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Iron chromium potential to model high-chromium ferritic alloys

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We present an Fe–Cr interatomic potential to model high-Cr ferritic alloys. The potential is fitted to thermodynamic and point-defect properties obtained from density functional theory (DFT) calculations and experiments. The developed potential is also benchmarked against other potentials available in literature. It shows particularly good agreement with the DFT obtained mixing enthalpy of the random alloy, the formation energy of intermetallics and experimental excess vibrational entropy and phase diagram. In addition, DFT calculated point-defect properties, both interstitial and substitutional, are well reproduced, as is the screw dislocation core structure. As a first validation of the potential, we study the precipitation hardening of Fe–Cr alloys via static simulations of the interaction between Cr precipitates and screw dislocations. It is concluded that the description of the dislocation core modification near a precipitate might have a significant influence on the interaction mechanisms observed in dynamic simulations.

Keywords: interatomic potential; atomistic modelling; iron chromium alloys; thermodynamics; dislocations

1. Introduction

High-chromium ferritic–martensitic steels (\sim 9–12 at% Cr) are the materials of choice for high temperature applications in aggressive environments (e.g. corrosion and/or irradiation). As a consequence, they are the commonly proposed structural materials for advanced nuclear reactors. This choice is supported by their superior thermal, corrosion and radiation resistance compared to austenitic steels. A first approximation to describe such steels in a modelling framework is the Fe–Cr binary alloy.

The addition of chromium to iron significantly influences the response to irradiation. It has been shown that the swelling in Fe–Cr is about one order of magnitude lower than in pure Fe for the same dose [1–5]. A remarkable effect of Cr is also reported in the shift of the ductile-to-brittle transition temperature (DBTT) in irradiated ferritic–martensitic steels. This shift is found to reach a minimum

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around 9 at% Cr [6,7], in a range of irradiation temperatures from 300 to 410° C and for doses from 7 to 36 dpa. This result is, in fact, an important reason for choosing Cr concentrations around 9 at% in most steels proposed for nuclear applications [7].

Even in the absence of irradiation, the Fe–Cr system exhibits a number of peculiarities. Density functional theory (DFT) calculations have shown that the mixing enthalpy exhibits a change of sign [8–14], which is negative below a critical concentration of ~10 at% Cr and positive above it. A negative mixing enthalpy indicates solubility and a tendency to partial ordering, while a positive value indicates that Cr atoms cluster into a separate Cr-rich phase (α' phase). These results are compatible with experimental observations that show an inversion of sign of the short-range order (SRO) parameter from negative to positive, with the zero crossing occurring at about 10 at% Cr [15–17]. However, this behaviour is not reflected in the standard Fe–Cr equilibrium phase diagram [18], whose revision was therefore recently proposed [19] by shifting the concentration of the Fe-rich phase boundary of the meta-stable (when ignoring the sigma phase) miscibility gap to a larger Cr value (~8 at% Cr), even at low temperature.

To better understand the behaviour of Fe–Cr alloys under irradiation, a description at the atomic level is necessary. To address problems of technological interest (e.g. plastic flow behaviour and evolution of the nano- and microstructure); however, description at large time and space scales are necessary. Both small and large-scale atomistic simulations essentially depend on the interatomic potentials employed, which must strike a balance between the conflicting needs of computational speed and reliability. Thus, an interatomic potential describing the Fe–Cr alloy reasonably well in the temperature range of interest is needed.

The complex behaviour of Fe–Cr is related to magnetic interactions [11]. Yet, in the literature, two interatomic potentials have been developed in an embedded atom method (EAM) like formalism, which are capable of reproducing Fe–Cr's heat of mixing behaviour as a function of composition, even without explicitly introducing magnetism. Although both formalisms succeed at describing the complexity of the ferro-magnetic phase in an effective way, temperature-dependent effects attributed to magnetic transitions cannot be captured within these models. These potentials are the two-band model (2BM) potential developed by Olsson et al. [20] (henceforth OLS) and the concentration dependent model (CDM) potential developed by Caro et al. [21] (henceforth CAR). Both formalisms are adaptations of the EAM, introducing local concentration dependence. Both potentials, however, suffer from some important drawbacks regarding thermodynamic properties, point-defect properties and the description of screw-dislocations. In this work, we develop a 2BM potential to correct those short-comings, while keeping the known useful qualities of both potentials.

The paper is organised as follows. In Section 2, we describe the 2BM formalism and the fitting methodology used; in Section 3, calculated properties for the pure potentials are presented and compared with existing potentials. Sections 4 and 5 report, respectively, thermodynamic and point-defect properties of our potential, and compare them to experiment and DFT data; also included is a comparison against OLS (strictly, the one fitted to PAW data from [20,22]) and CAR. In Section 6, as a first application of our potential, we study Cr precipitation hardening by analysing the interaction between a screw dislocation and a Cr-precipitate. Finally, Section 7 summarises the advantages and drawbacks of the different potentials.

