

A machine learning interatomic potential for the structural properties of hematite: supplemental material

IMMEDIATE

1. EMPIRICAL POTENTIALS

A. Interface

The Interface Force Field (IFF) is a set of parameters and a methodology that is compatible with the functional forms of major existing force fields [1]. In this sense, there exist many variants for the IFF and in this work we have used the “IFF-CHARMM”, which refers to the use of IFF-derived parameters within a simulation engine that uses the CHARMM force field functional forms [2]. The interaction between species i and j is composed by a van der Waals term (Lennard-Jones) plus an electrostatic contribution, namely,

$$U_{ij} = 4\epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left(\frac{\sigma_{ij}}{r_{ij}} \right)^6 \right] + \frac{e^2}{4\pi\epsilon_0} \frac{z_i z_j}{r_{ij}^2}, \quad (\text{S1})$$

where $1/4\pi\epsilon_0$ is the usual Coulomb constant and r_{ij} is the center-to-center separation between atoms i and j . Charges and Lennard-Jones (LJ) parameters were derived by Kanhaiya *et al.* [3]. Charge numbers are $z_{\text{Fe}} = 1.74$ and $z_{\text{O}} = -1.16$ and $e = 1.602 \times 10^{-19}$ C is the elementary charge. The other parameters are summarized in Table S1.

B. Clay

The functional form of the Clay potential is identical to IFF’s: it is composed by non-bonded interactions only, a sum of a LJ term and a Coulombic contribution [see Eq. (S1)] [4]. Charge numbers for Fe and O atoms in the Clay parametrization are set to 1.575 and -1.05 , respectively. The LJ parameters are mistyped in its original reference (Ref. [4]) and in this work we have used their corrected values [5], shown in Table S2.

C. Tersoff

The potential we refer as Tersoff belongs to a class of analytical bond-order potentials (ABOP), with origins in Abell’s [6] discussion of bond order. Tersoff [7, 8] first developed these ideas into an empirical potential for covalent materials [9, 10]. The ABOP formalism closely follows Tersoff’s work but is extended in a style similar to Brenner’s [11]. Notably, this entire family of potentials for covalent systems is fundamentally connected to the embedded atom method (EAM) potentials [12, 13], used for metals [14].

In this work, we used a slightly modified version of Tersoff equations, fully described in Bygmästar *et al.* work, where parameters for the Fe-O interaction were also derived [15]. Parameters for Fe-Fe were developed by Müller *et al.* [16] and Bjorkas and Nordlund [17], while the O-O interaction parameters are from Erhart *et al.* [18].

Table S1. Parameters for the Interface Force Field, Eq. (S1), used in this work.

Pair	ϵ_{ij} (kcal/mol)	σ_{ij} (Å)
Fe-Fe	0.2	1.8085
O-O	0.2	3.1003
Fe-O	0.2	2.4544

Table S2. Parameters for Clay potential, Eq. (S1), used in this work.

Pair	ϵ_{ij} (kcal/mol)	σ_{ij} (Å)
Fe-Fe	9.0298×10^{-7}	4.9062
O-O	0.1554	3.1650
Fe-O	3.7459×10^{-8}	4.0356

D. Reax

The Reactive Force Field, in this work denoted as Reax, was developed by Adri van Duin, William A. Goddard III, and collaborators. It is a bond-order-based interatomic interaction designed to model chemical reactions within classical molecular dynamics simulations [19].

The Reax methodology is founded on a general relationship between bond distance and bond order, and subsequently between bond order and bond energy, which ensures the correct dissociation of bonds to separated atoms. The contributions from other valence terms, such as bond angles and torsions, are also formulated as a function of bond order, causing them to decay smoothly to zero during dissociation events. Non-bonded interactions are handled by Coulomb and Morse (van der Waals) potentials applied between all atom pairs, with no exclusions. To avoid nonphysical behavior at close proximity, these non-bonded interactions are shielded at short ranges, causing them to plateau at a constant value as the interatomic distance approaches zero.

For the complete mathematical description of the Reax Force Field, we invite the reader to check the aforementioned original references. For the Reax parameters used in this work we used the Shin *et. al* derivations [20].

E. Core-Shell

The core-shell model is a classical approach that extends the rigid ion Born model of solids [21] to include the effects of electronic polarizability [22]. A polarizable ion, typically the anion (O^{2-}), is modeled as a massive core (representing the nucleus and core electrons) connected to a massless shell (representing the valence electrons) by a harmonic spring [23]. The polarizability of the ion is thus determined by the charges of the core and shell and the stiffness of the spring constant connecting them. Cations are generally treated as unpolarizable point charges [23]. Table S3 details the specific charges assigned to each particle type.

Table S3. Particle charges for the Core-Shell model [23].

Particle	Charge (e)
Fe	+3.00
O _{core}	+0.21
O _{shell}	-2.21

Interatomic forces in the model are described by a combination of long-range and short-range potentials. The long-range electrostatic interactions are calculated between all cores and shells using a Coulomb potential. Short-range interactions are modeled with a pair-potential function acting between the shells of neighboring ions. For this model, a Buckingham potential is used, with the functional form:

$$U_{ij}(r_{ij}) = A_{ij} \exp(-r_{ij}/\rho_{ij}) - \frac{C_{ij}}{r_{ij}^6}. \quad (S2)$$

These short-range potentials are defined only for interactions involving shells, as detailed in Table S4. Core-core interactions are purely Coulombic.

Finally, the intramolecular interaction between an oxygen core and its own shell is represented by the aforementioned harmonic spring [23]. The spring constant for this core-shell (cs) bond is $k_{cs} = 27.4 \text{ eV}/\text{\AA}^2$. Note that for implementation in the simulation package LAMMPS (Imp)

Table S4. Core-shell parameters for the Buckingham contribution (Eq. S2) [23].

Pair	A_{ij} (eV)	ρ_{ij} (Å)	C_{ij} (eV·Å ⁶)
Fe-Fe	0.0	–	0.0
Fe-O _{core}	0.0	–	0.0
Fe-O _{shell}	1102.4	0.3299	0.0
O _{core} -O _{core}	0.0	–	0.0
O _{core} -O _{shell}	0.0	–	0.0
O _{shell} -O _{shell}	22764.0	0.1490	27.88

(through the harmonic bond style), this corresponds to a bond coefficient $k_{\text{Imp}} = 13.7 \text{ eV}/\text{\AA}^2$, since $k_{\text{Imp}} = k_{\text{CS}}/2$.

F. Surface Energy Calculation

The surface energies (γ) were determined following an established methodology based on slab models. The procedure first required establishing a reference bulk energy per formula unit (E_b). This was calculated from a $3 \times 3 \times 1$ supercell containing $n = 54 \text{ Fe}_2\text{O}_3$ units, constructed from a fully relaxed primitive cell. Subsequently, symmetric slabs were cleaved for the (0001) and (0112) surfaces, with respective surface areas (A) of $208.7 \text{ \AA}^2$ and $242.6 \text{ \AA}^2$. The slabs were modeled with a thickness of approximately $40.0 \text{ \AA}$ to ensure convergence of the surface properties. To prevent interactions between periodic images, a vacuum gap of $100 \text{ \AA}$ was added. After performing a full geometric relaxation of the slab atoms, the resulting total energy (E_s) was used to derive the surface energy as:

$$\gamma = \frac{E_s - nE_b}{2A}.$$

G. Relative Deviations for Calculated Quantities

Here we present tables containing the relative deviation for the structural and elastic properties calculated in this work, shown in tables 1-6 of the main text.

Table S5. Average deviations from experimental values (%) for lattice parameters of hematite (see Tab. 1 of main text). $\langle \sigma \rangle$ is the absolute average deviation [see Eq. (1) of the main text]

Parameter	Tersoff	Reax	CS	IFF	Clay	Fe-MLIP	DFT
a	2.70	-0.89	0.50	0.14	0.93	2.44	2.52
c	-7.01	-13.27	-2.98	-0.90	0.29	1.92	0.49
c/a	-9.44	-12.48	-3.44	-1.02	-0.62	-0.48	-1.28
$\langle \sigma \rangle$	6.38	8.88	2.31	0.69	0.61	1.61	1.43

2. ELASTIC TENSOR AND ELASTIC MODULI OF HEMATITE

The primitive unit cell of hematite belongs to the trigonal crystal system and to the $R\bar{3}c$ (nr. 167) space group. For this symmetry, the elastic tensor $\mathbf{C}$ can be expressed in matrix form (in Voigt notation) as

Table S6. Average deviations from experimental values (%) for elastic constants of hematite (see Tab. 2 of main text).

