vacancies. The screening mechanism however is very efficient and accordingly the value of the polarization charge is sensibly different from zero only on the atoms NN to the vacancies.

The xc–energy in pure $\text{ZrO}_2$ instead is lower in the (M) phase and on average $d\Delta E_{\text{xc}}/dx < 0$. To understand this point we consider the perturbations induced in the electronic part of the Hamiltonian of pure $\text{ZrO}_2$, that is $V_{\text{pert}} = V^{\text{ext}}_{\text{doping}} + V^{\text{ind}}_{\text{ions}}$, where $V^{\text{ind}}_{\text{ions}}$ is the additional perturbation generated from the dislocation of the atoms. $V_{\text{pert}}$ has the shape of a random potential in a perfect periodic system which tends to destroy the collective behavior of the
Figure 5. (colors online) 3D plot of the polarization charge, in the tetragonal phase, defined as:
\[ \rho_{\text{pol}} = [32\rho(Zr_{0.94}O_{1.97}Y_{0.06}V_{0.03}) + \rho(O)] - 32\rho(ZrO_2) \]: iso-surfaces for \( \rho_{\text{pol}} = \pm \rho_0 \) are represented, with the positive ones in red (lighter), the negatives in blue (darker). Oxygen atoms are the small spheres (red), Zirconium atoms the dark big spheres (magenta) and Yttrium atoms the light biggest spheres (yellow). The (green) shadowed regions highlight the regions around an oxygen vacancy (small black sphere in center of the region), where \( \rho_{\text{pol}} \) is different from zero.

electron gas and thus electrons lose correlation. This is not an “on site” correlation, but rather a “ranged” correlation energy and so we expect that GGA can better describe this effect than LDA. While \( V_{\text{doping}}^{\text{ext}} \) is similar for the two phases, i.e. we are considering the same kind of doping, the \( V_{\text{ions}}^{\text{ind}} \) term is greater in the (T) phase due to its higher screening (see note 31). Accordingly the \( xc \)-energy is more penalized, that is \( dE_{xc}^{(M)}/dx < dE_{xc}^{(T)}/dx \). This explains \( d\Delta E_{xc}/dx < 0 \) and thus we can infer that in the screening process electrons lose
The emerging picture is that the \( \text{(M) \to (T)} \) DIPT is a balance between different mechanisms with a key role played by vacancies\(^{32}\), though the role of Yttrium atoms is not only to induce vacancies (see Fig. 4 and caption).

### III. CONCLUSIONS

The phase transition of Zirconia induced with Yttria doping is very well known experimentally and indeed Yttria stabilized Zirconia is commonly used in many applications. Our results show that an ab initio approach is able to correctly describe this mechanism, with a predicted phase transition at a doping concentration of \( \approx 7.5\% \), in good agreement with experimental data. This result confirms the opportunity to predict the effect of other less characterize kind of doping which could be of potential interest for new applications, as for example doping with magnetic materials\(^2\).

Moreover, within density functional theory we can explore how physical properties of the system are influenced by doping. As an example we considered how different components of the energy changes and in particular the behavior of the exchange–correlation energy. We showed that oxygen defects play a major role in the phase transition and how the perturbation induced by oxygen vacancies to the system is screened in the high-k tetragonal phase. The same approach could be used to check the effect of doping on other quantities, which are not easily accessible experimentally; among others, the value of the dielectric constant, which has a key importance for applications in the field of micro–electronics.

Finally these results, improving our understanding of the phase transition, can also be used to tune the parameters of models for the description of realistic devices based on Zirconia, whose dimensions are still beyond the capabilities of first–principles simulations.

### IV. ACKNOWLEDGMENTS

This work was funded by the Cariplo Foundation through the OSEA project (n. 2009-2552). The authors would like to acknowledge Professor Giovanni Onida and the ETSP\(^{32}\) Milan node for the opportunity of running simulations on the “etsfmi cluster”. We also acknowledge computational resources provided under the project OSEA by the Consorzio
1 Science and Technologies of Zirconia and Science and Technologies of Zirconia II, edited by A. Heurer and L. W. Hobbs in Advances in Ceramics, Vol. 3 (1981) and Vol. 12 (1984) (The American Ceramic Society, Westerville, OH)

2 S. Ostanin, A. Ernst, L. M. Sandratskii, P. Bruno, M. Däne, I. D. Hughes, J. B. Staunton, W. Hergert, I. Mertig, and J. Kudrnovský, PRL 98, 016101 (2007)