2. Formalism and fitting methodology

It has been shown that both the 2BM and the CDM formalisms are adequate models to describe Fe–Cr's complex mixing enthalpy without accounting for magnetism explicitly [23]. Here, we choose the 2BM formalism, where the total energy is given as in the EAM with an extra embedding term, F^{s} :

$$E = \frac{1}{2} \sum_{i,j(i \neq j)} V_{t_i t_j}(r_{ij}) + \sum_i F_{t_i}^{d}(\rho_i^{d}) + \sum_i F_{t_i}^{s}(\rho_i^{s}).$$
(1)

The additional embedding term on the right-hand side is assumed to stem from the s-band electrons, while the standard embedding, F^{d} , is thought of as resulting from the d-band electrons [20]; t_i denotes the atom type (in our case Fe or Cr) on site *i*. The electron densities, ρ^{d} and ρ^{s} , for the d- and s-band, respectively, are calculated as

$$\rho_i^{\lambda} = \sum_{j(j \neq i)} \varphi_{t_i t_j}^{\lambda}(r_{ij}).$$
⁽²⁾

Here, φ^d and φ^s are the d- and s-density functions, respectively, which comply with the relations

$$\begin{cases} \varphi_{AA}^{d} = \varphi_{BA}^{d} = \varphi_{A}^{d} \\ \varphi_{BB}^{d} = \varphi_{AB}^{d} = \varphi_{B}^{d} \\ \varphi_{AA}^{s} = \varphi_{BB}^{s} = 0 \\ \varphi_{AB}^{s} = \varphi_{BA}^{s}. \end{cases}$$
(3)

This choice is in line with [20] so that the d-density reduces to the EAM density, and the s-density samples the local concentration [23]. In this way, the s-embedding terms are only relevant for the alloy and do not contribute to the pure elements' energy, which keeps the standard EAM expression.

Thus, for the Fe–Cr system and within the 2BM, 10 functions need to be determined: V_{FeFe} , F_{Fe}^{d} , φ_{Fe}^{d} , V_{CrCr} , F_{Cr}^{d} , φ_{Cr}^{d} , V_{FeCr} , F_{Fe}^{s} , F_{Cr}^{s} and $\varphi_{\text{FeCr}}^{s}$. The first three functions determine the Fe potential, the second three the Cr potential and the last four the mixed Fe–Cr interactions. For pure Fe, we chose the potential developed by Mendelev et al. [24], while the pure Cr potential and the mixed Fe–Cr interactions were fitted as described later. As is also detailed later, the main reason to refit the potential for pure Cr is to provide a better description of dislocation properties in this metal. Generally, the fitting of an interatomic potential to material properties is a problem of matching data obtained from a trial function to a given data set. This problem can be cast into one of minimising the overall squared deviation, so-called objective function (OF), between the calculated properties and the associated reference data, possibly also imposing constraints. Within our methodology, those properties become linear in the fitting parameters, reducing the scheme

to a quadratic programming problem [25], which is exactly solvable; more details are given below.

2.1. Chromium potential

Prior to fitting the pair interaction V and the embedding term F, we define a reasonable form for the electron density function φ . For this purpose, a simple Thomas–Fermi screening function is selected, which is shifted and truncated for a smooth cut-off between third and fourth nearest neighbour distance (on the bcc equilibrium crystal):

$$\varphi(x) = \varphi_0 \left(\frac{\exp(-\beta x)}{x} - \frac{\exp(-\beta x_{\text{cut}})}{x_{\text{cut}}} \right) f_{\text{cut}}(x), \tag{4}$$

with $x = r/r_0^{\ln n}$, *r* the distance in Å, $r_0^{\ln n}$ the nearest neighbour distance for the bcc equilibrium crystal (with lattice parameter $a_0 = 2.878$ Å), $x_{\text{cut}} = 1.65 r_0^{\ln n}$ the cut-off distance, $\varphi_0 = 0.0676504617$ a normalisation factor and $\beta = 5$. The cut-off function f_{cut} is defined as

$$f_{\rm cut}(x) = \begin{cases} 1, & x \le 1\\ 1 - \frac{(x-1)^3}{(x_{\rm cut} - 1)^3}, & 1 < x \le x_{\rm cut} \\ 0, & x_{\rm cut} < x \end{cases}$$
(5)

The pair interaction V is parameterised by the cubic spline expansion:

$$V(r) = \sum_{k=1}^{N} a_k (r_k - r)^3 \Theta(r_k - r),$$
(6)

where N denotes the number of knots, a_k are the fitting parameters and Θ the Heaviside unit step function. As explained below, the embedding term F is only available in numerical format.

The essential material properties considered in the fit for pure Cr (antiferromagnetic phase unless stated otherwise) are the correct stability of different crystallographic structures, the cohesive energy E_{coh} , the equilibrium lattice constant a_0 , the elastic constants (for cubic lattice structures) C_{11} , C_{12} , C_{44} [26] (paramagnetic phase, see Section 3), and the vacancy formation energy, $E_{\rm f}^{\rm V}$. In addition to these, also included in the fit is the formation energy of self-interstitial configurations (dumbbells oriented along the $\langle 100 \rangle$, $\langle 110 \rangle$ and $\langle 111 \rangle$ directions), the vacancy migration energy, $E_{\rm m}^{\rm V}$, the {110} (111) gamma surface cut, γ , and the 1/2 (111) screwdislocation core structure. In particular, the gamma surface cut (~ 10 points) was fitted through the OF, only using the effective pair potential (i.e. neglecting higher order many-body contributions) [27]. The lattice parameter (zero pressure condition), unrelaxed vacancy formation energy, $E_{\rm f}^{\rm V}$, and the three elastic constants were fitted through equality constraints. The bcc lattice stability against fcc ($E_{\rm coh}(\rm bcc)$ - $E_{\rm coh}({\rm fcc}) > 0$) and the Duesbery and Vitek condition for a compact $1/2 \langle 111 \rangle$ screw dislocation core $(\gamma(b/3) - 2\gamma(b/6) > 0$ [28], with b the length of the burgers vector) were fitted through inequality constraints.