Parameter	Tersoff	Reax	CS	IFF	Clay	Fe-MLIP	DFT
C_{11}	2.02	68.67	43.64	17.45	-11.18	-16.35	-5.14
C_{33}	27.14	97.09	34.36	28.88	0.90	-11.60	-6.32
C_{44}	35.00	102.07	110.17	144.31	71.03	31.55	24.31
C_{12}	-1.68	-4.00	11.30	1.95	-27.89	-24.65	-8.59
C_{13}	-21.63	-55.35	-29.37	-8.50	-36.29	-44.75	-31.81
$ C_{14} $	142.31	865.38	325.00	-87.50	-43.27	11.54	-45.19
$\langle\sigma\rangle$	38.30	198.76	92.31	48.10	31.76	23.41	20.23

Table S7. Average deviations from experimental values (%) for elastic moduli and related quantities for hematite (see Tab. S10).

Parameter	Tersoff	Reax	CS	IFF	Clay	Fe-MLIP	DFT
Y	0.04	83.52	51.04	34.14	2.16	-12.75	-11.27
K	14.65	35.52	29.26	27.41	-6.50	-14.02	-0.24
G	-2.64	91.76	53.41	33.96	2.64	-13.30	-13.52
β	-14.29	-26.53	-22.45	-22.45	6.12	14.29	0.00
ν	-13.16	-36.84	-26.32	-21.05	-23.68	-18.42	-13.16
K/G	18.14	-29.20	-15.49	-4.87	-8.85	-0.44	15.49
H	-14.06	148.27	73.66	37.43	19.55	-3.81	-18.95
$\langle\sigma\rangle$	11.00	64.52	38.80	25.90	9.93	11.00	10.38

Table S8. Average deviations from experimental values (%) for lattice and elastic parameters of maghemite (γ -Fe₂O₃) (see Tab. 5 of main text).

Parameter	Tersoff	CS	IFF	Clay	Fe-MLIP
a	0.29	1.05	6.04	4.08	2.59
c	0.68	-0.60	-12.0	-1.65	3.19
c/a	0.40	-1.64	-17.0	-5.52	0.60
C_{11}	-59.7	8.88	34.91	-20.6	-28.2
C_{12}	-43.3	26.90	48.23	3.36	-4.06
C_{13}	-46.0	3.85	-16.5	-13.1	-8.83
C_{33}	-49.6	-42.6	-61.9	-74.6	-36.3
C_{44}	-76.2	-30.7	-30.8	-41.8	-34.8
C_{66}	-76.7	27.54	-15.8	-28.3	-30.0
K	-45.1	-6.62	-6.65	-44.3	-9.86
$\langle\sigma\rangle$	39.8	15.0	25.0	23.7	15.8

Table S9. Average deviations from experimental values (%) for lattice and elastic parameters of magnetite (Fe₃O₄) (see Tab. 6 of main text).

Parameter	Tersoff	Reax	Fe-MLIP
a	2.12	-3.31	2.82
C_{11}	-51.6	-6.63	-13.5
C_{12}	-36.4	-53.2	-13.1
C_{44}	-74.4	8.12	1.87
Y	-66.4	19.22	2.81
K	-42.7	-27.5	-11.1
G	-63.7	49.09	19.73
β	87.27	49.09	20.00
ν	25.81	-32.3	-6.45
K/G	163.7	-18.5	23.81
H	-76.0	147.6	31.11
$\langle\sigma\rangle$	53.6	37.7	13.3

$$\mathbf{C} = \begin{pmatrix} C_{11} & C_{12} & C_{13} & C_{14} & 0 & 0 \\ C_{12} & C_{11} & C_{13} & -C_{14} & 0 & 0 \\ C_{13} & C_{13} & C_{33} & 0 & 0 & 0 \\ C_{14} & -C_{14} & 0 & C_{44} & 0 & 0 \\ 0 & 0 & 0 & 0 & C_{44} & C_{14} \\ 0 & 0 & 0 & 0 & C_{14} & \frac{1}{2}(C_{11} - C_{12}) \end{pmatrix}.$$

The relationship between stress (σ_j) and strain (ε_i) follows Hooke's law, $\sigma_i = C_{ij}\varepsilon_j$, where C_{ij} are the elements of $\mathbf{C}$ [24]. Indices refer to the standard directions in strain and stress theory, as defined in Voigt notation [25], specifically $xx = 1$, $yy = 2$, $zz = 3$, $yz = 4$, $xz = 5$, and $xy = 6$. The inverse relationship is given by $\varepsilon_i = S_{ij}\sigma_j$, yields the compliance tensor. The explicit equations for S_{ij} in terms of C_{ij} are

$$\begin{aligned} S_{11} &= (C_{22}C_{33} - C_{23}^2)/\Delta \\ S_{22} &= (C_{11}C_{33} - C_{13}^2)/\Delta \\ S_{33} &= (C_{11}C_{22} - C_{12}^2)/\Delta \\ S_{12} &= (C_{13}C_{23} - C_{12}C_{33})/\Delta \\ S_{13} &= (C_{12}C_{23} - C_{13}C_{22})/\Delta \\ S_{23} &= (C_{12}C_{13} - C_{11}C_{23})/\Delta \\ S_{44} &= \frac{1}{C_{44}}, \quad S_{55} = \frac{1}{C_{55}}, \quad S_{66} = \frac{1}{C_{66}}, \end{aligned} \tag{S3}$$

where $\Delta = C_{11}C_{22}C_{33} + 2C_{12}C_{13}C_{23} - C_{11}C_{23}^2 - C_{22}C_{13}^2 - C_{33}C_{12}^2$.

For obtaining the Bulk (K) and Shear (G) moduli we used the Voigt-Reuss-Hill averaging scheme [26].

$$\begin{aligned} K &= \frac{K_v + K_r}{2} \\ G &= \frac{G_v + G_r}{2}, \end{aligned} \tag{S4}$$

where

$$\begin{aligned}
K_v &= \frac{f_1 + 2f_2}{9} \\
K_r &= \frac{1}{\tilde{f}_1 + 2\tilde{f}_2} \\
G_v &= \frac{f_1 - f_2 + 3f_3}{15} \\
G_r &= \frac{15}{4\tilde{f}_1 - 4\tilde{f}_2 - 3\tilde{f}_3}
\end{aligned}$$

are their corresponding Voigt (v) [25] and Reuss (r) [27] approximations, with auxiliary functions f_k and $\tilde{f}_k$ given by

$$\begin{aligned}
f_1 &= C_{11} + C_{22} + C_{33} \\
f_2 &= C_{12} + C_{13} + C_{23} \\
f_3 &= C_{44} + C_{55} + C_{66} \\
\tilde{f}_1 &= S_{11} + S_{22} + S_{33} \\
\tilde{f}_2 &= S_{12} + S_{13} + S_{23} \\
\tilde{f}_3 &= S_{44} + S_{55} + S_{66}.
\end{aligned}$$

The elements C_{ij} were obtained from simulations and S_{ij} are calculated through Eqs. (S3).

The isotropic Young's modulus (Y), Poisson's ratio (ν), and compressibility (β) can be derived from the bulk modulus (K) and shear modulus (G) – as shown in Eq. (S4) – using the standard relations:

$$\begin{aligned}
Y &= \frac{9KG}{3K + G} \\
\nu &= \frac{3K - 2G}{6K + 2G} \\
\beta &= \frac{1}{K}.
\end{aligned}$$

Table S10. Elastic moduli and related quantities for empirical potentials and our trained ML potential (Fe-MLIP) for hematite. DFT data are from Ref. [28] and experimental data is a calculated consensus from literature (see the Suppl. Mat. for details). Young's (Y), Bulk (K) and Shear (G) moduli and Vickers hardness (H) are given in GPa units, while the compressibility (β) in GPa^{-1} . Poisson (ν) and Pugh (K/G) ratios are dimensionless. Colors help to visualize overestimation (red) and underestimation (blue) in relation to experiments.

