Analytic bond-order potential for atomistic simulations of zinc oxide

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Abstract. An interatomic potential for zinc oxide and its elemental constituents is derived based on an analytical bond-order formalism. The model potential provides a good description of the bulk properties of various solid structures of zinc oxide including cohesive energies, lattice parameters, and elastic constants. For the pure elements zinc and oxygen the energetics and structural parameters of a variety of bulk phases and in the case of oxygen also molecular structures is reproduced. The dependence of thermal and point defect properties on the cutoff parameters is discussed. As exemplary applications the irradiation of bulk zinc oxide and the elastic response of individual nanorods are studied.

1. Introduction

Zinc oxide is a traditional semiconductor with manifold applications in electronic and optoelectronic devices [1, 2], which more recently attracted much attention, because nanostructures such as rods, wires, belts, and rings [3, 4] were experimentally prepared.

In order to improve the understanding of the physical and chemical properties of zinc oxide, computer simulations can be a valuable tool complementing experiments or aiding their interpretation. Quantum-mechanical methods such as Hartree-Fock (HF) or density-functional theory (DFT) have been extensively used to investigate e.g., surfaces [5, 6], high-pressure phase transitions [7, 8, 9], elastic properties [9], phonon dispersion [10], and point defects (see e.g., references [11, 12, 13, 14, 15, 16]) in ZnO. While such methods are highly transferable and allow for accurate and reliable calculations, they very computationally demanding. Atomistic simulations of multi-million atom systems that go beyond nanoseconds in time are therefore only possible if the electronic degrees of freedom are removed and interatomic potential are introduced, which are computationally more efficient. Upon reducing the complexity of the system some information is inevitably lost, but a sensibly chosen potential scheme in combination with a carefully constructed fitting database can nevertheless realistically mimic chemical bonding and structure of solids.

Metal oxides represent an especially difficult case for atomistic modelling because of their complex electronic structure and intricate mixture of ionicity and covalency. Therefore, very few potentials have been devised and most of them are shell-model potentials (see section 2) that are not capable of describing the elemental components (see section 2). The situation is further complicated if the system of interest features such diverse phases as a hexagonal close packed metal (zinc), a gaseous phase (dimeric oxygen), and a ionic-covalent semiconductor (zinc oxide). In the following, we briefly review potential schemes which have been previously used to model compounds with ionic and ionic-covalent bonding characteristics. In doing so we motivate our interest in exploring the applicability of a purely covalent model to zinc oxide, which can be considered a prototype for other oxide systems. In section 4 a concise summary of the potential formalism adopted in the present work is given. Sections 5, 6, and 7 introduce and discuss the potentials for Zn–O, Zn–Zn, and O–O interactions, respectively. Finally, the potential is applied in simulations of the irradiation of bulk zinc oxide and to study the elastic response of individual nanorods (section 9).

2. Review of potential schemes

Bonding and structure of many elemental metals and semiconductors can be reasonably described by analytic potentials that have been used in a wide range of applications [17, 18, 19, 20, 21, 22, 23]. With regard to compounds, the situation is more complex as in most systems cohesion arises from a mixture of covalent, ionic and van-der-Waals interactions. Formally, the cohesive energy can be written as a sum of these contributions, and if a realistic description of the bonding behaviour is pursued, one needs to assess the relative weight and the mathematical representations of each of these terms. In the following, potential schemes from the literature are compared focusing on the treatment of covalency vs ionicity, and, where applicable, charge equilibration.

The alkaline-halides and alkaline earth-oxides are typical example for ionic compounds in which long-range Coulomb interactions between the ions dominate. Simple Born-Mayer potentials which combine a Coulomb potential with a shortranged, spherically symmetric repulsive potential are usually sufficient in order to obtain a satisfactory description. Shell-model potentials (see e.g., references [24, 25] and [26]) represent a somewhat more refined approach combining short-ranged repulsive, long-ranged dispersive, and ionic interactions with a simple, usually harmonic, model for atomic polarisability [27]. The ionic charges are kept fixed at their nominal values (and independent on the atomic environment). In the past, shell-model potentials have been applied to a wide range of materials since they offer a very handy formalism with few fitting parameters and yield useful models when only a small section of configuration space is of relevance. Since three-body interactions are not explicitly taken into account covalent contributions either have to be neglected or are subsumed in the fitting parameters. The formalism is incapable of describing pure elements which renders it inapt for simulations which include the boundary phases of the material. Furthermore, due to the long range of the Coulomb potential the treatment of electrostatic interactions is computationally very demanding.

In contrast to ionic interactions, covalent bonding is characterised by strong directional bonds, and therefore angle dependent many-body terms need to be taken into account. By merging Stillinger-Weber type [28] two and three-body potentials with terms describing Coulomb, monopole-dipole, and van-der-Waals interactions,

Vashishta and coworkers arrived at a scheme capable of describing materials such as silicon carbide [29, 30], gallium arsenide [31], and zinc oxide [32]. The ions are assigned fixed, effective charges and the long-range electrostatic interactions are truncated at some intermediate distance reducing the computational effort substantially. The potentials are designed such that the boundary phases can be described on the same basis as the compound.

In an effort to simulate metal/metal-oxide systems Streitz and Mintmire devised a scheme (ES+) which merges the embedded-atom method (EAM) with an ionic potential [33]. The model explicitly accounts for charge transfer between anions and cations by equilibrating the ionic charge for each configuration which renders it computationally very demanding. The scheme has been applied with some success for modelling alumina [33] and with modifications to titania [34].

More recently, Duin *et al* constructed a reactive force field (ReaxFF) which features a combination of many-body, van-der-Waals, and Coulomb terms [35]. Originally, designed for hydrocarbons the scheme is sufficiently flexible for describing also oxidic compounds and metals as shown for the case of alumina [36]. Similar to the Vashishta potentials, the Coulomb potential is truncated at some distance while the effective ionic charges are determined via charge equilibration akin the Streitz-Mintmire approach. The full functional form features 93 parameters. Many degrees of freedom for fitting increase the flexibility of the potential scheme but parameter space sampling becomes tremendously complex and the risk to generate spurious minima in the potential hypersurface grows.