Given the effective pair potential, the embedding function is computed by subtraction from Rose's equation [29], which describes the energy per atom $E(\hat{a})$ for the lattice under uniform expansion or contraction. The expression as a function of the nearest neighbour distance, r^{1nn} , is given as

$$E(\tilde{a}) = -E_{\rm coh}(1+\tilde{a})\exp(-\tilde{a}). \tag{7}$$

Here, $\tilde{a} = \alpha (r^{1nn}/r_0^{1nn} - 1)$ and $\alpha^2 = 9\Omega B/E_{\rm coh}$, where Ω is the atomic volume and B is the bulk modulus. This embedding function, however, is not the one entering the potential. To ensure a smooth behaviour and, particularly, to constrain the function curvature to remain positive throughout and decreasing at high density, the former embedding is re-fitted on a discrete number of points (~100) taken as reference data. The fit is performed in a minimum square sense and by a cubic spline expansion (also fulfilling F'(0) = 0). It is the latter expansion that constitutes the embedding function. The price paid for a smooth behaviour is a violation of Rose's equation; however, in the applications of interest the lattice will never be subjected to the large, uniform contractions and dilations that are probed by Rose's equation. Finally, the last fitting step consisted in accommodating manually (i.e. by trial and error) the short distance region to fit the DFT obtained values for self-interstitial formation energies.

2.2. Alloy potential

Given the EAM potentials for the pure elements and prior to fitting V_{FeCr} , F_{Fe}^{s} and F_{Cr}^{s} , we define a reasonable form for the s-electron density function $\varphi_{\text{FeCr}}^{\text{s}}$. In line with [20], the square of a 4s-type Slater function is chosen:

$$\varphi_{\rm FeCr}^{\rm s}(r) = Kr^6 \exp(-2\zeta r) g_{\rm cut}(r). \tag{8}$$

Here, K = 20.34075425 is a normalisation constant, $\zeta = 2.5001$ and g_{cut} is a cut-off function to provide a smooth cut-off between fifth and sixth nearest neighbour distance:

$$g_{\text{cut}}(r) = \begin{cases} 1, & r \le r_c^i \\ \frac{1}{2} \left\{ 1 - \sin\left[\frac{\pi \left(r - r_m\right)}{2}\right] \right\}, & r_c^i < r \le r_c^f, \\ 0, & r_c^f < r \end{cases}$$
(9)

with $r_c^i = 5.1$ Å the inner cut-off, $r_c^f = 5.3$ Å the outer cut-off, $r_m = (r_c^i + r_c^f)/2$ and $d = (r_c^f - r_c^i)/2$. The mixed pair interaction V_{FeCr} is parameterised by a cubic spline expansion as in Equation (6), while the s-embedding functions F^s are parameterised by an extended second moment expression similar to [20,30]:

$$F^{s}(\rho) = A_{1}\sqrt{\rho} + A_{2}\rho^{2},$$
 (10)

with A_1 and A_2 fitting parameters. The square root proved to be essential in reproducing the correct trends in Cr–Cr and Cr–vacancy interactions. In fact, former attempts using spline expansions, as in [23], were met with failure. In this work,

we normalised the (maximum) s-density in such a way that unity corresponds to a single Cr hosted in the Fe matrix.

The mixed pair interaction, as well as both s-embedding functions, were fitted simultaneously to all properties of interest. The mixing enthalpy was fitted in the whole concentration range (\sim 50 points) using a variance expansion as described in [23], while, at each concentration, a zero pressure condition was imposed to obtain a lattice parameter as close as possible to the linear interpolation between the equilibrium lattice parameters of the pure elements. In addition, the Cr-Cr and Cr-vacancy interaction energies up to second nearest neighbour were fitted, as well as the Cr-vacancy migration energy in Fe, using DFT reference values as guides [12]. All the mentioned properties were fitted using appropriate weights to obtain reasonable values. In addition to the expansion coefficients of the s-embedding functions and the mixed pair potential, there remains one degree of freedom that expresses the relative weight between the d-densities of Fe and Cr, $\rho_{\rm Fe}/\rho_{\rm Cr}$. By manually changing this fraction and consequently solving the quadratic programming problem, we fitted the excess vibrational entropy, computed in the harmonic approximation [31], to a value as close as possible to the experimental one, without losing the above properties and without stabilising unphysical intermetallic compounds. Finally, the potential was manually adjusted in the short distance region to reproduce the DFT-obtained binding energy of the mixed (110) Fe–Cr dumbbell. A description of the full Fe-Cr potential is given in Appendix 1.

3. Properties of the iron and chromium potentials

From the EAM-type potentials available in the literature to describe pure Fe that are capable of reproducing the $\langle 110 \rangle$ self-interstitial configuration as the most stable, we considered those from [24,32,33], finally selecting "potential 2" developed by Mendelev et al. [24], which has been widely used and tested. In [34], it has been demonstrated that the potentials from [24] and [32] succeed best in reproducing the properties of iron that are of interest for radiation-damage studies. However, of all, our choice is the only one for which a clear proof exists of being able to consistently describe stable glide of a screw dislocation in a {110} plane [35].

Among the potentials for Cr, we considered those from [36] and [20]. The latter, henceforth O05, was used in combination with the Fe potential by Ackland et al. [32] to produce the OLS and CAR potentials. The basic properties of our potential are summarised and compared to O05 and DFT data in Table 1. There, $B = (C_{11} + 2C_{12})/3$ is the bulk modulus, $C' = (C_{11} - C_{12})/2$ is the tetragonal shear and C_{44} is the shear modulus [26]. Clearly, the experimental elastic properties are well reproduced by both potentials. Note, however, that, at zero Kelvin, Cr is antiferromagnetic and has a negative Cauchy pressure [37], which cannot be reproduced within a standard central force framework [38–40]. Above its Néel temperature (~310 K), however, Cr is paramagnetic with a positive Cauchy pressure [41]. Since we focus on technological applications above room temperatures, our potential was fitted to the elastic constants of paramagnetic Cr, linearly extrapolated down to zero Kelvin (the values given in Table 1, see e.g. [36]). Furthermore, our potential also closely fits a_0 , E_f^V , E_{coh} and E_m^V , obtaining improved values for the latter two.