Parameter	Tersoff	Reax	CS	IFF	Clay	Fe-MLIP	DFT	Exp.
Y	236.2	433.3	356.6	316.7	241.2	206.0	209.5	236.1
K	236.3	279.3	266.4	262.6	192.7	177.2	205.6	206.1
G	88.6	174.5	139.6	121.9	93.4	78.9	78.7	91.0
β	0.0042	0.0036	0.0038	0.0038	0.0052	0.0056	0.0049	0.0049
ν	0.33	0.24	0.28	0.30	0.29	0.31	0.33	0.38
K/G	2.67	1.60	1.91	2.15	2.06	2.25	2.61	2.26
H	7.21	20.83	14.57	11.53	10.03	8.07	6.80	8.39

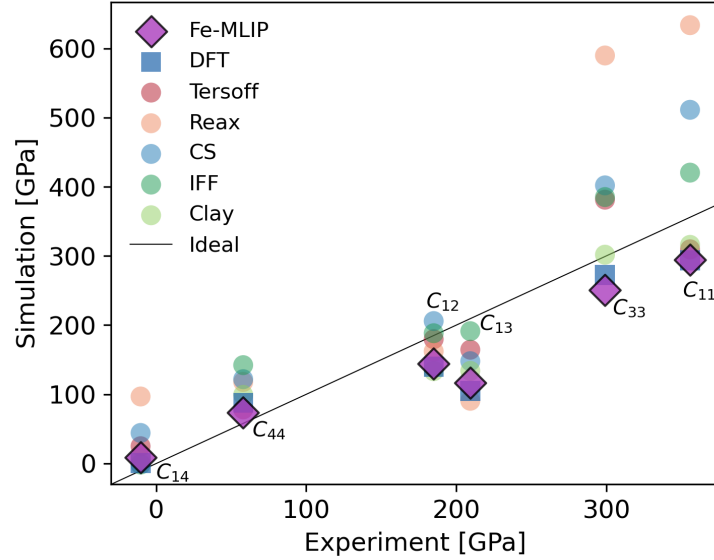


Fig. S1. Elastic tensor constants, C_{ij} for hematite calculated with different potentials. The continuous line (ideal) is given by Liebermann's experimental results [29].

Table S10 shows the bulk, K , Young, Y , and shear, G , moduli, the compressibility, β , Poisson's ratio, ν , Pugh ratio, G/K , and the Vickers hardness, H , for hematite.

Figure S1 shows a parity plot of the elastic tensor elements, C_{ij} , comparing the results of the potentials to experimental data (Ref. [29]).

3. THERMAL EXPANSION

The thermal expansion of hematite was calculated using the quasi-harmonic approximation (QHA) using the Phonopy code [30, 31]. Within the QHA, temperature effects are calculated including part of the anharmonic effects by the volume dependence of the (harmonic) phonon frequencies, which allows computing temperature-dependent thermodynamic properties from the corresponding free energy. The phononic contribution to the Helmholtz free energy is then given by [30]

$$F_{\text{ph}}^{\text{QHA}}(V, T) = \sum_q \frac{\hbar \omega_q(V)}{2} - k_B T \sum_q \ln(1 + n_q(V, T)), \quad (\text{S5})$$

where

$$n_q(V, T) = \frac{1}{\exp(\hbar \omega_q(V)/k_B T) - 1} \quad (\text{S6})$$

is the Bose-Einstein distribution. The total free energy of the system is

$$F(V, T) = E_{\text{latt}}(V) + F_{\text{ph}}^{\text{QHA}}(V, T), \quad (\text{S7})$$

where $E_{\text{latt}}(V)$ is the total energy given by DFT or potential calculations.

The frequencies $\omega_q(V)$ were calculated using a supercell with $3 \times 3 \times 3$ repetitions of the primitive unit cell of hematite for lattice parameters ranging from 5.50 Å to 5.68 Å, and a mesh of $21 \times 21 \times 21$ q -points. The results calculated with DFT and our Fe-MLIP are shown in Figures S2 and S3, respectively. From the minima of the free energy curves, $F(V, T)$ (Eq. S7), shown in the left panels of the Figs., the volume-temperature relation, $V(T)$, is obtained, shown in the middle panels. The thermal expansion coefficient is given by $(1/V)(\partial V/\partial T)$.

4. EXPERIMENTAL ELASTIC PROPERTIES CONSENSUS

A. Hematite

Surprisingly, considering the economic importance of hematite, we have found relatively few works in the literature addressing its elastic properties. In Table S11 we summarize the exper-

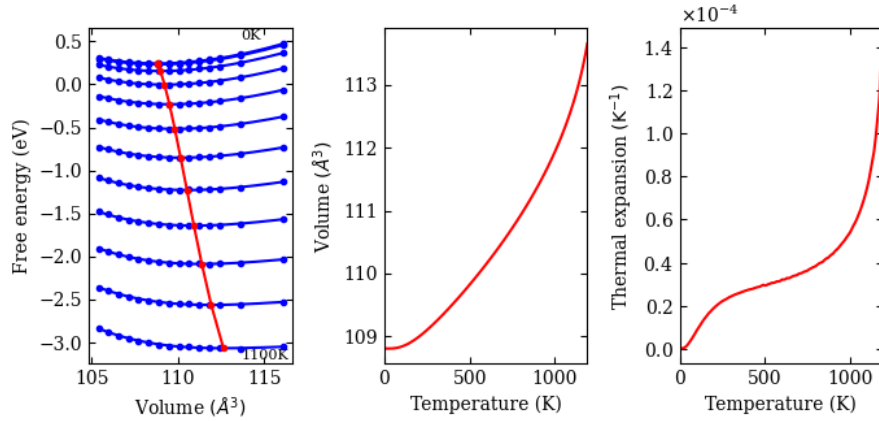


Fig. S2. Quasi harmonic thermal expansion of hematite calculated with DFT+U: (left panel) free energy vs. volume for various temperatures and their minima (red); (middle panel) volume vs. temperature; (right panel) thermal expansion coefficient.

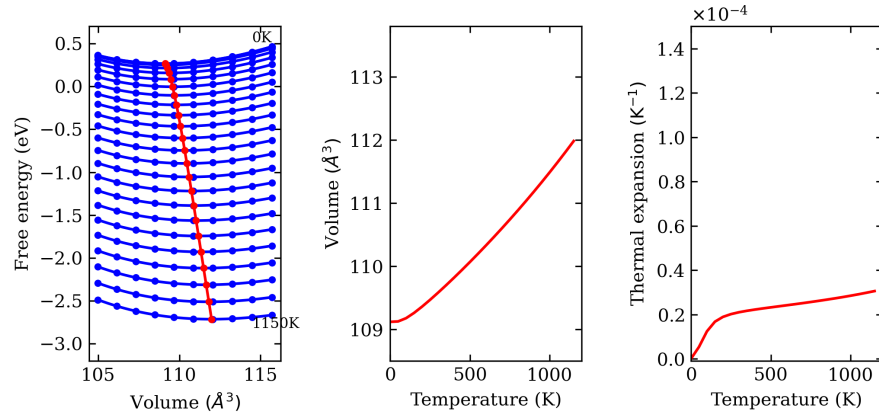


Fig. S3. Quasi harmonic thermal expansion of hematite calculated with our Fe-MLIP: (left panel) free energy vs. volume for various temperatures; (middle panel) volume vs. temperature; (right panel) thermal expansion coefficient.

imental results for the lattice constants and elastic moduli found in the literature. As can be seen, the property that has been measured the most is the isotropic bulk modulus (K), while the Young (E) and shear (G) moduli have been scarcely investigated. The experimental results shown in Table S11 were measured over a range of different conditions and samples. Thus, we use the bootstrap median estimation of the properties reported in the literature as a consensus for the reported experimental data, and use these estimates as a reference. We performed the median estimation by sampling the median with replacement ($n = 10000$) over the experimental data, and consider the uncertainty as two standard deviations, approximating a 95% confidence interval. This procedure for the median estimation is robust against outliers, and allows us to estimate the uncertainty associated with the different experiments as well.