Purely covalent bonding exemplified by elemental semiconductors such as carbon, silicon, or germanium, can be well described within the bond-order approach pioneered by Abell [37] and Tersoff [38] and later extended by Brenner [39]. It turns out the formalism works similarly well for strongly covalent compounds. The bond-order formalism was shown to be formally equivalent to the embedded-atom method (EAM) which has been very successful in modelling metallic bonding [40]. In fact, it turns out that the bond-order ansatz is capable of describing many pure metals (including transition metals) on the same footing as covalent materials [21, 23]. Therefore, it is possible to derive a consistent set of potentials which describe the compound as well as its components within a single formalism.

Pettifor and coworkers were able to show that bond-order potentials can be rigorously derived within tight-binding theory based on a moment expansion of the density of states [41, 42]. They later extended their approach to higher moments and multi-component systems [43]. The scheme has been applied to obtain potentials for example for hydrocarbons [44], but oxide materials have not been considered so far.

3. Motivation

In this work, we pursue the question whether it is possible to treat a highly ionic compound such as zinc oxide within a purely covalent model. The neglect of charges avoids the computationally expensive treatment of long-range interactions and therefore, allows to obtain an atomistic model which is significantly more efficient than any of the formally ionic potentials described in the foregoing section.

We employ an analytic bond-order potential (ABOP) scheme [21, 20] which has already been successfully applied to a number of systems including purely metallic [21, 23], predominantly covalent [45, 20, 18], molecular [22] as well as mixed ioniccovalent bonding [23, 22]. Thus, the ABOP scheme in principle offers the possibility to describe the Zn–O, O–O as well as Zn–Zn interactions within a single framework. Although, it turns out that for many situations the neglect of charges is legitimate, it needs to be acknowledged that the applicability of the ABOP is limited if internal or external electric fields become dominant as in the case of interfaces or surfaces. The restriction to first-nearest neighbours also implies that the energy difference between the zinc blende and the wurtzite structures becomes zero, since their local environments are indistinguishable. The ABOP performs, however, very well in reproducing a variety of bulk properties including cohesive energies, structures, and elastic properties as will be shown in the next sections.

We anticipate the analytic bond-order potential presented in this work to be useful for atomistic simulations of a variety of phenomena and processes as exemplified in section 9.

4. Methodology

The bond-order potential scheme and the fitting strategy used in this work have been extensively described elsewhere (see e.g., references [21] and [20]). Differences in the fitting strategy are pointed out where applicable. For completeness we briefly summarise the functions needed to calculate the potential energy.

The total energy is written as a sum over individual bond energies

$$E = \sum_{i>j} f_{ij}^c(r_{ij}) \left[V_{ij}^R(r_{ij}) - \underbrace{\frac{b_{ij} + b_{ji}}{2}}_{\overline{b_{ij}}} V_{ij}^A(r_{ij}) \right]$$
(1)

where for the pair-like attractive and repulsive terms Morse-like pair potentials are adopted

$$V^{R}(r) = \frac{D_{0}}{S-1} \exp\left(-\beta\sqrt{2S}(r-r_{0})\right),$$

$$V^{A}(r) = \frac{SD_{0}}{S-1} \exp\left(-\beta\sqrt{2/S}(r-r_{0})\right).$$
(2)

The parameters in these equations are the dimer bond energy D_0 , the dimer bond distance r_0 and the parameter S which determines the slope of the Pauling plot (see reference [21] and figures 1 and 5 below). The parameter β can be obtained from the ground-state oscillation frequency of the dimer. The cutoff function restricts the range of the interaction to first or second nearest neighbours

$$f^{c}(r) = \begin{cases} 1 & r \leq R_{c} - D_{c} \\ \frac{1}{2} - \frac{1}{2} \sin\left[\frac{\pi}{2}(r - R_{c})/D_{c}\right] & |r - R_{c}| \leq D_{c}, \\ 0 & r \geq R_{c} + D_{c} \end{cases}$$
(3)

where D_c and R_c are adjustable parameters. Explicit three-body interactions are included via the bond-order parameter

$$b_{ij} = (1 + \chi_{ij})^{-\frac{1}{2}},\tag{4}$$

$$\chi_{ij} = \sum_{k(\neq i,j)} f_{ik}^c(r_{ik}) \ g_{ik}(\theta_{ijk}) \ \exp[2\mu_{ik}(r_{ij} - r_{ik})].$$
(5)

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Parameter	\mathbf{Zn} - \mathbf{Zn}	0–0	Zn–O
$D_0 (eV)$	0.6470	5.166	3.60
r_0 (Å)	2.4388	1.2075	1.7240
S	1.8154	1.3864	1.0455
β (Å ⁻¹)	1.7116	2.3090	1.8174
γ	4.3909×10^{-5}	0.82595	0.019335
c	77.916	0.035608	0.014108
d	0.91344	0.046496	0.084028
$h \left[\theta_c \right]$	1.0	0.45056	0.30545
	$[180^{\circ}]$	$[116.8^{\circ}]$	$[107.8^{\circ}]$
$2\mu ({\rm \AA}^{-1})$	0.0	0.0	0.0
R_c^e (Å)	3.00	2.10, 2.60	-
D_c^e (Å)	0.20	0.20	-
R_c (Å)	2.85	2.45	2.60
D_c (Å)	0.20	0.20	0.20

Table 1. Parameter sets for describing Zn–Zn, O–O, and Zn–O interactions. R_c^e , D_c^e : cutoff parameters derived for the pure elements; R_c , D_c : default cutoff parameters for simulations of ZnO (see section 8).