| | Experiment or DFT | This study | O05 |
|---|--|------------|-------|
| a_0 (Å) | 2.878 ^a /2.834 ^b | 2.866 | 2.878 |
| B (GPa) | 208° | 215 | 208 |
| C' (GPa) | 152 ^c | 155 | 152 |
| C_{44} (GPa) | 105° | 108 | 105 |
| $E_{\rm coh}$ (eV) | 4.10 | 4.10 | 3.84 |
| $E_{\rm f}^{\rm V}$ (eV) | $2.0\pm 0.2^{d}/2.59^{b}$ | 2.52 | 2.56 |
| $E_{\rm m}^{\rm V}$ (eV) | 0.95 ^e | 0.94 | 0.99 |
| $E_{\rm coh}^{\rm fcc} - E_{\rm coh}^{\rm bcc} ~({\rm eV})$ | -0.4^{f} | -0.16 | -0.03 |
| $E_{\rm f}^{\langle 110 \rangle} - E_{\rm f}^{\langle 111 \rangle} ~({\rm eV})$ | -0.02^{b} | -0.07 | -0.02 |
| $\hat{E_{\rm f}^{(110)}} - \hat{E_{\rm f}^{(100)}}$ (eV) | -1.12 ^b | -0.58 | -1.23 |

Table 1. Summary of the basic properties of both Cr potentials.

Notes: ^aExperimental [63]. ^bDFT [20]. ^cExperimental [41]. ^dExperimental [64]. ^eExperimental [65].

^fDFT [66].

The relative stability between the different self interstitial configurations is not equally well reproduced as by O05, but the important trends given by DFT are reproduced, i.e. the $\langle 110 \rangle$ and $\langle 111 \rangle$ configurations are almost degenerate and more stable than the $\langle 100 \rangle$ configuration. Finally, our potential is closer to DFT data regarding the energy difference between the body-centred cubic (bcc) and face-centred cubic (fcc) phase, thus providing enhanced stability of the bcc phase at high temperature. In summary, from the values presented in Table 1, we can conclude that the basic properties of Cr are equally well reproduced by both potentials.

The O05 potential, however, does not reproduce the correct structure for a 1/2 (111) screw-dislocation core, which is the main reason to refit the Cr potential. This is depicted in Figure 1 via differential displacement maps [42]. Clearly O05 shows a three-fold symmetry core, the opposite of the compact one, as predicted by DFT [43] and our potential.

4. Thermodynamic properties

Prior to presenting the phase diagrams resulting from the different potentials, we compare and discuss statically calculated properties that influence the Fe–Cr phase diagram. In Figure 2, the mixing enthalpy calculated with the three potentials is compared to DFT data [8,12,44]. For the mixing enthalpy resulting from the potentials, 50 compositions in the complete concentration range were sampled. For each composition, the solutes were randomly distributed in cubic bcc crystals containing 2000 atoms and relaxed at zero pressure using a conjugate gradient method. Although different DFT techniques give different values for the mixing enthalpy, the DFT curves show the same behaviour, i.e. a change of sign in the



Figure 1. Comparison of the screw component of differential displacement maps obtained with our potential (a) and the O05 potential (b).



Figure 2. Comparison of the mixing enthalpies obtained from the potentials with DFT data [8,12,44].

mixing enthalpy, being negative below ~ 5 at% Cr and positive above. Both 2BM potentials were fitted to DFT data [12,44] obtained from special quasi-random structures (SQSs) [45] while CAR was fitted to DFT data [8] obtained from a coherent potential approximation (CPA) [46]. The mixing enthalpy curves from our potential and CAR reproduce the DFT mixing enthalpy very well, both qualitatively and quantitatively. Below ~ 25 at% Cr, this is also true for OLS; the complete curve, however, takes a symmetric shape, in contrast to the asymmetric DFT shape. As a consequence, the mixing enthalpy takes negative values at the Cr-rich side too, leading to short-range order and high Fe solubility in the Cr-rich phase. These observations contradict DFT data and have never been reported in experimental studies.



Figure 3. Comparison of the formation energy obtained by the different potentials with the corresponding DFT values [11,47,48] for various intermetallic compounds.

The formation energy of intermetallic compounds is another issue to be considered. In Figure 3, the formation energy of 57 intermetallic compounds (as described in [11,47]) calculated by the three potentials and DFT are compared. In both cases, the configurations were relaxed at zero pressure using a conjugate gradient method. Below 10 at% Cr, DFT predicts stable intermetallic compounds, consistent with the negative mixing enthalpy for the disordered alloy. Above 10 at% Cr, the formation energy of the intermetallic compounds becomes positive and remains so throughout the composition range.

In the Fe-rich region (<10 at% Cr), the most stable compounds predicted by DFT are also the most stable ones predicted by all three potentials (see Figure 3), with most importantly the lowest compound at 6.67 at% Cr. This compound is described by a unit cell with axes oriented along the [210], [120] and [001] directions with dimensions $\sqrt{5}a_0 \times \sqrt{5}a_0 \times 3a_0$. It contains 30 atoms, of which 2 Cr are separated by 7 nn distance $(\sqrt{19}/2a_0)$ along [133]. It should be noted that this compound is consistent with the ordered phase observed in the Monte Carlo simulations by Pareige et al. [48] using OLS. Among the three, both 2BM potentials also give good quantitative agreement in that concentration range. At higher Cr composition (>10 at%), OLS generally overstabilises the compounds thereby even giving negative values at 50 and 93.33 at% Cr. The latter is consistent with the low (compared to DFT) and negative mixing enthalpy of the disordered alloy at equiatomic and Cr-rich compositions, respectively. Such negative values in turn may lead to unphysical configurations in the course of, for example, Monte Carlo simulations, and thus constitute a drawback of the potential. The other two potentials do not suffer from the latter problems, although ours still somewhat overstabilises the compounds.