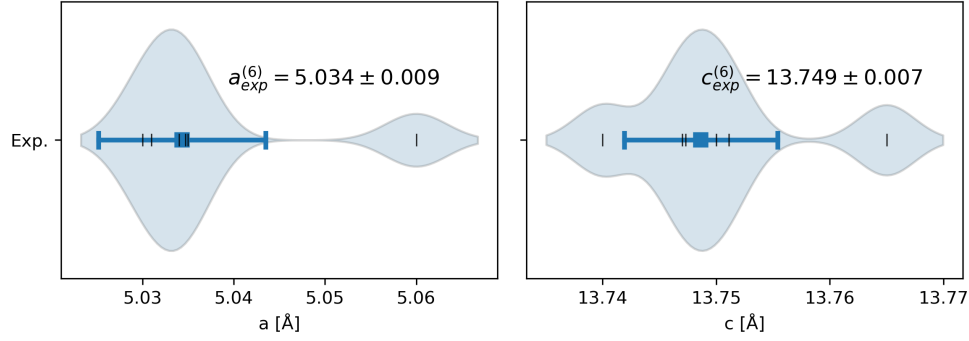


Fig. S4. Experimental lattice constants violin plot distributions for α -Fe₂O₃. The blue squares indicate the median estimation with two standard deviations error bars. The number in parenthesis in the labels reports the number of measurements.

Table S11. Experimental lattice constants, a and c (in Å), the elastic moduli for hematite (in GPa): namely, the bulk modulus and its volume derivative, K and K' , the Young modulus, E , the shear modulus, G , and the Poisson ratio, ν (adimensional). Values indicated with apostrophes (') were considered fixed by the authors. ^a) Calculated by us from the elastic tensor constants c_{ij} . ^b) Obtained by the relationships between elastic moduli.

Ref.	Material % purity	Conditions	a	c	K	K'	E	G	ν
[32] ^d					99.7		216.3	95.0	0.139
[33]		25.0°C, 1 bar			202.66	4.53	237.46 ^b	0.386	
[34]		298 K					260		
[35]	Synthetic	<30 kbar			231	4.0'			
		>30 kbar			178	4.0'			
[36]		25°C, 1 bar			199(6)	4.0'			
[37]		298 K, 1 bar	5.0347(4)	13.7473(15)	225(4)	4.0'			
[38]	High purity	298 K, 1 bar			246.1(4)	4.5(2)	179.98 ^b	65.3(1.5)	0.378 ^b
[39]					212		250.2 ^b	96	0.303 ^b
[40]	Powder	P=0	5.031(7)	13.765(7)	230(5)	3.5(6)			
[41]	Bulk	298 K, 1 bar	5.034(5)	13.747(2)	205(5)	4.0'			
	Nanocrystal				211(10)	4.0'			
[42]	Synthetic 99.998%				202.66(20)				
[43]	99.99%		5.049(5)	13.73(2)	206.1(4.8)	4.0'			
					202.1(3.8)	4.3(3)			
[44]	Bulk	298 K, 1 bar			300(30)				
	7 nm nanopart.				336(5)				
[45]		298 K, 1 bar	5.060(5)	13.75(2)	268(14)	4.0'			
[46]	Powder, 99.99%				201(4)	4.3(2)			
[47]	Powder, 99.99%				201	4.3'			
[48]	3% magnetite, ~3.2% poros.	298 K	5.0350(1)	13.7511(3)	184.0(9)		214.8(3)	82.00(7)	0.306
					201		228	87	0.311
[49]	99.999%	$P < 25.3$ GPa	5.0354(17)	13.7477(48)	207(3)	4.0'			
					199(8)	5.3(9)			
[50]			5.0993	13.8040	235.4(8)	3.29(25)	234.748 ^b	88.0	0.334 ^b

In Figures S4 and S5 we show violin plots for the distributions of the lattice constants and elastic moduli reported in the literature (shown in Table S11) along with the *consensus* estimation.

For the elastic tensor constants, C_{ij} , only a handful of works are available in the literature, shown in Table S12. In this case, it should be noted that we have found only three experimental measurements, with one of them (Voigt [32]) being substantially softer (lower) than the other two. This disparity is also clear for the lattice constants (max. values) shown in Fig. S4, and for the elastic moduli (lower values) shown in Fig. S5. In that regard, Voigt pointed out that the sample for hematite "was more scarcer and not entirely free of defects" (free translation from

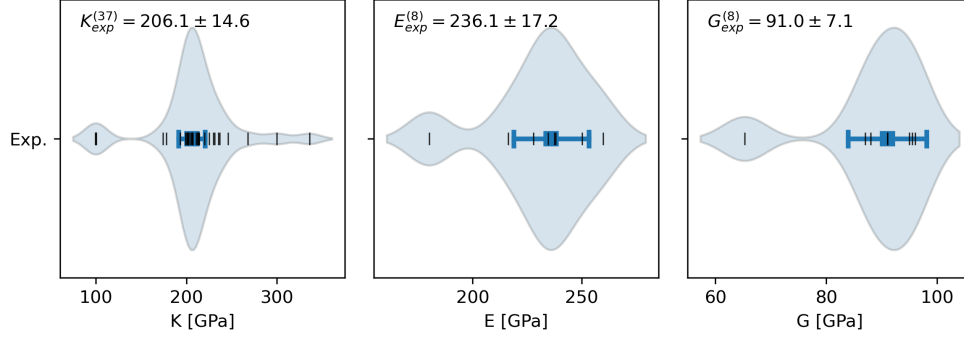


Fig. S5. Experimental elastic moduli (bulk, Young and shear modulus, K , E , and G , respectively) violin plot distributions for α -Fe₂O₃. The blue squares indicate the median estimation with two standard deviations error bars. The number in parenthesis in the labels reports the number of measurements.

Ref.	c_{11}	c_{33}	c_{44}	c_{12}	c_{13}	c_{14}
Voigt (1928) [32]*	247	232	87	56	16	-13
Birch (1966) [51]*	352.0	318.0				
Liebermann (1986) [38]* ^a	355.9	299.2	58.0	185.0	209.4	-10.4
Tomar (2004) [52] ^{+b}	273.54	184.05	106.6	56.8	16.52	-12.6
Tomar (2006) [53] ^{+b}	238.2	223.08	78.05	50.75	28.75	-13.12
Guo (2011) [54] [†]	319.6	294.4	80.1	125.6	104.5	5.5
Zhang (2020) [28] [†]	337.56	280.26	72.11	169.13	142.76	5.71

Table S12. Elastic tensor constants, C_{ij} [GPa], for hematite (Voigt notation). Symmetries: $c_{11} = c_{22}$, $c_{44} = c_{55}$, $c_{13} = c_{23}$, $c_{14} = c_{56} = -c_{24}$, $c_{66} = \frac{1}{2}(c_{11} - c_{12})$. *) Experimental results. [†]) Theoretical results. ^a) Based in magnetic field extrapolation $H \rightarrow 0$, except for c_{11} and c_{33} . ^b) Fitted to Voigt's data [32].

Ref. [32]). Other authors agreed that Voigt's sample was questionable [33, 51]. Despite this, these results were used as a reference in later theoretical works [52, 53]. In this work we use the values obtained by Liebermann [38] as reference for the elastic tensor constants.

For the Vicker's hardness, we used the average of the results obtained by Bhaumik [55], of 8.58 GPa (at 100 g load, based on literature standards [56]), and the result of 8.2 GPa from Chicot *et al.* [57], resulting in an average value of 8.39 GPa.

B. Magnetite

Consensus data for magnetite is shown in the Tab. S13.