The indices monitor the type-dependence of the parameters, which is of importance for the description of compounds. The angular dependence is described by

$$g(\theta) = \gamma \left(1 + \frac{c^2}{d^2} - \frac{c^2}{d^2 + (h + \cos \theta)^2} \right).$$
 (6)

The parameter sets for Zn–Zn, O–O, and Zn–O derived in this work are given in table 1 and will be discussed in the following sections. Cutoff parameters derived for the pure elements are denoted R_c^e and D_c^e , while the parameters appropriate for simulations of the compound ZnO are given in rows R_c and D_c .

5. Zinc oxide

Our fitting methodology requires data on cohesive energies, lattice constants, and elastic constants of a variety of structures in order to cover a range of coordinations as wide as possible. For zinc oxide a plethora of data from experiment and quantummechanical calculations is available which could be used for fitting and benchmarking the potential. Therefore, no additional reference calculations were necessary. In order to simplify fitting we transformed the experimentally measured hexagonal elastic constants to the cubic system by means of Martin's transformation method [46] and included the latter values in the fitting database.

5.1. Dimer properties

As the pair parameters of the ABOP are usually adjusted to dimer data, we begin by discussing the properties of the ZnO molecule. The dimer behaves peculiar in the way that its ground state dissociates into excited atomic states (T1: ZnO(X¹ Σ) \rightarrow Zn(¹S) + O(³P)) while the dissociation into the atomic ground states occurs from an excited state of the dimer (T2: ZnO(a³\Pi) \rightarrow Zn(¹S) + O(¹D)). This implies that the lowest experimentally determined dimer energy (associated with the extrinsic bond energy, EBE) cannot be described by a single pair potential but corresponds to

Table 2. Summary of properties of the ZnO dimer. Experimental data from references [47], data from quantum-mechanical calculations from references [48, 49]; EBE: extrinsic bond energy (eV); IBE: intrinsic bond energy (eV); r_0 : bond length (Å); ω_0 : ground state oscillation frequency (cm⁻¹); IBE, r_0 and ω_0 are given for the dissociation of the dimer ground state into higher energy atomic states (T1: $\text{ZnO}(X^1\Sigma) \rightarrow \text{Zn}(^1\text{S}) + \text{O}(^3\text{P})$).

	Expt.	Theory	ABOP
EBE	1.61 ± 0.04	1.20 - 1.63	
IBE	3.58	3.31, 3.60	3.60
r_0		1.679 - 1.771	1.724
ω_0	805 ± 40	646 - 913	708

a crossing-over of the potentials describing the transitions T1 and T2, respectively [47, 49]. For consistency, we only consider the dimer energy, bond length and oscillation frequency which describe transition T1, that is the decomposition of the dimer ground state into the excited atomic states (intrinsic bond energy, IBE). The parameter set given in table 1 yields very good agreement with both the experimental as well as the quantum-mechanically computed data for this transition as shown in table 2.

5.2. Bulk Properties

The performance of the ABOP with respect to bulk properties is compared to experiment and DFT calculations in table 3. The energy difference between the wurtzite and zinc blende structures is zero due to the neglect of long-range interactions. For the same reason the axial ratio $(c/a = \sqrt{8/3})$ as well as the internal relaxation parameter (u = 3/8) of the wurtzite structure are restricted to their ideal values. Otherwise, the agreement with respect to cohesive energies, volumes and bulk moduli is excellent. In particular, the elastic constants of the wurtzite phase compare very well with experiment. In table 3 we also give the elastic constants of wurtzite calculated directly with the values obtained by Martin's transformation method from the elastic constants of the zinc blende structure and find very good agreement.

The semi-logarithmic plot of bond energies vs bond lengths in figure 1 reveals an almost perfect agreement with the Pauling relation. By applying the commontangent construction to the energy-volume curves shown in figure 2 one obtains the transition pressure for the wurtzite-rock salt transformation as 10 GPa in good agreement with experimental values in the range of 8.6 to 10 GPa as well as firstprinciples calculations predicting transition pressures between 8.6 and 14.5 GPa (see reference [55] and references therein).

The comparison also includes the shell-model potential by Lewis and Catlow (LC) [24, 25] which is computationally much more expensive than the ABOP. For our calculations with the LC potential we employed the General Utility Lattice Program (GULP) [59]. The LC potential yields elastic constants in good agreement with experiment but reveals some deficiencies in the description of the equilibrium volumes and energies of the rock salt and zinc blende phases.

As a final test, we have evaluated the phonon dispersion for wurtzite with the ABOP and the LC potential and compared the results with experiment and quantummechanical calculations as shown in figure 3 (reference [10] and references therein). **Table 3.** Summary of bulk properties of zinc oxide as obtained from the analytic bond-order potential (ABOP) in comparison with data from experiment [50, 51, 52, 53, 54], density-functional theory (DFT) [9, 55, 7, 8, 56, 57] and Hartree-Fock (HF) [7, 56] calculations as well as the shell-model potential due to Lewis and Catlow (LC) [24, 25]. E_c : cohesive energy (eV/f.u.); ΔE : energy difference with respect to ground-state structure (eV/f.u.); V_0 : equilibrium volume (Å³/f.u.); a: lattice constant (Å); c/a: axial ratio; u: internal parameter of wurtzite structure; B, B': bulk modulus (GPa) and its pressure derivative. c_{ij} : elastic constants (GPa); c_{44}^0 : unrelaxed shear modulus of zinc blende structure (GPa); ζ : Kleinman parameter [58].