Experiments have shown the excess vibrational entropy to be non-negligible for the Fe–Cr system [49–51]; also, it was predicted to have a significant influence on the location of the solubility limit [44]. We calculated the excess vibrational entropy with all three potentials for disordered alloys, based on the harmonic approximation [31] in the Γ point applied to cells containing 1024 atoms, prepared as purely random samples.



Figure 4. Comparison of the excess vibrational entropy obtained from the different potentials with the experimental values from [49,51].

In Figure 4, excess vibrational entropies for all three potentials are compared with experimental values for the ferro-magnetic phase [49,51]. The experimental curves suggest a non-negligible contribution to the total excess entropy, amounting to 30–40% of the configurational entropy for the disordered alloy. As shown in the figure, none of the potentials gives a close fit to the experimental values. From the three, however, both 2BM potentials agree best, with ours and OLS closest to the data from [49] and [51], respectively. In the former case, the experimental data is underestimated by a factor of about 1.5; in the latter, it is overestimated (below ~75 at% Cr) by about the same factor. The CAR potential, on the other hand, underestimates S_{xc}^{vib} by about a factor 3 and gives negative contributions below ~12 at% Cr, which contradicts experiments (see also [44,52]).

The phase diagrams based on all three potentials are compared to a recently proposed parameterisation based on the CALPHAD methodology [53] in Figure 5. The CALPHAD miscibility gap is calculated from an experimentally based [53,54] Gibbs free energy parameterisation. To indicate the ferro to paramagnetic phase transition, the CALPHAD calculated Curie temperature is also added. The phase boundaries for our potential and OLS are obtained from isobaric Monte Carlo simulations in the semi-grand canonical ensemble [55]. In the figure, the data points are interpolated by smooth curves to guide the eye. These curves are based on the data points and the fact that full solubility was verified to occur starting from 1200 and 800 K for our potential and OLS, respectively. The phase boundaries for CAR were obtained from free energy calculations using a thermodynamic integration technique (see [52] for more details). Thus in all cases vibrational and configurational entropy were accounted for.

When focussed on the Fe-rich side, we observe large Cr solubility at low temperature, due to the negative heat of mixing. As shown in Figure 5, this behaviour is well reproduced by all potentials, with a particularly close agreement



Figure 5. Comparison of the phase diagrams resulting from the different potentials against a CALPHAD-based calculation. The curves for OLS and our potential were taken from [55], while, for CAR and CALPHAD, they were taken from [52] and [53], respectively.

for ours. At about 750 K, the Fe-rich solubility limit increases until full solubility is observed above the critical temperature \sim 900 K within the CALPHAD calculation. The curve resulting from OLS underestimates this critical temperature by \sim 200 K, while our potential overestimates it by roughly the same amount. The critical temperature resulting from CAR, on the other hand, lies above the melting temperature. Note that the difference in critical temperature between OLS and ours closely follows the different trends observed in the excess vibrational entropy. Good agreement of the critical temperature obtained with potentials and CALPHAD should not be expected since magnetic interactions are not explicitly included in the former. Below \sim 25 at%, however, direct comparison with the CALPHAD phase diagram is justified since in this concentration range the CALPHAD phase boundary is well below the Curie temperature. For higher Cr concentrations, however, care should be taken as the magnetic transition might significantly affect the solubility; thus close agreement between experiment and potential prediction might not be a concern for potentials that effectively describe the ferro-magnetic phase.

On the Cr-rich side, the Fe solubility approaches zero with decreasing temperature, which is well reproduced by both our potential and CAR. The OLS potential, on the other hand, predicts high Fe-solubility as an unphysical artefact of the negative heat of mixing at the Cr-rich side.

5. Point-defect properties

The formation, migration and binding energies reported in the current section were calculated in cubic bcc boxes containing 2000 atoms at constant volume



Figure 6. Comparison between DFT [12] and all three potentials of the interaction energy between Cr–Cr and Cr–vacancy pairs in bulk Fe.

 $(a_0 = 2.8553 \text{ Å})$ up to a precision of 1 meV. For the formation and binding energies, the boxes were relaxed using a conjugate gradient method and the migration energies were estimated using the nudged elastic band method [56].

In Figure 6, the Cr–Cr and Cr–vacancy interaction energy (E_1) in bulk Fe, calculated using the three potentials and DFT [12], is presented. The DFT data show Cr–Cr repulsion in bulk Fe that decreases fast with increasing distance. This behaviour is mainly responsible for the experimentally observed SRO and is acceptably well reproduced by all potentials. Both 2BM potentials, however, underestimate the difference in repulsion between first and second nearest neighbour Cr pairs; an effect that is well reproduced by CAR. Concerning the Cr–vacancy interaction, according to both DFT calculations [12] and experiments [57], it is essentially negligible. As shown in Figure 6, this feature is also qualitatively reproduced by all potentials, which predict weak repulsion between 1 nn and 2 nn Cr–vacancy pairs increases, as reproduced by all potentials. This is a prerequisite to obtain the correct order between forward and backward Cr–vacancy exchanges for several chemical environments in the Fe matrix, as discussed further.

