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Magnetite Fe$_3$O$_4$ (111) Surfaces: Impact of Defects on Structure, Stability, and Electronic Properties

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Abstract

We present a comprehensive investigation, via first-principles density functional theory (DFT) calculations, of various surface terminations of magnetite, Fe₃O₄ (111), a major iron oxide which has also a number of applications in electronics and spintronics. We compare the thermodynamic stability and electronic structure among the different surfaces terminations. Interestingly, we find that surfaces modified with point defects and adatoms are close in surface energy and in the oxygen-rich and oxygen-poor regimes can be more stable than bulk-like terminations. These surfaces show different surface chemistry and electronic structures as well as distinctive spin polarization features near the Fermi level with regard to those previously considered in the literature. Our studies provide an atomic level insight for magnetite surfaces, which is a necessary step to understanding their interfaces with organic layers in OLED and spintronic devices.

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I. Introduction

Iron oxides, particularly magnetite (Fe$_3$O$_4$), have attracted interest in numerous fields including environmental remediation, biophysics, electrochemistry, sensors, and catalysis. Fe$_3$O$_4$ has also been exploited in organic light-emitting diodes (OLEDs). For example, magnetite electrodes can prevent the short-circuit problems seen in conventional top metallic electrodes as the diffusivity of the oxide species is much lower than its metallic counterparts, and have been successfully deposited on top of tris (8-hydroxyquinoline) aluminum(III) (Alq$_3$). Moreover, the use of a thin Fe$_3$O$_4$ layer on indium-tin-oxide (ITO) and Ag anodes has been shown to enhance hole injection and thus reduce turn-on voltage and enhance luminance and current density in devices.

A major potential application of this iron oxide is as an electrode in spintronic devices where spin functionality is built into hybrid organic devices. While there have been many materials considered so far as a source of spin injection, including ferromagnetic (FM) metals, dilute magnetic semiconductors, and Heusler alloys, half-metallic ferro- (or ferri-) magnetic oxides can produce a very high magnetoresistive (MR) response. In particular, advantages for using magnetite as an electrode in such devices include its high degree of spin polarization at the Fermi level, high Curie temperature (850 K), and the ability to grow high-quality films at room temperature, although device performance remains underwhelming. As the performance of these devices is dictated by the electronic structure immediately at the interface, a fundamental knowledge of the surface chemistry is required. For a spin field-effect transistor, the spin— injection and detection efficiencies are required to be 99.9995% to achieve the $10^5$ on-off ratio needed in mainstream applications. In the case of surface electronic states on a clean
Fe₃O₄ (001) surface, from 65%\(^{18}\) to less than 5% of electron-spin polarization at the Fermi-level has been observed.\(^{19-20}\) Thus, even small contributions to the electrode’s electronic structure from defects are of essential importance. However, the study of hybrid interfaces between magnetite and organic layers remains in its infancy. The electronic structure of magnetite varies qualitatively depending on the surface orientation and termination.\(^{21-22}\) Additional complexity is introduced as the control of the stoichiometry is highly dependent on the preparation conditions.\(^{23}\)

Bulk magnetite is a ferrimagnet with a cubic inverse spinel structure, with a chemical formula often written as \([\text{Fe}^{3+}]_A[\text{Fe}^{3+},\text{Fe}^{2+}]_B\text{O}_4\). This formula indicates that the tetrahedral sites denoted as A are occupied by ferric ions while octahedral sites denoted as B contain an equal number of ferric and ferrous ions. In magnetite, the tetrahedral and octahedral sites form two magnetic sublattices with the spin moments on the A sublattice antiparallel to those on the B sublattice. The proposed electronic structure of the octahedral \(\text{Fe}^{2+}\) cations corresponds to a situation where an extra electron resides in the lowest unoccupied \(t_{2g}\) orbital located at the Fermi level. The high electrical conductivity of \(2 \times 10^2\ \text{S/cm}\)\(^{24}\) in the thermodynamic standard state can be understood as resulting from rapid electron hopping processes between \(\text{Fe}^{2+}\) and \(\text{Fe}^{3+}\) ions. However, upon cooling below \(\sim 121\ K (T_v)\), magnetite undergoes a Verwey phase transition,\(^{25}\) with the electronic conductivity decreasing by two orders of magnitude and an associated opening of the optical gap.\(^{26}\) This transition is also accompanied by a structural distortion from the room temperature cubic system.\(^{27-29}\)