	$\mathbf{Experiment}$	ABOP	$\mathbf{Quantum}$	\mathbf{LC}		
			LDA-DFT	GGA-DFT	HF	
zinc ble	nde (F $\overline{4}3m$), n	o. 216, B3				
ΔE		0.0^a	0.015	0.013	0.052	0.097
a		4.552	4.509	4.633	4.614	4.547
B	145^{b}	144	162	135	157	144
B'		4.4	4.0	3.7	3.6	3.6
c_{11}	193^{b}	192				175
c_{12}	121^{b}	122				129
$c^{\prime c}$	36^{b}	36				23
c_{44}	54^{b}	56				106
c_{44}^{0}		149				
ζ		0.72				
wurtzite	e (P6 ₃ mc), no.	186, B4				
E_c	-7.52	-7.52	-9.769	-7.692	-5.658	-39.92^{d}
V_0	23.61	23.59	22.874	24.834	24.570	23.46
a	3.242^{c}	3.219	3.199	3.292	3.290	3.268
c/a	1.6003^{c}	1.6330	1.6138, 1.604	1.6076	1.593	1.553
u	0.3819^{c}	0.3750	0.3790	0.3802, 0.381	0.3856	0.3920
B	136 - 183	$14 \ (144^b)$	162,138,160	134, 125	154	143
B'	3.6 - 4	4.4	4.1, 4.4	3.8	3.6	3.3
c_{11}	207, 210	$212 \ (210^b)$	209	230		236
c_{12}	121	$116 (117^b)$	85	82		112
c_{13}	106, 105	$109 \ (108^b)$	95	64		105
c_{33}	210, 211	$219 (220^b)$	270	247		188
c_{44}	43, 45	$43 (41^b)$	46	75		74
$c_{66}{}^{c}$	45, 44	$48~(47^b)$	62	74		62
rock sal	rock salt (Fm3m), no. 225, B1					
ΔE		0.237	0.158	0.237	0.242	0.095
a	4.271 - 4.283	4.275	4.229	4.345	4.225, 4.294	4.267
B	170 - 228	200	206, 205	173	203, 132	192
B'	3.5 - 4	4.7	3.9, 4.9	3.7	3.6, 3.8	3.7
caesium	chloride (Pm	Β̃m), no. 221	1, B2			
ΔE		0.976	1.307	1.358		1.555
a		2.642	2.624	2.705		2.605
B		218	194	157		216
B'		5.0	4.0	3.8		3.8

 a due to restriction to first neighbours.

^b transformed from the hexagonal/cubic elastic constants using Martin's transformation method [46]. ^c cubic: $c' = (c_{11} - c_{12})/2$; hexagonal: $c_{66} = (c_{11} - c_{12})/2$.

 d This value includes the atomic ionisation energy



Figure 1. Pauling plot for zinc oxide comparing the analytic bond-order potential (ABOP) with data from experiment, Hartree-Fock (HF) and density-functional theory (DFT) calculations.



Figure 2. Energy-volume curves for bulk structures of zinc oxide illustrating the good agreement between the analytic bond-order potential (ABOP) and the reference curves which were obtained by combining data from experiment and first-principles calculations. The transition pressure can be obtained from this plot by means of the common tangent construction which is equivalent to finding the crossing point of the enthalpy curves. For the Lewis-Catlow shell-model potential (LC) the cohesive energy of the wurtzite structure was set to the experimental value (see footnote in table 3).

The ABOP shows a very good agreement with the experimental and first-principles data for the lower six branches of the dispersion relation. The deviations are somewhat larger for the LC potential but the overall agreement is still reasonable. On the other hand, the differences are more significant for the upper six (optical) branches. The LC potential yields qualitatively the correct shapes of these bands but fails to predict the phononic band gap and largely overestimates the splittings. In contrast, the ABOP underestimates the splitting of the bands but successfully predicts the existence of a phononic band gap. The shortcomings of both potential in the description of the higher lying branches are related to an overestimation (LC) or respectively an underestimation (ABOP) of the ionicity of the bond and the atomic polarisabilities.



Figure 3. Phonon dispersion relations for (a) the analytic bond-order potential (---) developed in the present work and (b) the shell-model potential by Lewis and Catlow. Dashed lines (- - -) and black circles (•) show data from density-functional theory calculations and experiment, respectively [10].

In particular the very good description of the lower branches is very encouraging with respect to the applicability of the ABOP.

6. Zinc

Since Zn–Zn interactions are practically absent in the compound, they have to be fitted independently of the Zn–O parameter set. Although the role of *d*-electrons is *per-se* not taken into account in the ABOP scheme (compare section 2), it has turned out that the transition metals platinum [21] and tungsten [23] can be very well described within the ABOP framework. While this experience is encouraging with respect to fitting a similar potential for zinc, it must be acknowledged that the 3*d*-electrons in zinc have a much more pronounced effect on the bonding behaviour than in platinum or tungsten most prominently embodied by the unusually large axial ratio of hcp-Zn. Keeping these considerations in mind, we are aiming to devise a parametrisation that can be used in conjunction with the Zn–O parameter set but is not primarily intended to be employed for simulations of pure zinc. It should also be recalled that in the past attempts to model zinc using the embedded-atom method (EAM) and modified EAM schemes have essentially failed [64, 65].

Within the ABOP scheme it turns to be impossible to reproduce all properties equally well with a single parameter set. In particular due to the short-ranged nature of the potential it is very difficult to reproduce the fcc-hcp energy difference (and thus the stacking fault energy) realistically. We have therefore focused on obtaining a reasonable fit to the energies and equilibrium volumes of various structures and accepted larger deviations in the elastic constants and the hcp axial ratio.

The fitting database comprised data from experiment and calculations. As information on low-coordinated structures is not available in the literature, we performed additional density-functional theory (DFT) calculations on several hypothetical bulk phases as described in the appendix.