5. RAMAN AND INFRARED ACTIVE MODES' FREQUENCIES

Ref.	Conditions	a	c	K	K'	E	G	ν
Kataoka (1984) [58]						180		0.35
Smasonov (1973) [59]	100 GPa, 273 K			181.8 ^a				
Straffelini (2002) [60]	32-43°C					210		0.3
Seo (2001) [61]	Room					174		
Asada (1983) [62]						141		0.33
Hearmon (1984) [63]						230.33	91.3	
Mao (1974) [64]				185				
				175				
				183				
				159				
				161				
				170				
				160				
				177				
				139				
				183				
Reichmann (2004) [65]	Ambient	8.39639(14)		180(1)	5.2(4)			
Lin (2014) [66]				186(3)	4'		66.3(1.8)	

Table S13. Experimental lattice constants and elastic moduli (units for length and pressure: Å, GPa) for hematite. ^a) Calculated by the inverse of the compressibility. Values indicated with apostrophes (') were considered fixed by the authors.

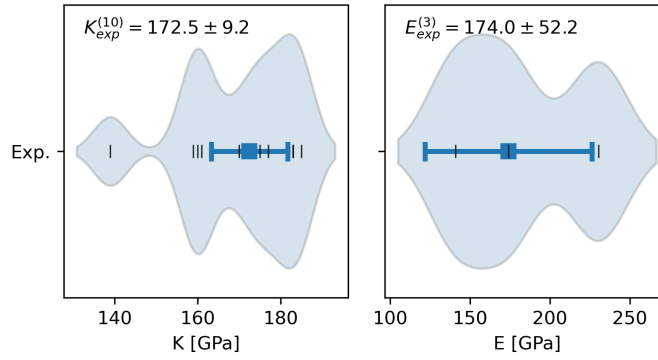


Fig. S6. Experimental elastic moduli (bulk and Young, K and E , respectively) violin plot distributions for magnetite. The blue squares indicate the median estimation with two standard deviations error bars. The number in parenthesis in the labels reports the number of measurements.

Ref.	c_{11}	c_{44}	c_{12}
[67]	272.5	97.1	106.0
[65]	260.5(1.0)	63.3(1.5)	148.3(3.0)
[66]	240.6(4.9)	68.3(1.6)	113.8(5.9)

Table S14. Elastic tensor constants, c_{ij} [GPa], for magnetite (Voigt notation).

Fe-MLIP			Clay			Reax			Tersoff			IFF			Exp.	
n	Sym.	ω	n	Sym.	ω	n	Sym.	ω	n	Sym.	ω	n	Sym.	ω	Sym.	ω_{avg}
7	Eg	219.4	7	A1g	256.5	5	Eg	317.3	7	Eg	172.1	7	A1g	302.7	A1g	226.3 ± 2.5
9	A1g	221.8	8	Eg	272.7	12	Eg	403.5	9	Eg	215.8	8	Eg	324.7	Eg	245.7 ± 3.1
11	Eg	267.9	10	Eg	281.1	15	A1g	432.2	11	A1g	227.1	10	Eg	332.7	Eg	292.7 ± 2.5
15	Eg	286.9	14	Eg	325.0	16	Eg	446.8	16	Eg	433.1	14	Eg	375.1	Eg	299.0 ± 2.6
19	Eg	377.9	18	Eg	437.4	18	Eg	476.3	21	Eg	492.6	18	Eg	523.0	Eg	411.7 ± 3.2
26	A1g	481.4	26	A1g	612.6	25	Eg	587.0	26	Eg	628.4	26	A1g	712.8	A1g	498.7 ± 2.3
28	Eg	548.9	28	Eg	691.5	27	A1g	606.2	29	A1g	732.8	28	Eg	800.6	Eg	612.0 ± 3.0

Table S15. Calculated Raman active modes: number of the mode (n), irreducible representation symmetry, and frequency (ω [cm^{-1}]) for active Raman modes. Experimental average (ω_{avg}) taken from the data of Refs. [42, 68, 69]

Fe-MLIP			Clay			Reax			Tersoff			IFF			Exp.	
n	Sym.	ω	n	Sym.	ω	n	Sym.	ω	n	Sym.	ω	n	Sym.	ω	Sym.	ω_{avg}
5	Eu	212.2	5	Eu	236.3	10	A2u	411.5	5	Eu	169.7	5	Eu	292.3	Eu	224.0 ± 7.0
10	A2u	233.1	12	A2u	290.6	10	Eu	390.5	14	Eu	365.4	12	A2u	350.1	Eu	292.7 ± 7.0
13	Eu	286.4	16	Eu	356.0	8	Eu	345.9	13	A2u	338.1	16	Eu	426.7	A2u	300.0 ± 1.4
21	Eu	385.6	21	Eu	492.6	20	Eu	486.5	18	Eu	453.5	21	Eu	594.2	Eu	436.5 ± 2.9
23	A2u	432.0	23	A2u	517.4	23	A2u	509.2	23	A2u	519.8	23	A2u	618.3	Eu	523.0 ± 4.1
24	Eu	456.3	24	Eu	585.8	28	Eu	618.1	24	Eu	621.8	24	Eu	673.6	A2u	525.7 ± 0.6

Table S16. Calculated IR active modes: number of the mode (n), irreducible representation symmetry, and frequency (ω [cm^{-1}]) for active IR modes. Experimental average (ω_{avg}) taken from the data of Refs. [68, 70–72]

6. TRAINING

The configurations that comprised our dataset, used to train our potential, were sampled by molecular dynamics simulations with temperatures ranging from 100 to 2000 K and pressures from 1 to 100 atm. We also generated configurations by applying strains of $\pm 1\%$ and $\pm 3\%$ in each direction (x , y , and z), and combined directions (xy , yz , xz , and xyz) concomitantly with random displacements with standard deviations of 0.01, 0.02, and 0.03 Å. After sampling this first set of configurations, we trained a preliminary GNN potential from which we calculated the thermal expansion coefficient in the quasi-harmonic approximation, and random thermal displacements for the atoms at temperatures from 300 to 2100 K (with volumes corresponding to each given temperature) using the Phonopy code [31]. Figure S7 shows the energy-pressure diagram of the configurations of our data.

We trained 4 different message-passing graph neural networks (MPGNN) with different initial random weights, in order to use the models in a committee by averaging them. The node states of the GNNs were represented by 64 channels up to $L = 1$ equivariant features. Two message passing interactions were used, with correlation order $\eta_c = 3$. A hybrid scheme that smoothly interpolates the Ziegler-Biersack-Littmark (ZBL) screened nuclear repulsion potential with the GNN models was employed to avoid divergences in predicted forces and energies if atoms get too close to one another [73]. A Huber loss function was used for energies, forces, and stresses in