In Figure 7, the average (of forward and backward) vacancy migration barriers for different local Cr configurations calculated with the three potentials are compared to the corresponding DFT values [58]. The DFT results show that the barriers are largest for both Cr and Fe exchanges when there are three Cr and three Fe atoms occupying the six nearest neighbour positions around the saddle-point. This effect is only reproduced by our potential, although OLS and CAR also give reasonable trends. Note, however, that in the case of CAR potential, the value for Cr migration in bulk Fe is significantly overestimated. As a consequence, Cr migration barriers calculated with CAR are somewhat higher as compared to



Figure 7. Comparison between DFT [58] and the three potentials of the average vacancy migration barrier in bulk Fe for different local Cr configurations around the saddle.



Figure 8. Formation energies of different interstitial configurations in bulk Fe and Cr, calculated using DFT [12] and the three different potentials.

the corresponding DFT values. For the Fe jumps, on the other hand, the migration barriers are largely underestimated when locally enriched by three or more Cr atoms. For completeness, a table summarising the precise configurations and values of both forward and backward jumps on which Figure 7 is based is given in Appendix 2.



Figure 9. Comparison between DFT [12] and the three potentials of the binding energy of Cr interstitial configurations.

In Figure 8, we show the formation energy of interstitial defects in bulk Fe, taking the $\langle 110 \rangle$ mixed dumbbell as reference; values predicted by DFT [12] and the three potentials are compared. From the DFT data [12], it follows that the $\langle 110 \rangle$ mixed dumbbell configuration in bulk Fe has the lowest formation energy. However, the inclusion of Cr in the defect diminishes the difference in formation energy between the $\langle 111 \rangle$ and $\langle 110 \rangle$ configurations.

In the case of pure Fe, all potentials follow the DFT trend reasonably well, with a proper reproduction of the relative differences between the formation energies of the $\langle 100 \rangle$, $\langle 110 \rangle$ and $\langle 111 \rangle$ configurations being the most important feature. With the introduction of Cr, the data from our potential still captures the shape of the DFT curves reasonably well, while the data from OLS give almost degenerate values for all configurations. The CAR potential, on the other hand, fails to reproduce the $\langle 110 \rangle$ mixed dumbbell as the lowest energy configuration.

Next, we compare the binding energy of Cr interstitials in bulk Fe obtained from the three potentials with the corresponding DFT values [12]. Such a comparison is presented in Figure 9. As expected from the above results, the CAR potential shows poor agreement with DFT data. Given that both 2BM potentials were only fitted to the binding energy between a single Cr and the $\langle 110 \rangle$ dumbbell, they show reasonable agreement with DFT. In particular, our potential is the only one that correctly reproduces the sign for all reported binding energies.

6. Dislocation-precipitate interaction

Here, we address the problem of dislocations in Fe and Cr and of the interaction of a 1/2(111) screw dislocation with Cr precipitates in the Fe matrix, as obtained from static simulations. The dislocation core energy and radius, shear modulus,

| | Iron | | Chromium | | |
|--|---------|------------|----------|------------|--|
| Property | OLS | This study | OLS | This study | |
| Lattice unit a_0 (nm) | 0.28553 | 0.28553 | 0.2878 | 0.2866 | |
| Core radius $r_0(b)$ | 3 | 3 | 3 | 3 | |
| Core energy \tilde{E}_C screw (eV/nm) | 4.24 | 4.27 | 6.27 | 6.23 | |
| Core energy $E_{\rm C}$ edge (eV/nm) | 6.97 | 7.01 | 9.71 | 9.6 | |
| Shear modulus* μ (GPa) | 73 | 73 | 130 | 89 | |
| Poisson** ratio v | 0.49 | 0.49 | 0.21 | 0.4 | |
| Peierls stress for 1/2(111) screw dislocation for the glide in a (110) plane (GPa) | 1.4 | 1.3 | 1.95 | 2.02 | |

Table 2. Properties of Fe and Cr as obtained from the potentials.

Notes: *The shear modulus was estimated by straining a crystal on a $\{110\}$ plane in a $\langle 111 \rangle$ direction, as was done in [67].

**The Poisson ratio was determined following the method used in [68] for pure Fe.

Poisson ratio and Peierls stress for a 1/2(111) screw and a $1/2(111)\{110\}$ edge dislocation in the pure metals are summarised in Table 2. Details regarding the calculations can be found in [59], where the interaction of an edge dislocation with Cr precipitates was characterised. From Table 2, it appears that the two Fe potentials provide very similar results, whereas the Cr potentials exhibit at least two essential differences. The shear modulus calculated with our potential is ~30% smaller, while the Poisson ratio is two times larger than according to OLS. The characteristics of the core of both edge and screw dislocations are similar with both potentials. Finally, we note that the two Cr potentials predict approximately the same Peierls stress (τ_P) for the screw dislocation, which is higher than τ_P in Fe by 35 and 55% according to OLS and our potentials, respectively.

The interaction of a screw dislocation with Cr precipitates was studied in a bcc simulation box with dimensions $20 \times 12 \times 28 \text{ nm}^3$ with axes oriented along the [112], [110] and [111] directions. Periodic boundary conditions were applied in the [111] direction and a straight screw dislocation with left-handed thread and Burgers vector b = 1/2[111] was created. A Cr precipitate was inserted at a defined distance from the dislocation line and placed so that its centre coincides with the (110) glide plane of the screw dislocation. Then the relaxation was performed to calculate the total energy of the crystal as a function of the distance between the precipitate centre and dislocation line. The precipitate–dislocation interaction energy $E_{\rm I}$ was estimated as $E_{\rm I} = (E^{\rm D+P} + E^{\rm Fe}) - (E^{\rm D} - E^{\rm P})$, following [60] (positive values indicate repulsive interaction). Here, $E^{\rm D+P}$, $E^{\rm D}$ and $E^{\rm P}$ are the total energy of the crystal containing the dislocation interacting with the precipitate, dislocation only and precipitate only, respectively. $E^{\rm Fe}$ is the total energy of the Fe crystal with the same geometry and boundary conditions as the crystals used to calculate the above mentioned energies. All crystals were relaxed to reach a convergence of at least 5 meV in the total energy.