	Experiment	DFT		ABOP	
		Literature	This work		
hexagonal-close packed (P6 ₃ /mmc, no. 194, A3)					
E_c	-1.359			-1.359	
V_0	14.90^{a}	14.53	14.90	14.88	
a	2.660^{a}	2.607	2.581	2.764	
c	4.863^{a}	4.937	4.791	4.498	
c/a	1.8282^{a}	1.8937	1.8563	1.627	
\dot{B}	73	78	(100^{b})	73	
B'	4.8		5.4	5.6	
face-ce	ntred cubic (Fr	$n\bar{3}m$, no. 22	25, A1)		
ΔE		0.033	0.032	0.005	
a		3.85	3.90	3.91	
B		95	95	72	
B'		4.2	5.1	5.6	
body-co	entred cubic (I	$m\bar{3}m$, no. 2	29, A2)		
ΔE		0.177	0.106	0.149	
a		3.07	3.11	3.07	
B		84	89	67	
B'		5.1	5.2	5.5	
diamond (Fd3m, no. 227, A4)					
ΔE			0.576	0.419	
a			5.72	5.86	
B			40	25	
B'			5.1	5.2	
simple cubic (Pm $\overline{3}$ m, no. 221, A_h)					
$\overline{\Delta}E$. , ,		0.270	0.281	
a			2.54	2.62	
B			65	43	
B'			5.1	5.4	

Table 4. Compilation of bulk properties of zinc as calculated using the analytic bond-order potential (ABOP) in comparison with experiment [60, 61, 52, 62] and density-functional theory (DFT, literature data from reference [63]) calculations. Symbols as in table 3 but energies and volumes are given in units per atom.

^a at 0 K as cited in reference [63].

^b energy-volume curve obtained at fixed axial ratio, c/a = 1.8563.

6.1. Dimer properties

The Zn₂ molecule is a van-der-Waals dimer with a very low binding energy on one side and a very large bond length on the other side. Since dispersion interactions are not taken into account in the ABOP scheme (compare equations (1), and (2)), we made no attempt to fit the dimer properties but instead treated D_0 and r_0 as adjustable parameters. It is noteworthy that the final values of $D_0 = 0.647 \text{ eV}$ and $r_0 = 2.439 \text{ Å}$ are comparable in magnitude to the Morse parameters describing the second excited state [66] $(4p)^1 \Sigma_u^+$ (the first excited state is also of the van-der-Waals type). For this state *ab initio* calculations yield bond energies between 1.00 and 1.13 eV (bond lengths between 2.65 and 2.97 Å), while values of 1.12 and 1.30 eV have been derived from experiment (bond length 3.30 Å, reference [66] and references therein).



Figure 4. Energy-volume curves for bulk structures of zinc as obtained from the analytic bond-order potential (ABOP) in comparison with reference curves which were obtained by combining data from experiment and density-functional theory (DFT) calculations.

6.2. Bulk properties

Table 4 provides an overview of the performance of the ABOP with respect to bulk properties in comparison with experiment and DFT calculations. The potential reproduces the energetics very well; the largest deviation from the DFT calculations occurs for the low coordinated diamond structure. The lattice parameters and bulk moduli are also in good overall agreement with the reference data as illustrated in figure 4.

The vacancy formation enthalpy and volume have been determined as 0.4 eV and $-0.3 \Omega_0$ (Ω_0 : atomic volume) which is in reasonable agreement with the experimental values of 0.5 eV and $-0.6 \Omega_0$ [67]. For the interstitial a formation enthalpy of 2.7 eV and a formation volume of $1.7 \Omega_0$ have been calculated. Experimentally, a formation volume of $3.5 \Omega_0$ has been determined but the formation enthalpy is unknown [67].

6.3. Melting behaviour

The melting behaviour of elemental zinc has been investigated by means of molecular dynamics simulations of a solid-liquid interface [68]. The simulation cell contained 768 atoms. The system was equilibrated at zero pressure at temperatures between 0 and 1000 K for at least 0.5 ns. Temperature and pressure were controlled using the Berendsen temperature and pressure controls [69] with coupling constants of 100 and 200 fs, respectively. For temperatures below 650 K and above 685 K either complete solidification or complete melting was observed while for intermediate temperatures the interface was stable for at least 2 ns. This observation agrees very well with the experimental melting point of 693 K.

7. Oxygen

In order to reproduce the O_2 molecule a simple pair potential would be sufficient but then the thermodynamically most stable phase would be a close-packed structure



Figure 5. Pauling plot for oxygen. The solid line illustrates the the Pauling relation while the symbols correspond to data from experiment, density-functional theory (DFT) and the analytic bond-order potential (ABOP) calculations. The deviation of the ABOP data points from the Pauling plot for larger bond lengths is due to the penetration of the cutoff range ($R_c^e = 2.60 \text{ Å}$).

(either hexagonal or cubic). In the present work we aim to obtain a more general representation of the bonding behaviour of oxygen.

The fitting database comprised experimental data on the dimer and ozone molecules complemented by data from first-principles calculations (compare appendix). The bond energy, bond length and oscillation frequency of the dimer fix the parameters D_0 , r_0 and β . The bond angle of the ozone molecule determines h. Thus there are four parameter left to be fitted. The parameter set which gave the best overall agreement with the input data is given in table 1. The properties calculated with the ABOP are compared in table 5 to experiment and quantum-mechanical calculations. Data sets are given for two different choices of the cutoff parameter R_c^e . The first value ($R_c^e = 2.10 \text{ Å}$) reproduces the experimental data for the bond angle as well as the cohesive energy of the ozone molecule but the cohesive energies of the higher coordinated structures are systematically overestimated. The longer cutoff ($R_c^e = 2.60 \text{ Å}$) yields a superior description of the higher coordinated phases but sacrifices the ozone molecule. The good overall agreement of the oxygenoxygen parameter set with the reference data is further illustrated by the Pauling plot shown for $R_c^e = 2.60 \text{ Å}$ in figure 5.