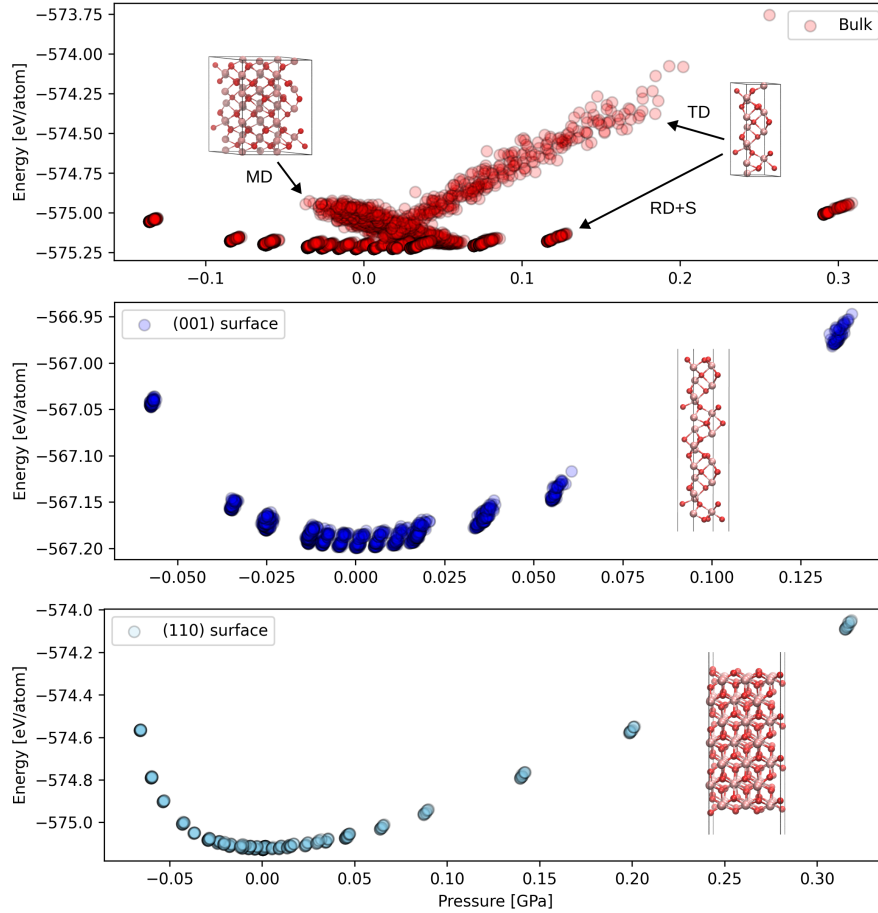


Fig. S7. Energy vs. pressure diagrams of the data sets for bulk configurations (top), (001) surface configurations (middle), and (110) surface configurations (bottom). The pictures show the supercell geometries of the configurations. The arrows indicate configurations generated by molecular dynamics (MD), random displacements and strains (RD+S), and random thermal displacements (TD).

a two-phase optimization. First, we applied the exponential moving average (EMA) weight decay with weight $\eta_{EMA} = 0.95$, and loss function component weights of $w_E^{EMA} = 1$, $w_F^{EMA} = 100$, and $w_\sigma^{EMA} = 1000$ for energies, forces and stresses, respectively. Then, we switch to stochastic weight averaging (SWA) with loss function component weights of $w_E^{SWA} = 100$, $w_F^{SWA} = 1$, and $w_\sigma^{SWA} = 100$ for another maximum 150 epochs with a learning rate of 10^{-3} , using early stopping if the models do not improve for 20 epochs. The schedulefree optimizer [74] was used during optimization.

Figure S8 shows the root mean square errors for the validation data set during training of energies, forces, and stresses. The vertical lines indicate the switching from EMA to SWA at epoch 150, and the horizontal lines are typical accuracy targets for interatomic potentials, 1 meV/atom for forces, 40 meV/Å for forces and 0.6 meV/Å³ (~ 0.1 GPa) for stresses. All models are well converged.

Figures S9-S12 show the parity graphs and the error distributions for the 4 individual trained Fe-MLIP models. All of the models display good performance on the test set, with RMS errors lesser than 1.2 meV/atom for energies, 32 meV/Å for forces, and 0.72 meV/Å³ for stresses.

The four different models shown above can be averaged in a committee. This procedure has two main advantages: first, there is some error cancellation between the models, improving robustness and generalization; and secondly, the dispersion of the models predictions (the standard deviation, for instance) can be used as a measure of the adequacy of the potential for a given configuration, which allows the assessment of the quality of the potential for different thermodynamic conditions, for example [75]. Figure S13 shows the parity graphs and the error distributions for the committee Fe-MLIP models. The RMS errors of the committee for energies, forces and stresses are 0.75 meV/atom, 21 meV/Å, and 0.33 meV/Å³, respectively.

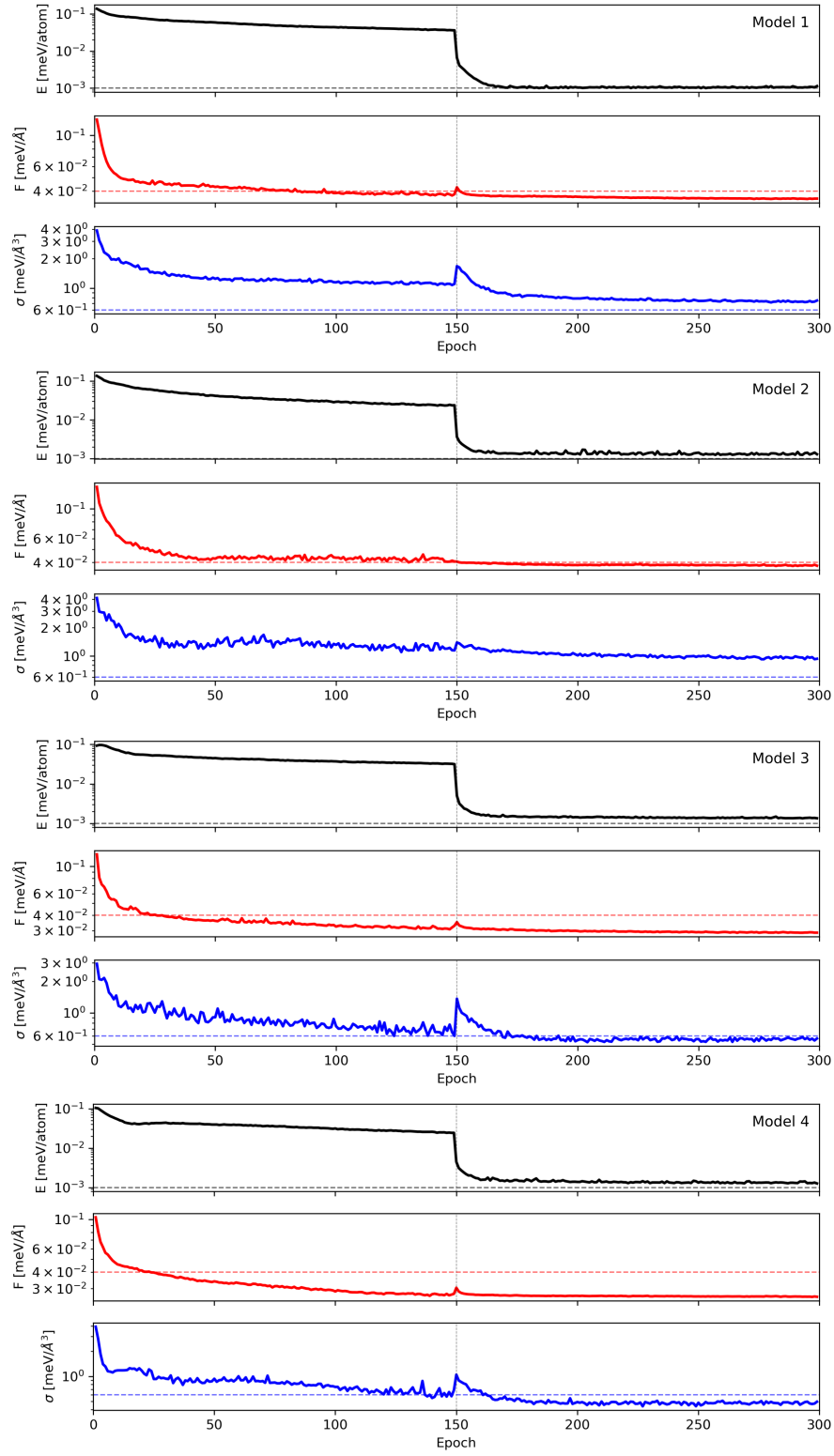


Fig. S8. Root mean square errors for the validation data set of energies E , forces F , and stresses σ . The vertical traced line indicates the switch from EMA to SWA (see text).