Several surface faces of magnetite have been prepared by varying the preparation conditions, including (100), (111), (110), (311), (331), and (511).\(^{30-34}\) Some of these
facets have been thoroughly investigated previously, for example, the (100) surface,\textsuperscript{30,35-36} which has recently been shown to have \((\sqrt{2} \times \sqrt{2})R45^\circ\)-reconstructed B termination with assistance of iron vacancies and interstitials in the subsurface.\textsuperscript{37} In the present study, we have chosen to focus on the Fe\textsubscript{3}O\textsubscript{4} (111) plane since: (i) it is the predominant natural cleavage plane; and (ii) it is the plane compatible with the hematite (\(\alpha\)-Fe\textsubscript{2}O\textsubscript{3}) (0001) surface that is regarded thermodynamically as the most stable at ambient conditions. However, \(\alpha\)-Fe\textsubscript{2}O\textsubscript{3} does not grow epitaxially on Fe\textsubscript{3}O\textsubscript{4}(111), the cause of which remains a topic of study.\textsuperscript{38} Additionally, in comparison to the other natural growth faces of Fe\textsubscript{3}O\textsubscript{4}, Yu \textit{et al.}\textsuperscript{39} found a higher stability of the (111) surface within a DFT+\(U\) scheme.

Here, our goal is to present a comprehensive DFT investigation, using generalized gradient approximation (GGA)+\(U\) approach, to describe the electronic and magnetic features of room-temperature magnetite surfaces. This understanding represents a necessary step prior to extending our calculations to interfaces with organic layers seen in OLED and spintronic devices. The calculations presented here provide a thorough analysis of the various magnetite (111) surface terminations in terms of thermodynamic stabilities, surface electronic structures, and surface chemical properties. Our results underline the possible stabilization of the Fe\textsubscript{3}O\textsubscript{4} (111) surface via introduction of point defects and adatoms. Importantly, our work demonstrates that four modified surfaces not considered in earlier theoretical studies show comparable surface stabilities to bulk truncated terminations but with significantly different surface chemistry and electronic/magnetic features. They thus appear to be of critical importance when developing atomistic models for magnetite interfaces.
II. Methodology

a) Terminations examined

The magnetite crystal contains layers of either only iron cations or oxygen anions along the [111] direction with the stacking sequence: O₁-\text{Fe}_{\text{oct1}}-O₂-\text{Fe}_{\text{tet2}}-\text{Fe}_{\text{oct2}}-\text{Fe}_{\text{tet1}}, as displayed in Figure 1; this leads to six unique atomic planes, where we follow the conventional notations to refer to each of the six bulk terminations.²³ The four iron terminations can present two different Fe sublayers; one is a dense monolayer (3/4ML) with all octahedrally coordinated Fe atoms (Fe\text{oct1}) on the same plane, while the other sublayer contains three distinct low-density monolayers (1/4ML) involving both octahedrally and tetrahedrally coordinated Fe atoms (Fe\text{tet2}-Fe\text{oct2}-Fe\text{tet1}). Those two iron sublayers are alternating between nearly close-packed oxygen stacking layers (1ML) that present a slight buckling. As the number and charge of the iron and oxygen ions on the surface differ, Fe₃O₄ (111) can be classified as a type-3 surface in the Tasker classification scheme,⁴⁰ with diverging electrostatic surface energy due to the presence of a non-zero dipole moment on all the repeat units throughout the material. In order to decrease the internal polarity perpendicular to the surface, major ionic relaxation and electron redistribution as well as surface reconstruction are expected;²² in spite of multiple studies involving various surface science techniques and computational approaches, the atomistic details of the Fe₃O₄ (111) surface are still a matter of debate as the exposed surface depends strongly on the preparation conditions and chemical environment. As the spin-polarization, resistivity, and Verwey transition are dependent
on the exact stoichiometry of thin films, development of an atomic-level description is vital.

![Figure 1](image)

**Figure 1** (a) Top view and (b) side view of the symmetric Fe$_3$O$_4$ (111) slab model. The stacking sequence, labeled O$_1$-Fe$_{\text{oct}1}$-O$_2$-Fe$_{\text{tet}2}$-Fe$_{\text{oct}2}$-Fe$_{\text{tet}1}$, is shown in panel (b). Fe atoms are colored blue, and oxygen ions are colored red.