8. Defect properties, melting, and the role of cutoff parameters

The geometries and the energetics of point defects in zinc oxide result from an intricate interplay of electronic effects leading to rather complicated configurations such as the oxygen dumbbell interstitials [75, 76, 77]. The existence of many different stable charge states furthermore leads to a strong dependence of the formation enthalpies on the Fermi level. Obviously, these complications are impossible to capture in any analytic potential scheme, which owes its efficiency to the neglect of the electronic structure. For modelling of an ensemble of several hundred or thousand atoms, it is, however, usually sufficient to have a reasonable description of the energetics, while defect geometries play a minor role. Therefore, no attempt has been made to reproduce

Table 5. Cohesive energies and structural parameters of oxygen molecules and bulk phases calculated with the analytic bond-order potential (ABOP) in comparison with experimental data [60] and quantum-mechanical (QM) calculations [70, 71, 72, 73, 74]. E_c : cohesive energy (eV/atom), ΔE : energy difference with respect to oxygen dimer (eV/atom); r_b : bond length (Å); θ : bond angle of ozone molecule (deg); a: lattice constant (Å).

Experiment	QM calculations		ABOP			
	Literature	This work	$\overline{R^e_c=2.1\text{\AA}}$	$2.6\mathrm{\AA}$		
Dimer						
r_b 1.21	1.22 - 1.25	1.21	1.21			
$E_c -2.58$	-2.243.07	-2.85	-2.58			
Ozone, ground-state,	C_{2v}					
r_b 1.28	1.27 - 1.39	1.28	1.35	1.40		
θ 116.8	116.0		116.8	178.7		
$\Delta E = 0.49$	0.48 - 0.67	0.47	0.50	0.83		
Ozone, equilateral tri	angle (ring)					
r_b	1.45	1.45	1.60			
ΔE	0.90 - 1.13	0.98	1.43			
O_4 molecule						
r_b		1.80	1.81			
ΔE		2.26	1.81			
graphene sheet						
r_b		1.71	1.75			
ΔE		1.60	1.60			
diamond (Fd3m, no. 227, A4)						
a		4.40	4.46			
ΔE		1.95	1.95			
simple cubic (Pm $\bar{3}$ m, no. 221, A_h)						
a		2.50	2.07	2.15		
ΔE		2.49	2.08	2.17		
body-centred cubic (Im3m, no. 229, A2)						
a		2.91	2.43	2.76		
ΔE		2.32	2.12	2.27		
face-centred cubic (Fi	$m\bar{3}m, no. 225$, A1)				
a	-	3.65	3.03	3.54		
ΔE		2.29	2.14	2.31		

the defect structures obtained from quantum mechanical calculations with the analytic bond-order potential.

The bond-order potential scheme used in the present work employs a cutoff function in order to scale the interaction to zero between the first and second nearest neighbour shells. Therefore, the cutoff parameters can be varied within a certain range without affecting the ground state properties considered during fitting (cohesive energies, lattice parameters, elastic constants). On the other hand, the cutoff parameters do affect other properties such as point defect formation enthalpies, melting behaviour, or migration barriers. In the past the cutoff parameters of existing potentials have been occasionally modified in order to optimise them for certain applications, e.g., in the context of point defect properties [78, 45] or in the study of large volumetric deformations [79].

The ABOP developed in the foregoing sections was therefore tested with respect to variations of the cutoff parameters. The impact on point defect formation enthalpies can be rationalised as follows:

- The vacancy formation enthalpies are unaffected by the cutoff distances. The extent of relaxation around the vacant site is obviously insufficient for any of the participating atoms to enter the cutoff region and/or for atoms of the same type to interact.
- The introduction of surplus zinc atoms leads to direct Zn–Zn interactions whence the formation enthalpies of zinc interstitials and antisites are strongly dependent on the zinc cutoff distance. The dependence is typically monotonic but discontinuous. On the other hand their dependence on the oxygen cutoff distance is comparably small but usually continuous. The formation enthalpies of oxygen interstitials and antisites which involve oxygen surplus display the inverse behaviour of their zinc-surplus counterparts.
- The formation enthalpies for oxygen-surplus defects are virtually independent of the Zn–O cutoff distance. On the other hand, the zinc-surplus defects display some variation which can, however, not be easily rationalised as the dependence on $R_c^{\rm Zn-O}$ itself is sensitive to the choice of $R_c^{\rm Zn-Zn}$ and $R_c^{\rm O-O}$.

These observations show that the formation enthalpies of zinc and oxygen-surplus point defects (in high-symmetry positions) to some extent can be controlled independently. Vacancies are, however, exempt from this possibility as their formation enthalpies are independent of the cutoff distances. Note that since the ABOP is an atomistic model, it cannot capture the Fermi level dependence of the formation enthalpies of charged defects (see e.g., references [11, 12, 13, 14]).

Zinc oxide does not melt congruently but dissociates into liquid zinc and gaseous oxygen. Therefore, the melting point cannot be obtained from simulations of a liquid-solid interface (as done e.g., in section 6.3 and reference [68]). Simulations of single-crystalline ZnO cells, however, show qualitatively an increase of thermal stability with increasing Zn–O cutoff distance.

In table 1 a set of cutoff parameters (R_c, D_c) is given suitable for simulations of compound systems, which has been tested in a variety of simulations and which has also been used for the applications presented below. Before the present potential is used in any simulation, it is nonetheless advisable to consider the effect of the cutoff parameters explicitly with respect to the application in mind.

9. Applications

9.1. Irradiation of bulk zinc oxide

The threshold displacement energy measures which kinetic energy given to a lattice atom in a certain direction is required in order to produce at least one stable lattice defect [80, 81]. It is a fundamental quantity in irradiation physics, and the first step towards understanding the extent and kind of damage produced by any kind of energetic particle irradiation of a lattice [82]. The definition of this quantity is, however, not unique since the probability to produce a defect does not always rise steeply from zero to one at a certain energy [81]. For a more profound understanding one needs to analyse the integrated displacement probability function [83], i.e. the



Figure 6. Probability to form a defect during a recoil event in ZnO as a function of energy. Note that the data illustrates specifically the probability to form *at least* one defect; of course at higher energies in many cases more than one defect is formed.

probability to form a defect at a given energy. In order to determine this function for zinc oxide, molecular dynamics simulations were carried out employing the analytic bond-order potential derived in this work (see also Ref. [84]).