3 X. Wu, P. Zhou, J. Li, L. Y. Chen, H. B. Lv, Y. Y. Li, and T. A. Tang, APL 90, 183507 (2007)

4 H. Zhang, B. Gao, B. Sun, G. Chen, L. Zeng, L. Liu, X. Liu, J. Lu, R. Han, J. Kang, B. Yu, APL 96, 123502 (2010)

5 G.-M. Rignanese, J. Phys.: Condens. Matter 17, R357 (2005)

6 J. K. Dewhurst, and J. E. Lowther, Phys. Rev. B 57, 741 (1998)

7 E. V. Stefanovich, A. L. Shluger, C. R. A. Catlow, Phys. Rev. B 49, 11560 (1994)

8 G. Stapper, M. Bernasconi, N. Nicoloso, and M. Parrinello, Phys. Rev. B 59, 797 (1999)

9 S. Ostanin, A. J. Craven, D. W. McComb, D. Vlachos, A. Alavi, A. T. Paxton, and M. W. Finnis, Phys. Rev. B 65, 224109 (2002)

10 S. Ostanin, E. Salamatov, A. J. Craven, D. W. McComb, and D. Vlachos, Phys. Rev. B 66, 132105 (2002)

11 K. C. Lau, and B. Dunlap, J. Phys.: Condens. Matter 21, 145402 (2009)

12 J. P. Goff, W. Hayes, S. Hull, M. T. Hutchings, and K. N. Clausen, Phys. Rev. B 59, 14202 (1999)

13 P. Aldebert, and J. P. Traverse, J. Am. Ceram. Soc. 68, 34 (1985)

14 C. J. Howard, R. J. Hill, and B. E. Reichert, Acta Crystallogr., Sect. B: Struct. Sci. 44, 116 (1988)

15 D. G. Lamas, N. E. Walsöe De Reca, Journal of Materials Science 35, 5563 (2000)

16 D.R. Clarke and C.G. Levi, Annu. Rev. Mater. Res. 33, 383 (2003)

17 H. Jiang, R. I. Gomez-Abal, P. Rinke, and M. Scheffler, Phys. Rev. B 81, 085119 (2010)

18 R. W. G. Wyckoff, Crystal Structure (Interscience, New York, 1996)

19 P. Hohenberg, and W. Kohn, Phys. Rev. 136, B864 (1964)

20 W. Kohn, and L. J. Sham, Phys. Rev. 140, A1133 (1965)
Also for the doped case we considered only the (M) and the (T) phases. Indeed the (C) phase is a special (T) phase with $c/a = 1$ and $d_z = 0$. The description of this phase is included in our work as we let the atoms relax their position and, in few cases, also the cell parameters.

Here what we consider is the KS kinetic energy.

Here we are studying the response of pure $\text{ZrO}_2$ to a static external perturbation $V_{doping}^{ext}$. The self–consistent solution, in the Bohr–Oppenheimer approximation, of the ionic plus the electronic Hamiltonian $H_{\text{ZrO}_2}^{I+el}[\rho] + V_{doping}^{ext}$ describes the full response of the system to this perturbation. For a comparison the static dielectric $\epsilon(0)$ constant of pure $\text{ZrO}_2$ can give an insight on how the system will react to this perturbation to linear order. In the self–consistent solution the phonon contribution to the dielectric constant, $\epsilon_{ph}(0)$, is accounted by the relaxation of the atomic positions, while the electronic contribution, $\epsilon_{el}(0)$, is accounted by the the solution electronic part of the Hamiltonian $H_{\text{ZrO}_2}^{el}[\rho] + V_{doping}^{ext} + V_{indions}^{ind}$. In high–k materials like Zirconia the contribution from the lattice is dominant. The (T) phase has higher dielectric constant and accordingly we expect a greater distortion of the crystal, provided that the linear order behavior is respected to all orders.

Recent experimental results seems to suggest that the (T) phase can be stabilized simply inducing oxygen vacancies, tailoring the growing conditions of the sample, without the inclusion of other dopants.\textsuperscript{35}

\textsuperscript{14}http://www.etsf.it; http://www.etsf.eu; Nature 450, 777 (2007)

\textsuperscript{34}M. Grünig, R. Shaltaf, and G.-M. Rignanese, Phys. Rev. B B 81, 035330 (2010)
A. Lamperti, L. Lamagna, G. Congedo, S. Spiga, J. of the Electr. Soc. 158(10), G221-G226 (2011)