At this stage, it is useful to recall a number of previous observations regarding magnetite surfaces. Scanning tunneling microscopy (STM) study of Fe$_3$O$_4$ (111) annealed at 1173 K in $2 \times 10^{-7}$ mbar O$_2$ reported a coexisting surface termination composed of a hexagonal array of 6.1 Å and a honeycomb pattern separated by 3.6 Å. The authors assigned the second pattern to termination to 2/4 ML of Fe atoms which is predicted to be more stable than the former, 3/4 ML capped by an O atom. In a later paper, the same authors reexamined the surface structure of single crystal UHV-prepared Fe$_3$O$_4$ (111) and concluded that the surface mainly consists of Fe$_{\text{tet}1}$, Fe$_{\text{oct}2}$,
and ferryl terminations.\textsuperscript{42} Full-potential density functional theory calculations by Zhu \textit{et al.}\textsuperscript{43} show that the Fe$_{\text{oct2}}$ termination is energetically more favorable using a local density approximation (LDA)+$U$ approach; their result is consistent with that from \textit{ab initio} periodic Hartree-Fock (HF) calculations,\textsuperscript{44} although HF bulk band-structure calculations fail to reproduce the known antiferromagnetic alignment of magnetic moments within the tetrahedral and octahedral sublattices.\textsuperscript{26}

STM and low-energy electron diffraction (LEED) intensity analysis at 1000 K in a 10$^{-6}$ mbar atmosphere indicates the Fe$_3$O$_4$ (111) surface forms an unreconstructed bulk termination that exposes $\frac{1}{4}$ ML Fe$_{\text{tet1}}$ atoms over a close-packed oxygen layer, with protrusions arranged in a hexagonal lattice with a 6 Å periodicity.\textsuperscript{45,46} Both Paul \textit{et al.}\textsuperscript{47} and Shimizu \textit{et al.}\textsuperscript{48} report that the Fe$_{\text{tet1}}$ termination is routinely observed in naturally grown single crystals at room temperature whereas Fe$_{\text{oct2}}$ appears only when the sample is prepared under oxygen-poor conditions, for instance, when exposed to UHV for a long period of time. This supports the results of GGA+$U$ calculations by Grillo \textit{et al.}\textsuperscript{49} and Kiejna \textit{et al.}\textsuperscript{50} where they found the two surfaces have comparable thermodynamic stabilities at this limit. Another type of coexistence of surface terminations has been observed by Berdunov \textit{et al.}\textsuperscript{51} Here the regular Fe$_{\text{tet1}}$ termination consists of the superstructure with an oxygen-rich surface for crystalline Fe$_3$O$_4$ (111) possibly with oxygen vacancies after annealing in an oxygen partial pressure of 10$^{-6}$ mbar at 950 K and subsequently cooling to room temperature.

An alternative to bulk termination reconstruction is the existence of defects, \textit{e.g.}, vacancies or adatoms, on the surface. Lennie \textit{et al.}\textsuperscript{41} have proposed that the irregular texture observed in their STM images indicates the presence of defects in the surface
layer. One of the possible defects is an iron vacancy formed on Fe\textsubscript{oct1} termination to stabilize the surface polarity. In the Fe\textsubscript{oct1} surface, a single V\textsubscript{Fe} is expected to be more stabilizing than two cationic vacancies as its surface charge (+5) from the remaining octahedral ions is closer to half the absolute value of the subsurface charge (-4), which fulfills the condition for the cancellation of the macroscopic dipole moments according to classical electrostatics.\textsuperscript{22} Ferric iron vacancies have also been observed on Fe\textsubscript{tet1}-terminated epitaxial Fe\textsubscript{3}O\textsubscript{4} (111) films grown on Pt (111) substrates in an 10\textsuperscript{-6} mbar oxygen partial pressure,\textsuperscript{46} as have inhomogeneities caused by FeO\textsubscript{x} agglomerates\textsuperscript{52} and FeO\textsubscript{1-x} overlayers\textsuperscript{46,47} on the surface. In order to evaluate the impact of defects, in addition to four bulk terminations in the [111] direction (Fe\textsubscript{tet1}, Fe\textsubscript{oct1}, Fe\textsubscript{oct2}, and O\textsubscript{1}), we have chosen to study single cationic vacancies formed at the octahedrally-coordinated iron layer (Fe\textsubscript{oct1}) on the surface, as shown in Figure 2a. Iron vacancies in octahedral sites are exceptionally noteworthy as they are associated with the oxidation redox cycles of magnetite(100)\textsuperscript{53} in contrast to oxygen vacancies in other metal oxides.\textsuperscript{54} We have also considered oxygen vacancies in the close-packed oxygen layer as proposed in earlier STM studies.\textsuperscript{51}