As mentioned in Section 3, the two Cr potentials predict different core structures for the screw dislocation in pure Cr. Therefore, it is of interest to investigate whether or not the non-degenerate-to-degenerate core structure modification occurs at the

Table 3. Differential displacement maps drawn in the (111) zone for a screw dislocation interacting with a 2-nm precipitate. The thick dashed line shows the edge of the precipitate and a double arrow shows the directions of the preferential split of the dislocation core. The centre of the dislocation core is shown by the symbol §.



precipitate-matrix interface and inside the precipitate. It should be noted that such a modification was found to occur with OLS in our previous study [61]. To address this issue, the core structure of different dislocation segments, selected according to the scheme presented in Table 3, was constructed using the differential displacement (DD) method [42] from the atomic positions obtained after relaxation. The corresponding DD maps are also presented in Table 3. Firstly, we observe that the core structure of the segment located near the centre of the precipitate was found



Figure 10. Precipitate dislocation interaction energy as function of the distance for our (left) and the OLS potential (right). The position of maximal force between precipitate and dislocation is indicated by F_{MAX} .

to be the same as in pure Cr for the corresponding potentials, i.e. the three-fold split core for OLS and isotropic for our potential (Figure 1). The core of the segment penetrating the precipitate normal to its surface (#1 of Table 3) has a two-fold structure with pronounced split over the precipitate surface, similar for both potentials. However, a difference in the core structure was found for the segments penetrating the precipitate at a tangent to its surface (#2 and #3 in Table 3 as the limiting case). With our potential no isotropic or three-fold structure modification occurs, contrary to OLS. Additional calculations, not reported here, proved that, with OLS, the isotropic-to-degenerate core transformation occurs not only inside the precipitate but also everywhere near the precipitate–matrix interface. The latter implies that, in MD simulations, such core transformation may manifest itself as emission of kink pairs in the {112} plane, thus causing cross-slip movement instead of stable glide (see [43] for details).

The interaction energy of the dislocation with precipitates of different sizes as a function of distance between precipitate centre and dislocation line is presented in Figure 10. Qualitatively, the curves look similar for both potentials, showing a maximum in the centre of the precipitate and a fast decrease beyond the precipitate radius. Quantitatively, however, the maximum interaction energy estimated with OLS is about twice as high considering precipitates of the same size. The derivative of the interaction energy with respect to the distance gives an estimate for the interaction force, from which the maximum force (and hence stress) necessary for the dislocation to shear a precipitate can be calculated. Irrespective of the precipitate size and applied potential, the maximum interaction force occurs at $\sim 65\%$ from the precipitate centre.

The difference in magnitude of the interaction energy between the two potentials can be attributed to the difference in the shear modulus of pure Cr (see Table 2). For a more correct treatment, however, contributions from the chemical energy (due to additional Fe–Cr interface) and dislocation core energy (which differs in Fe and Cr) should not be neglected. Therefore, we have performed additional calculations allowing the total interaction energy to be decomposed into the contributions coming from the shear modulus misfit, chemical and relaxation (related to the strain



Figure 11. Different contributions to the interaction energy (top row) and interaction force (bottom row) for our (left) and the OLS potential (right), for a 4-nm Cr precipitate. The position of the precipitate edge and maximal force between precipitate and dislocation are indicated by $R_{\rm P}$ and $F_{\rm MAX}$, respectively.

created around the precipitate) excess energy. The chemical component, E_S , was calculated statically, by shearing a precipitate in the (110) plane at its centre by the distance b, in a number of steps, each equal to $2R_P/b$, followed by relaxation. In this way, we mimic the shearing of the precipitate due to the advance of the screw dislocation. The shear modulus misfit component, E_{μ} , was calculated by taking the volume integral of the dislocation strain energy between the particle centred at $(h_x, h_y, 0)$ and a dislocation line parallel to the z-axis, following the expression [62]:

$$E_{\mu} = \frac{\mu_P - \mu_M}{\mu_M} \int_P \rho \cdot D_Z(\rho, \theta) \cdot e_M(\rho, \theta) \cdot d\rho \cdot d\theta.$$
(11)

Here, the subscripts *M* and *P* indicate, respectively, matrix and precipitate, $\rho = \sqrt{x^2 + y^2}$ and $\theta = \arctan(y/x)$ are the standard polar coordinates, $e(\rho, \theta) = \mu b^2/8\pi^2 \rho^2$ is the strain energy density and D_Z is expressed as

$$D_Z(\rho,\theta) = 2\sqrt{R_P^2 - (\rho\cos\theta - h_x)^2 - (\rho\sin\theta - h_y)^2}.$$
 (12)

In turn, the excess relaxation energy was calculated as $E_{xc} = E_I - (E_S + E_{\mu})$.

As a representative example, the three above determined components and their derivatives, calculated for a screw dislocation interacting with a 4-nm precipitate, are shown in Figure 11. The smallest contribution comes from the chemical component, whose curve is magnified 10 times for better observation while the contribution from the shear modulus misfit clearly dominates. As expected, its absolute contribution is the highest for OLS. Interestingly, the third term, i.e. the relaxation excess energy, appears to be approximately the same and negative for both potentials. In absolute terms, however, $E_{\rm xc}$ is higher for OLS.

7. Summary and concluding remarks

Based on the phase diagram, all potentials are capable of reproducing short-range order and α' precipitation. Our potential, however, seems to be the best compromise for thermodynamic modelling. It gives an asymmetric mixing enthalpy, does not stabilise unobserved intermetallic compounds (unlike OLS), provides (though underestimated) a significant amount of excess vibrational entropy (unlike CAR) and a reasonable agreement with the experimental miscibility gap, given that magnetic interactions are not explicitly included in the formalism.