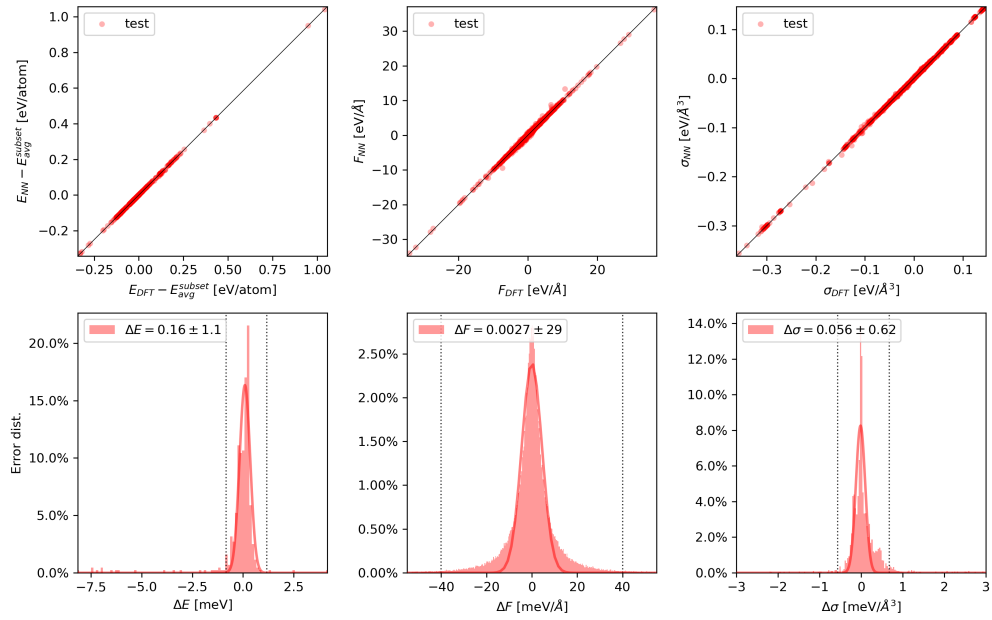


Fig. S9. Model 1 Fe-MLIP potential. Parity graph (top) and error distributions (bottom) for energies (left), forces (center), and stresses (right) calculated on the test set. The dotted lines indicate the ranges ± 1 meV/atom, ± 1 meV/Å, and ± 0.624 meV/Å³ (0.1 GPa).

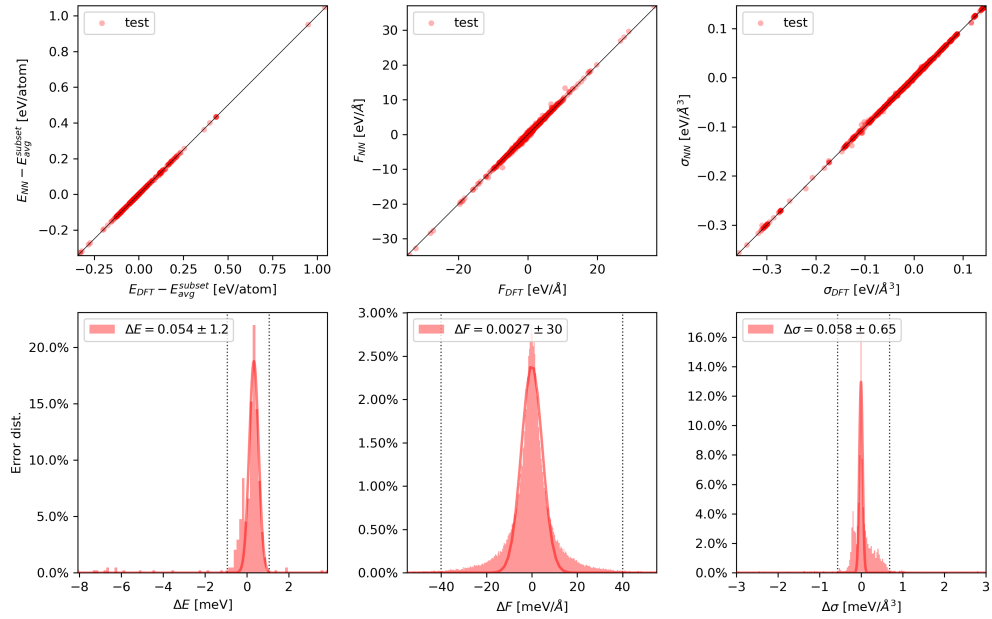


Fig. S10. Model 2 Fe-MLIP potential. Parity graph (top) and error distributions (bottom) for energies (left), forces (center), and stresses (right) calculated on the test set. The dotted lines indicate the ranges ± 1 meV/atom, ± 1 meV/Å, and ± 0.624 meV/Å³ (0.1 GPa).

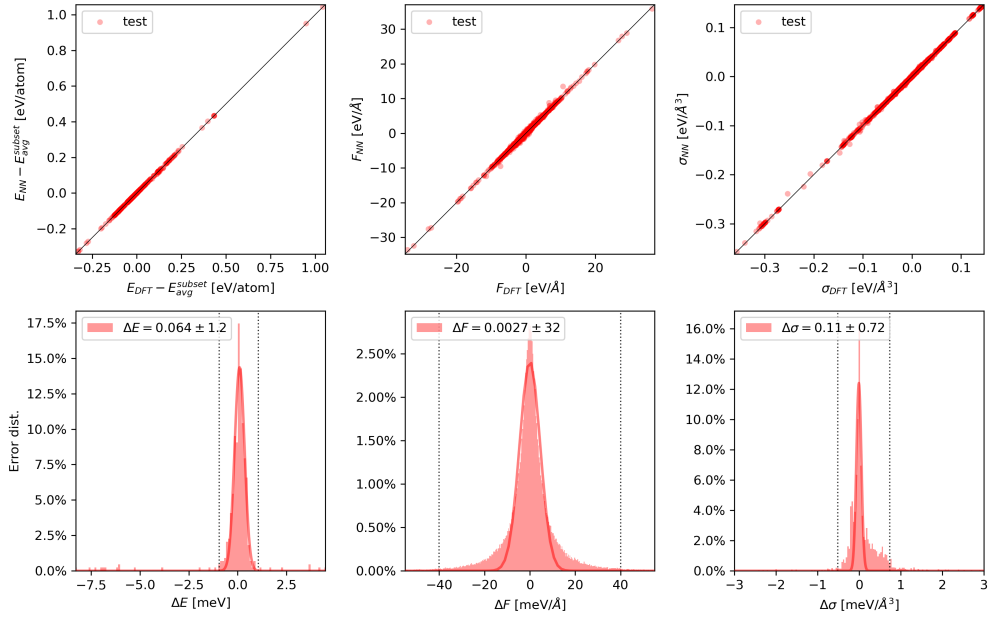


Fig. S11. Model 3 Fe-MLIP potential. Parity graph (top) and error distributions (bottom) for energies (left), forces (center), and stresses (right) calculated on the test set. The dotted lines indicate the ranges ± 1 meV/atom, ± 1 meV/Å, and ± 0.624 meV/Å³ (0.1 GPa).

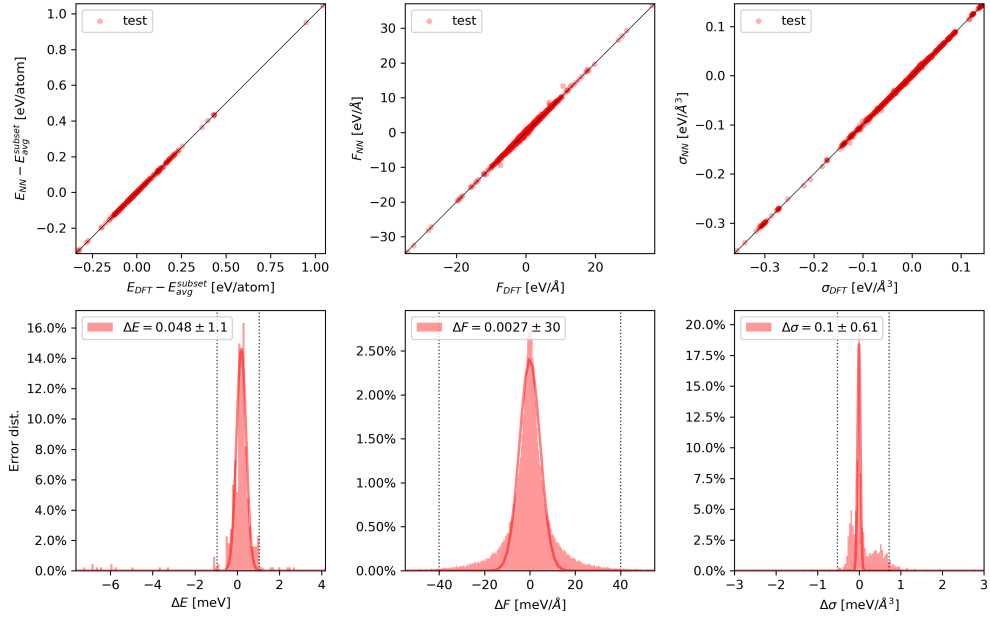


Fig. S12. Model 4 Fe-MLIP potential. Parity graph (top) and error distributions (bottom) for energies (left), forces (center), and stresses (right) calculated on the test set. The dotted lines indicate the ranges ± 1 meV/atom, ± 1 meV/Å, and ± 0.624 meV/Å³ (0.1 GPa).