Other possible surface modifications are attachments of foreign atoms or ions from the residual atmosphere due to experimental conditions. Several STM studies have observed adsorbates above the atomic layer.\textsuperscript{47, 52} Importantly, high-resolution STM measurements show there is a distinctive step of 1.2±0.1Å above the regular Fe\textsubscript{tet1} terminated surface, depicted in Figure 2b, suggesting that surface iron atoms may be capped by a single oxygen atom at the atop position.\textsuperscript{42} A similar termination, with a hydroxyl group on top of the surface terminating cations, has been proposed after water
The STM results indicate that dissociative water adsorption to form surface hydroxyls takes place on a termination of Fe$_3$O$_4$ (111) thought to contain a 1/4 monolayer of Fe$^{3+}$ ions on top of a close-packed oxygen monolayer. To evaluate this class of defects, we have modified the Fe$_{tet1}$ termination with two adsorbates, an oxygen atom and a hydroxyl group. Thus, in total, we will consider four defect-containing surface terminations; (i) iron vacancy in Fe$_{oct1}$ ($V_{Fe}$, Figure 2a); (ii) oxygen vacancy in O$_1$ ($V_{O}$); (iii) oxygen adatom on Fe$_{tet1}$ (ferryl, Figure 2b); and (iv) hydroxyl adsorbate on the Fe$_{tet1}$ termination.

Figure 2 (a) Top view of the Fe$_{oct1}$ termination with one Fe vacancy. The vacant site is indicated with an arrow. (b) Side view of the Fe$_{tet1}$ termination with ferryl group formation upon attachment of an additional oxygen atom, indicated with an arrow, on the surface. Fe atoms are colored blue, and oxygen ions are colored red.

b) Computational details

First-principles calculations have been performed using spin-polarized DFT as implemented in the Vienna Ab initio Simulation Package (VASP). The ionic potentials are described by the projector augmented wave (PAW) pseudopotential.
with valence configurations of $3d^74s^1$ and $2s^22p^4$ for Fe and O atoms, respectively. The
cutoff energy for the plane-wave basis set was chosen to be 550 eV.

On the basis of our previous theoretical study on the full electronic structure of cubic
$\text{Fe}_3\text{O}_4$,\textsuperscript{60} we take the DFT+$U$ approach\textsuperscript{61} to modify the intra-atomic Coulomb interaction
among strongly correlated Fe $3d$ electrons. We consider the GGA exchange-correlation
functional of Perdew, Burke, and Ernzerhof (PBE).\textsuperscript{62-63} The DFT+$U$ method used here
is a simplified rotationally invariant formulation by Dudarev et al.\textsuperscript{64} where the on-site
Coulomb parameter, $U$, and exchange parameter, $J$, are combined into a single
parameter, $U_{\text{eff}} \equiv U - J$. We applied $U_{\text{eff}} = 4$ eV for Fe $3d$ electrons in the calculations.

Brillouin-zone integration was performed on Monkhorst-Pack grids\textsuperscript{65} with a $7 \times 7 \times 1$
mesh where a Gaussian-smearing approach with $\sigma=0.05$ eV is used during the ionic
optimization, while total energies and densities of states (DOS) were calculated using
the tetrahedron method with Blöch corrections.\textsuperscript{66} Atomic charges were estimated within
the Bader scheme using converged FFT grids.\textsuperscript{67-68} All of the vacancies and adatoms are
uncharged. Bulk properties for $\text{Fe}_3\text{O}_4$ were calculated using a unit cell containing 24 Fe
and 32 O atoms which was fully relaxed while preserving cubic symmetry. We found
that PBE+$U$ successfully reproduced the electronic and magnetic properties of the bulk
$\text{Fe}_3\text{O}_4$ and describe the results from this methodology throughout our discussion. As the
energies calculated using different values of $U$ cannot be compared, the value for $U$
must be transferrable amongst both the different multivalent oxides and metallic
systems. An alternative to the empirical $U$ term is the use of hybrid functionals which
may provide a better treatment of exchange and correlation effects in both extended as
well as localized states. In our previous work on bulk magnetite,\textsuperscript{60} we compared the
PBE+$U$ results to those of both a conventional exchange correlation functional, PBE, and a modified version of the range-separated hybrid functional, HSE06.\textsuperscript{69-70} As the optimal amount of Hartree-Fock (HF) exchange is system dependent, we utilized HSE(15%), where the HF exchange mixing coefficient, $a$, was taken to be 0.15 and $\omega$, the parameter that defines the partitioning between the short- and long-range exchange components, was 0.11 bohr\textsuperscript{−1}. A comparison to these other DFT approaches is presented in the supporting information (SI). The effect of dispersion corrections was examined for a subset of the systems studied, and they introduce only extremely small changes in the electronic structure.