Based on the description of substitutional defect interactions and the phase diagram, both OLS and our potential *a priori* seem to be the most appropriate choice to simulate thermal annealing in Fe–Cr alloys. Besides a proper reproduction of the thermodynamic limit, a reasonable reproduction of vacancy migration barriers, Cr–Cr and Cr–vacancy interactions assure that the correct kinetic path is followed using the correct mechanisms. All potentials provide a reasonable description of the Cr–Cr and Cr–vacancy interactions, but only the 2BM potentials provide reasonable values for vacancy migration barriers, with ours following the DFT trends the closest among the two.

Based on the description of interstitial and substitutional defect interactions and the phase diagram, both 2BM potentials *a priori* seem to be the most appropriate choice to simulate thermal annealing of Fe–Cr alloys under irradiation. Besides a proper description of thermal annealing, a correct stabilisation of interstitial complexes is needed to assure that the correct micro-structure is obtained under irradiation. Both 2BM potentials provide reasonable interstitial stabilities (ours in particular), while CAR poorly reproduces interstitial properties.

Based on the description of the $1/2 \langle 111 \rangle$ screw dislocation core structure (in bulk Fe and Cr) and its stable glide in a $\{110\}$ plane in bulk Fe, our potential *a priori* seems to be the most suitable choice for the simulation of precipitate–screw dislocation interaction. From the three potentials, ours is the only one reproducing the compact $1/2 \langle 111 \rangle$ screw dislocation core structure in bulk Cr. In addition, our static simulations have revealed an isotropic-to-degenerate core transition in every position near the precipitate with OLS that is not seen with our potential. Such a core modification may lead to different interaction mechanisms in dynamic simulations, such as cross slip.

In summary, we have shown that the developed Fe–Cr potential reached the goal of removing the main shortcomings of previously existing ones, while conserving their "good" properties. It should be noted, however, that it is not stiffened to the

screened Coulomb interaction and is, therefore, not suitable to simulate collision cascades in its present form.

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Appendix 1. Tabulation of the potential functions

Tabulations containing up to 5000 points are available online as supplementary material related to this paper or at: http://www.ctcms.nist.gov/potentials/ or upon request to the authors. Note that a tabulation of the s-embedding functions is not provided due to the singular slope in the origin, so that care should be exercised if used in tabular form. The parameters for the s-embedding functions are $A_1 = -0.217009784$, $A_2 = 0.388002579$ and $A_1 = -0.00977557632$, $A_2 = 0.374570104$ for Fe and Cr, respectively.

Appendix 2. Vacancy migration barriers

In this Appendix, the migration barriers for Fe–Vacancy and Cr–Vacancy exchange for different local chemical environments are provided. The latter are built by adding Cr atoms at the six nearest neighbour positions around the migration saddle point. The details are as reported in Table 4 next.

Table 4. Vacancy migration barrier (eV) for different local Cr configurations, with the first and second value denoting the forward and backward jump, respectively. In the figure, black circles are Fe, white circles are Cr, grey is the migrating atom, and the square represents the vacancy.

| Configuration | | DFT | This study | OLS | CAR |
|---------------|--------------------|-----------|------------|-----------|-----------|
| | $E_{ m m}^{ m Cr}$ | 0.57/0.57 | 0.57/0.57 | 0.56/0.56 | 0.90/0.90 |
| | $E_{ m m}^{ m Fe}$ | 0.64/0.64 | 0.63/0.63 | 0.64/0.64 | 0.63/0.63 |
| | $E_{ m m}^{ m Cr}$ | 0.72/0.54 | 0.63/0.57 | 0.61/0.53 | 0.97/0.87 |
| | $E_{ m m}^{ m Fe}$ | 0.66/0.60 | 0.65/0.63 | 0.67/0.63 | 0.64/0.62 |
| | $E_{ m m}^{ m Cr}$ | 0.79/0.52 | 0.66/0.58 | 0.62/0.51 | 1.00/0.85 |
| | $E_{ m m}^{ m Fe}$ | 0.73/0.57 | 0.67/0.63 | 0.68/0.61 | 0.65/0.61 |

(Continued)

| Configuration | | DFT | This study | OLS | CAR |
|---------------|--------------------|-----------|------------|-----------|-----------|
| | $E_{ m m}^{ m Cr}$ | 1.09/0.69 | 0.68/0.58 | 0.59/0.49 | 0.99/0.82 |
| | $E_{ m m}^{ m Fe}$ | 0.84/0.55 | 0.69/0.63 | 0.67/0.60 | 0.65/0.59 |
| | $E_{ m m}^{ m Cr}$ | 0.88/0.55 | 0.67/0.61 | 0.56/0.51 | 0.86/0.77 |
| | $E_{ m m}^{ m Fe}$ | 0.78/0.58 | 0.68/0.64 | 0.66/0.62 | 0.57/0.55 |
| | $E_{ m m}^{ m Cr}$ | 0.74/0.56 | 0.66/0.62 | 0.54/0.52 | 0.76/0.72 |
| | $E_{ m m}^{ m Fe}$ | 0.70/0.61 | 0.67/0.65 | 0.65/0.64 | 0.50/0.50 |
| | $E_{ m m}^{ m Cr}$ | 0.56/0.56 | 0.64/0.64 | 0.52/0.52 | 0.67/0.67 |
| | $E_{ m m}^{ m Fe}$ | 0.64/0.64 | 0.66/0.66 | 0.64/0.64 | 0.47/0.47 |

| Table | 4. | Continued. |
|-------|----|------------|
| | | |