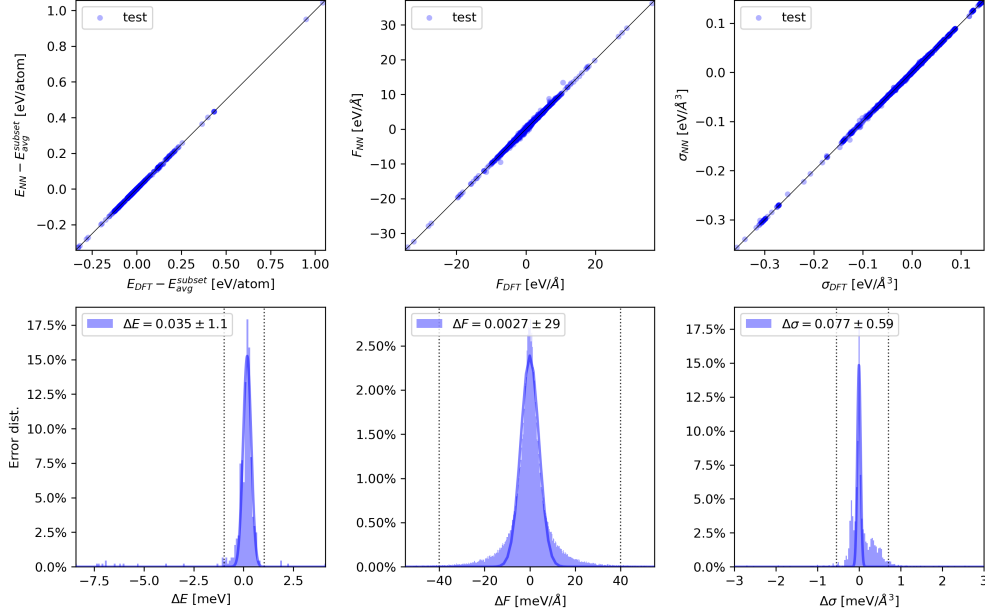


Fig. S13. Committee (4 models) Fe-MLIP potential. Parity graph (top) and error distributions (bottom) for energies (left), forces (center), and stresses (right) calculated on the test set. The dotted lines indicate the ranges ± 1 meV/atom, ± 1 meV/Å, and ± 0.624 meV/Å³ (0.1 GPa).

A. Hyperparameter testing

Table S17 shows the scaling of the Fe-MLIP with different cutoff radius, r_{cut} . Although the cutoff radius does not impact number of parameters of the model, the calculation of the atomic basis features in the MACE descriptor scale linearly with the number of neighbors, which is determined by r_{cut} (see Eqs. 8-9 in Ref. [76]). Considering constant density, the total number of neighbors scales with r_{cut}^3 . Therefore it is expected that the time, as well as the memory, required to evaluate the MACE descriptor to scale with $\sim \mathcal{O}(r_{cut}^3)$.

r_{cut}	E_{RMSE} (meV)	F_{RMSE} (meV/Å)	σ_{RMSE} (meV/Å ³)	Δt_{epoch}^{train} (s)
4.0	1.08	28.2	0.585	48.1 (1×)
5.0	1.13	28.7	0.595	77.9 (1.6×)
6.0	1.01	22.3	0.491	129.3 (2.7×)

Table S17. Impact of cutoff radius on the Fe-MLIP: RMS errors of energies, E , forces, F , stresses, σ , and time for training (per epoch). The training was carried out on a Nvidia GeForce RTX 4070 GPU. The numbers in parenthesis indicate the slowdown relative to $r_{cut} = 4.0$.

The parameter L_{max} determines the rank of the tensors used in the internal representations of the MACE GNN, or, in other words, the order of equivariance of the hidden nodes and message features carried through message passing ($L_{max} = 0$ amounts to only scalar-like features, while $L_{max} = 2$, to scalar-, vector-, and quadrupole-like features). Table S18 shows the influence of L_{max} on accuracy and computational costs.

The parameters $r_{cut} = 5.0$ and $L_{max} = 1$ show a good compromise between accuracy and computational efficiency.

B. Benchmarks

We have evaluated the computational performance of NPT molecular dynamics simulations performed with DFT, and the potentials Fe-MIP, Clay, and Reax. The Fe-MLIP calculations have been carried out on a Intel i7-14700KF and a Nvidia RTX 4070 GPU; and the rest, on a AMD Ryzen

L_{max}	E_{RMSE} (meV)	F_{RMSE} (meV/Å)	σ_{RMSE} (meV/Å ³)	Δt_{epoch}^{train} (s)	N_{params}
0	1.12	28.3	0.619	27.9 (1×)	124112 (1×)
1	1.13	28.7	0.595	77.9 (2.8×)	188880 (1.5×)
2	1.07	26.2	0.565	160.8 (5.8×)	263760 (2.1×)

Table S18. Impact of L_{max} on the Fe-MLIP: RMS errors of energies, E , forces, F , stresses, σ , time for training (per epoch), and number of MLIP parameters. The training was carried out on a Nvidia GeForce RTX 4070 GPU. The numbers in parenthesis indicate growth relative to $L_{max} = 0$.

9 7950X3D CPU using 16 cores. The MD simulations have been carried using SIESTA and the LAMMPS code.

Figure S14 shows the walltime per MD step versus the number of atoms for DFT, and the potentials Fe-MIP, Clay, and Reax. The Fe-MLIP has a performance on par with the empirical potentials Clay and Reax. Evaluation of the Fe-MLIP on a GPU is paramount to its computational efficiency, with a speedup of roughly one order of magnitude in comparison to its evaluation on a CPU.

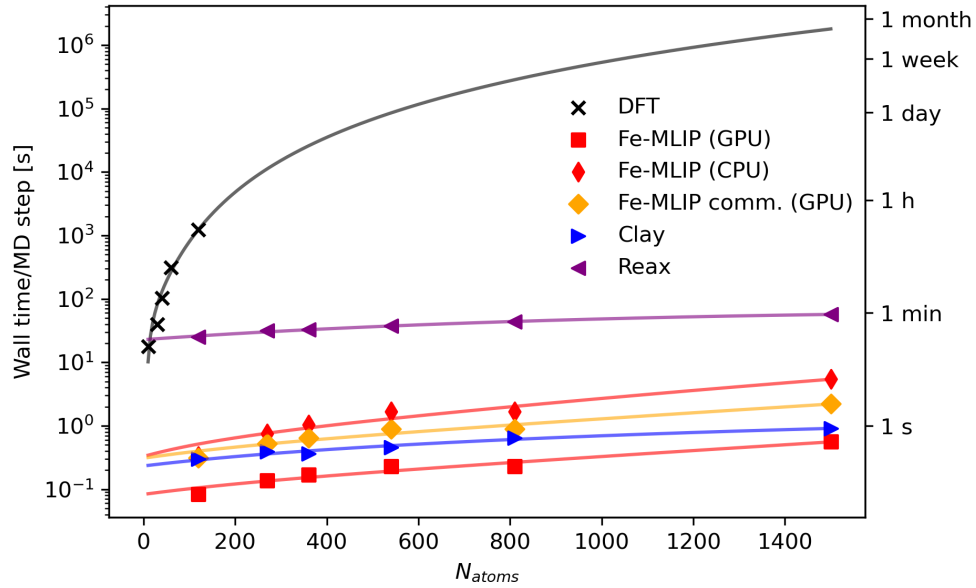


Fig. S14. Wall time per MD step versus number of atoms for DFT, Fe-MLIP (for a committee of 4 models, and for 1 model on CPU and GPU), Clay and Reax NPT simulations. The lines are cubic fits to the points.

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