A well-established tool to investigate the electronic structure of surfaces and interfaces at the DFT level is the repeated slab approach that allows one to take easy account of the two-dimensional periodic character of such systems. The surface has been modeled using a symmetric slab with of a $(1 \times 1)$ unit cell which is based on cubic-phase Fe$_3$O$_4$ ($a= 8.488$ Å). The slabs are separated by a vacuum space of approximately 20 Å. A top view and side view of the stacking sequence are shown in Figure 1. The optimized lattice vector within PBE+$U$\textsuperscript{60} are slightly overestimated compared to the experimental value ($a=8.396$).\textsuperscript{71} PBE+$U$ calculations provide a net magnetic moment of 3.99 $\mu_B$/f.u. compared to the experimental value of 4.05 $\mu_B$/f.u..\textsuperscript{72} In order to maintain bulk behavior below the surface, the central layers were kept fixed at the bulk crystal positions during the structural optimization while the outermost six layers on both sides of the slab were allowed to relax. The ion positions were optimized using a conjugate-gradient algorithm until the Hellmann-Feynman forces were less than 0.01 eV/Å and energy convergence was reached within $10^{-5}$ eV/atom. An antiferromagnetic ordering, where the sign of the
Fe\textsubscript{oct} atoms are antiparallel to those on the Fe\textsubscript{tet} atoms, was taken as the initial magnetic configuration and allowed to relax for all the surface models. The surface energy, atomic charge, magnetic moment, and relaxation pattern were analyzed using a series of Fe\textsubscript{tet1} terminations with different stoichiometry by changing the slab thickness and the results are shown in Table S1 in SI. The data presented in the text are for slabs consisting of more than 17 layers where these surface characteristics have converged.

c) Thermodynamic stability of surface terminations

The stability of various surface terminations of iron oxides can be determined using a simple thermodynamic approach. If the surface system is modeled in DFT by a slab with two equivalent surfaces, the surface energy can be obtained by the following equation where it is assumed that the Gibbs free energy ($G$) can be approximated by the calculated internal energy ($E$):

$$\gamma = \frac{1}{2A}(E_{\text{slab}} - E_{\text{bulk}})$$

In Equation 1, $\gamma$ denotes the surface energy, $A$ the surface area of the unit cell, $E_{\text{slab}}$ the total energy of the slab, and $E_{\text{bulk}}$ the total energy of the bulk system with the same number of atoms as in the slab: $E_{\text{bulk}} = N_{Fe}\mu_{Fe} + N_{O}\mu_{O}$; here, $N_{Fe}$ and $N_{O}$ represent the numbers of each type of atoms while $\mu_{Fe}$ and $\mu_{O}$ denote the chemical potentials of a Fe atom and an O atom. When there is enough bulk material to act as a thermodynamic reservoir, the potentials of the components are no longer independent but are related by the condition of thermal equilibrium in bulk Fe\textsubscript{3}O\textsubscript{4}:

$$3\mu_{Fe} + 4\mu_{O} = \mu_{Fe_{3}O_{4}}$$

(2)
where $\mu_{Fe_3O_4}$ is the DFT energy of bulk Fe$_3$O$_4$ per formula unit. This constraint can be used to eliminate the dependence of the surface energy on $\mu_{Fe}$. In this approximation and at equilibrium, the surface energy can be expressed as a function of the oxygen chemical potential as:

$$\gamma = \frac{1}{2A} \left[ E_{slab} - \frac{1}{3} N_{Fe} \mu_{Fe_3O_4} + \left( \frac{4}{3} N_{Fe} - N_{O} \right) \mu_{O} \right]$$

(3)

In this way, it is possible to determine the relative stabilities of slabs with different stoichiometries.