Firstly, a cell with 2280 atoms was equilibrated at 5 K and zero pressure. Then a randomly chosen atom near the centre of the cell was assigned a recoil energy in a randomly chosen lattice directions. The evolution of the system was followed for 6 ps. For each recoil atom and direction the energy was successively raised starting from 4 eV in steps of 4 eV until a defect was found. About one thousand O and Zn (atom type and direction) combinations were simulated. Since some threshold experiments were carried out with electron irradiation on single crystal samples in the *c*-axis direction, another set of simulations was carried out with recoils directed along the positive ("Zn face") *c*-axis direction, assuming an electron beam angular spread of 15 degrees about the [0001] direction. Due to the 4 eV step size all threshold values reported below have an uncertainty of $\pm 2 \,\text{eV}$.

Voronoy polyhedron defect analysis centred on the perfect atom sites was used to determine whether a defect had formed at the end of the simulation [85]. Both Voronoy interstitials and vacancies as well as antisites were counted as defects. In addition, if the final energy of the cell was at least 3.7 eV (the minimum energy to form a Frenkel pair) higher than the initial one, a defect was assumed to have formed as well.

Figure 6 shows the integrated displacement probability function [83], i.e. the probability to form a defect at a given energy considering all (atom and direction) combinations simulated. The figure shows that although the minimum threshold energy is about the same for both Zn and O recoils, the defect production efficiency increases much more slowly for O recoils. The defect production probabilities for Zn recoils along arbitrary directions and for recoils along the *c*-axis are similar.

Threshold values for Zn recoils of 57 eV at 313 K [86] and 56 eV at 5 K [87] have been determined from electron irradiation experiments. These studies were carried out on ZnO single crystals in the *c*-axis direction. More recently using ion irradiation an effective upper limit threshold for Zn recoils of $65 \,\mathrm{eV}$ was determined [88]. In electron irradiation experiments the high energy data is fit to a linear function and the threshold energy is obtained as the intersection with the energy axis. Since at energies clearly above the threshold, the overall damage production is determined by the average threshold displacement energy, the electron irradiation threshold is most appropriately compared to the average threshold energy. Note that in reference [87] the actual defect production data is clearly higher than the linear fit to the lowest data points, which can be regarded as an indication for a small defect production probability even below the average threshold, consistent with the observations in the simulations. The present simulations give an average threshold energy of 42 eV for all directions, and 45 eV along the *c*-axis direction. These values are thus in reasonable agreement with the experimental values of $< 65 \,\mathrm{eV}$ [88] and $56-57 \,\mathrm{eV}$ [86, 87], respectively.

For oxygen recoils the experimental data is more difficult to interpret. Optical absorption methods have not detected any defects which can be attributed to the O threshold [87]. (The threshold energy interpretations reported in [89] were later disputed [86]). Electron paramagnetic resonance and electrical measurements have detected an onset of damage at 310 keV electron energy, corresponding to an O recoil energy of 55 eV [86]. It should be noted, however, that the maximum electron irradiation energy in this experiment was only about 680 keV, corresponding to an O recoil energy of 155 eV. This signal may thus be related to the low fractional defect production probabilities observed in the simulations for oxygen recoils in the energy range 20-155 eV.

9.2. Elastic properties of nanorods

In recent years a variety of zinc oxide nanostructures such as rings, belts, rods, wires, and pillars with intriguing properties have been fabricated (see e.g., references [3, 90, 91, 92]). They are currently intensively investigated because of potential applications in nanoscale photonics and sensing devices. For many applications the elastic properties of these structures are crucial, which on the nanometre scale are strongly size dependent. Experimentally, the mechanical properties of individual nanorods or -tubes can be probed by means of atomic force microscopy as demonstrated e.g., for silicon carbide nanorods, carbon nanotubes [93], and molybdenum sulfide nanotube ropes [94]. To the best of our knowledge, the elastic response of individual zinc oxide nanorods has, however, not been experimentally characterised. Previous theoretical studies investigated the response of zinc oxide nanowires under tensile loading [95]. In the present work, we are interested in the behaviour of individual nanorods under shear load resembling the situation in an atomic force microscopy experiment [93, 94]. To this end, we employed conjugated gradients minimisation and analysed the results in terms of linear elastic beam theory.

Prismatic zinc oxide nanorods oriented along [0001] were created with diameters ranging from 3 to 12 nm and heights between 8 and 156 nm. The nanorods were placed on [0001] oriented ZnO substrates with fixed bottom layers. This mimics the effect of a stiff bulk layer underneath. In order to test the impact of the elastic deformation substrate all simulations were also carried out with all substrate atoms fixed. Periodic boundary conditions were imposed in the (0001) plane. First, the atoms in the nanorod were displaced to impose a specified shear strain. Then, the entire system was relaxed using conjugated gradients minimisation with the atoms in the top layer tethered in the $[2\overline{110}]$ direction. This procedure was repeated for different shear strains.



Figure 7. Dependence of the effective shear modulus on the aspect ratio of the nanorods. Open symbols indicate simulations in which all substrate atoms were fixed, while filled symbols correspond to simulations in which only the bottom layer of the substrate was fixed.