The oxygen chemical potential varies with its partial pressure ($p$) and temperature ($T$):

$$\mu_{O}(T, p) = \mu_{O}(T, p^*) + 1/2 k_B T \ln \left( \frac{p}{p^*} \right)$$

(4)

where $p^*$ is the pressure in the reference state and $k_B$, the Boltzmann constant. As experimental conditions have a considerable impact on the oxygen chemical potential, it is necessary to evaluate the surface energies for various surface terminations as a function of the chemical potential of oxygen to determine the surface stability in a given environment. For example, the oxygen chemical potential at $p = 10^{-6}$ mbar and $T = 900$ K, one of the typical conditions for annealing of magnetite, is calculated to be -1.78 eV with the $\mu_{O}(T, p)$ reference state defined as the total energy of molecular oxygen at the standard pressure, $p^* = 1$ bar. It is useful to restrict the range of oxygen chemical potentials by considering the limiting conditions. The oxygen-rich limit can be defined as the chemical state in which molecular oxygen starts to condense on the surface. By
assuming that such condensed oxygen is in equilibrium with gas-phase oxygen, the upper limit of the oxygen chemical potential becomes:

\[ \mu_O(T, p) = \frac{1}{2} E_{O_2} \]  

(5)

where \( E_{O_2} \) is the DFT energy of an \( O_2 \) molecule.

At the oxygen-poor limit, reduction of the oxide from magnetite to wüstite (FeO) and further to metallic iron occurs. Thus, a reasonable minimum \( \mu_O \) corresponds to the condition where Fe crystallizes and oxygen separates onto the surface according to the equilibrium of bulk magnetite, that is:

\[ \mu_O = \frac{1}{4}(\mu_{Fe_2O_4} - 3\mu_{Fe,bulk}) \]  

(6)

where \( \mu_{Fe,bulk} \) is the DFT-calculated chemical potential of metallic Fe.

**III. Results and Discussion**

We have selectively calculated four bulk terminations in the [111] direction; \( Fe_{tet1} \), \( Fe_{oct1} \), \( O_1 \), \( Fe_{oct2} \), omitting the \( Fe_{tet2} \) and \( O_2 \) surfaces that were reported to be highly unstable.\(^{31}\)

While it must be emphasized that the bulk-terminations have been examined previously,\(^{49-60}\) it is important to include a brief discussion here in order to illustrate the large differences between these surfaces and the four modified surfaces incorporating surface defects (iron vacancy in \( Fe_{oct1} \), oxygen vacancy in \( O_1 \); oxygen adatom (ferryl) and hydroxyl adsorbate on the bulk \( Fe_{tet1} \) termination). Below, we will turn successively to a discussion of: surface stability, ionic relaxation, surface electronic structure and chemistry, and spin polarization near the Fermi level for each of these surface models.
a) Surface stability

Using Equation 3, we have evaluated the relative stabilities of the surface models. The results are plotted as a function of oxygen partial pressure in Figure 3 for the PBE+U functional. Previous theoretical studies containing only bulk terminations have concluded that the Fe\textsubscript{tet1} termination has the lowest surface energy in the oxygen-rich regime, while the Fe\textsubscript{oct2} and Fe\textsubscript{oct1} surfaces are competitive in the oxygen-poor regime.\textsuperscript{39, 49-50, 74} The defect-containing surfaces that we examine here are viable in comparison to the bulk-like terminations. For example, as both the V\textsubscript{O} and ferryl-terminated surfaces are within 100 meV/Å\textsuperscript{2} (1.60 J/m\textsuperscript{2}) of the Fe\textsubscript{tet1} surface near the oxygen-rich limit, specific sample preparation methods or environmental conditions may lead to the observation of these oxygen-rich surfaces. The ferryl termination is more stable than the formally oxygen-terminated surfaces over most of the pressure range. Experimental observations by Berdunov \textit{et al}.\textsuperscript{51} indicate the presence of such oxygen-terminated surfaces, which are higher in surface energy than the defect-containing surfaces. This is in agreement with the calculated large binding energy of an oxygen atom located on a bridge position between Fe\textsubscript{tet1} and Fe\textsubscript{oct2} atoms by Santos-Carballal \textit{et al}.\textsuperscript{74} We note that there is a well-known overbinding problem\textsuperscript{75} in small molecules within DFT (and DFT+\textit{U}), which can lead to an inaccurate energy for molecular oxygen. If we replace the calculated binding energy of oxygen gas with the experimental value, 5.23 eV\textsuperscript{76} ($E_{B,exp}$), the chemical potential limit can be rewritten as $\mu_{O_2,exp} = E_{B,exp} + 2\mu_{O,DFT}$. This equates to shifting the oxygen-rich limit to a more positive value by 0.4 eV, shown as the rightmost vertical line in Figure 3. This is smaller than the 1.36 eV shift determined by
differences in experimental and calculated formation energies over a series of oxides, which was used previously.$^{77}$ At this extreme, the ferryl termination becomes isoenergetic with the Fe_{tet1} termination.