The elastic energy stored in a beam subject to a force F which causes a displacement u is

$$E_{st} = \frac{1}{2}Fu.$$

Taking into account both shearing and bending of the beam, and assuming constant stiffnesses EI and $GA\kappa$, one obtains [96]

$$E_{st} = \frac{1}{2} \left(\underbrace{\frac{h^3}{3EI}}_{\text{bending}} + \underbrace{\frac{h}{GA\kappa}}_{\text{shearing}} \right)^{-1} u^2$$

where E and G are Young's modulus and the shear modulus, I is the moment of inertia of the beam, d and h are the diameter and the height of the beam, respectively, A is the cross section, and κ is a geometry factor which takes into account the shape of the cross section. Replacing the displacement by the shear strain $\gamma = u/h$ and dividing by the volume V = Ah yields

$$E_{st}/V = \frac{1}{2}c^*\gamma^2$$

with the effective elastic modulus

$$c^* = \left(\frac{1}{G\kappa} + \frac{Ah^2}{3EI}\right)^{-1}$$

For a hexagonal cross section, the moment of inertia is $I = 5\sqrt{3}d^4/512$, the geometry factor is $\kappa = 0.901$, and the cross section is $A = 3\sqrt{3}d^2/8$. Introducing the aspect ratio $\beta = h/d$ the effective modulus is given by

$$c^* = \left(\frac{1}{G\kappa} + \frac{64\beta^2}{5E}\right)^{-1}.$$
(7)

For the present geometry the ABOP yields E = 178 GPa and $G = c_{44} = 43$ GPa.

As shown in figure 7 the elastic response of nanorods is strongly size dependent. Open symbols indicate the results obtained for a completely rigid substrate while filled symbols correspond to simulations in which only the bottom layer was fixed. Since in the latter case the substrate can be deformed, the effective shear moduli are systematically lower than in the case of a fully rigid substrate.

The solution obtained from classical beam theory given by equation (7) is indicated by the solid line in figure 7. On the present length scales one would assume that surface relaxations and non-linear contributions, which are captured by the ABOP, lead to a pronounced deviation from a continuum description. The agreement between the atomistic simulations and the continuum model is, however, very good even for very small dimensions. Since equation (7) works reliably for the very small nanorods investigated in the present work, it is reasonable to assume that it is also applicable for much larger nanorods.

10. Conclusions

We have derived an analytic bond-order potential for zinc oxide which describes ZnO as a predominantly covalently bonded system. Since the interatomic potential developed in this work is short-ranged it is computationally more efficient than ionic potentials which require the evaluation of long-ranged Coulombic interactions. The neglect of charges has some drawbacks in the sense that explicit electrostatic effects which play a role for example at surfaces due to unscreened charges cannot be captured. Otherwise the ABOP provides a very good description of many properties of zinc oxide.

The Zn–O parameterisation is complemented by parameter sets for the elemental phases of zinc and oxygen. The zinc potential yields a good description of the coordination dependence of the bond lengths and cohesive energies. The oxygen potential is capable of describing oxygen molecules as well as several hypothetical bulk phases. Thereby, the present potential is applicable in simulations which require a thermodynamic model for the entire Zn–O system.

The applicability of the potential has been demonstrated in an exemplary cases, namely the determination of threshold displacement energies in ZnO and the study of the elastic response of individual nanorods.

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Appendix A. Total energy calculations

In order to apply the fitting strategy outlined previously [21, 20], the energy differences and bond lengths of various bulk structures with different coordinations have to be included in the fitting database. For zinc oxide a broad data basis is available but for zinc and oxygen, further input was required.

Appendix A.1. Zinc

Singh and Papaconstantopoulos [63] calculated cohesive energies and lattice parameters of the face-centred cubic (fcc), body-centred cubic (bcc) and hexagonalclose packed (hcp) structures based on density-functional theory (DFT) calculations using the linearised-augmented plane-waves (LAPW) method. We complemented these data with plane-wave pseudo-potential (PWPP) calculations on the lower coordinated simple cubic (sc) and diamond (dia) structures. These calculations were carried out within the local-density approximation (LDA) in the Teter-Pade parameterisation [97] using the PWPP code ABINIT [98, 99]. We employed the normconserving pseudo potentials due to Troullier and Martins (TM) [100] including the 3d-electrons as part of the valence. We used a plane-wave cutoff-energy of 70 Ha. Brillouin zone sampling was performed employing 572 (hcp), 570 (fcc), 240 (bcc), 570 (dia) and 220 (sc) k-points distributed on shifted Monkhorst-Pack meshes. With these parameters a convergence better than 1 meV/atom was achieved. The calculated energy-volume curves shown in figure 4 were fitted to the Birch-Murnaghan equation of state [101]. The results are summarised in table 4. The agreement with experimental data and previous theoretical work is very good. The systematic underestimation of the lattice constants as compared to experiment is a well know deficiency of the LDA. In order to compensate for the systematic overbinding of the LDA.

Appendix A.2. Oxygen

A wealth of experimental as well as theoretical data is available for the O_2 and O_3 molecules (see e.g., references [60, 70, 71, 72, 73, 74]). Additional calculations were performed on higher coordinated molecules and bulk structures. The CASTEP code [102, 103] was used to perform DFT calculations within the spin polarised generalised gradient approximation (GGSA) based on the Perdew-Wang parameterisation [104] (PW91), and the Gaussian94 code [105] for DFT calculations using the BLYP [106] and B3LYP [107] functionals as well as for Hartree-Fock calculations. Preliminary tests proved the GGSA-PW91 DFT method to be reliable whence it was selected for the further computations.

For the calculations ultrasoft pseudopotentials were used employing a plane wave cutoff energy of 380 eV (norm-conserving pseudopotentials were also considered and gave basically identical results); finite basis set corrections were included to compensate for imperfect Brillouin zone sampling. The number of k-points was chosen to obtain convergence of the total energy better than 15 meV/atom. Using these parameters the relaxed structures and cohesive energies of several molecules and bulk structures were computed. In the case of the O₃ molecule the ground state as well as the equilateral triangle geometry, which is known to be a local minimum on the potential energy surface (see e.g., reference [74]), were considered. For the O₄ molecule the tetrahedron geometry was considered. The higher-coordinated structures comprised graphene as well as diamond, sc, bcc, and fcc. The data is shown in table 5. Where comparison is possible the DFT data computed in the present work is in very good agreement with experiment and previous quantum mechanical calculations.

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