In the oxygen-poor limit, the Fe_{oct2} termination is shown to be more stable than the Fe_{tet1} termination, as suggested before.$^{41, 47-48}$ The Fe_{oct1} termination becomes more energetically favored than the Fe_{oct2} termination only at extremely low oxygen chemical potentials, beyond the estimated oxygen-poor limit, because of the higher density of Fe ions on the Fe_{oct1} surface. However, missing Fe atoms on this termination, labeled as V_{Fe}, moderately stabilize the surface, leading to competitive surface energies with the Fe_{oct2} termination over the entire range of oxygen chemical potentials.

The sensitivity of our results to choice of functional is presented in Figure S1 of the SI. Despite the limitation of using a constant empirical $U$ correction for both the oxide and the metallic species, this method reproduces the enthalpy of formation for binary-manganese oxides within 4%.$^{78}$ Additional quantitative accuracy could be obtained by utilizing a method systematically combining GGA and GGA+$U$ energies (however, further experimental data would then be required$^{79}$). In general, such additional numerical accuracy is not needed as the surface stabilities using different functionals show the same ordering as the PBE+$U$ method but with further stabilization of the ferryl termination. The hydroxyl surface is also dependent on the hydrogen (or water) partial pressure and is not plotted in Figure 3. However, the observation of hydroxyl groups on epitaxial Fe$_3$O$_4$ (001) surfaces$^{80}$ indicate that this termination can appear not only in humid conditions,$^{81}$ but is competitive in ultrahigh-vacuum environments.
Figure 3 PBE+U derived surface energies of the different terminations plotted against the oxygen chemical potential. The vertical black lines indicate the allowed range of the chemical potential. The bottom two horizontal axes indicate corresponding oxygen pressure in log scale at 300 K and 900 K, respectively ($p^o = 1$ bar).

The modified surfaces presented above are high in defect density, with one defect or adlayer in each unit cell. The stability of the ferryl surface with a lower defect coverage on the Fe$_{tet1}$ termination ($\theta = 1/4$) is shown in Figure 4. This approach has previously been used to study mixed terminations of hematite surfaces. The resulting mixed surface is nearly equal in energy to the Fe$_{tet1}$ termination over much of the potential range, and the point at which it becomes lower in energy shifts further toward lower oxygen chemical potentials. In a macroscopic surface, these defects may be present, even if higher in energy in a small unit cell. Given the stringent requirements for spin-injection and detection in devices, contributions from these slightly higher-energy surfaces may determine the overall success of a device.
Figure 4 Surface free energy change as a function of the oxygen chemical potential for two different ferryl concentrations on the surface: 25% (dotted black) and 100% (solid black). The blue line indicates Fe\textsubscript{tet} termination. The arrows in inset images correspond to the positions where oxygen adatoms are attached.

b) Ionic relaxation

As described above, (111) surfaces created from cleavage of bulk cubic magnetite are unstable as two inequivalent layers of opposite charge densities alternate perpendicular to the surface. A high dipole moment, 29.5 D, is estimated within the repeat unit of six distinct layers when using formal charges and the bulk interlayer spacing.\textsuperscript{22} One pattern of stabilization of these surfaces is through significant changes in the nuclear positions of the atoms comprising the surface layers.

The interlayer relaxations are determined mostly by a response to the large dipole moments of alternating polar layers in the surface layers. The relaxations between
adjacent layers (ij) of the outmost six layers as a percentage ($\Delta_{ij}$) of the bulk interlayer distance are shown in Figure 5 for terminations based on (a) Fe$_{tet1}$, (b) O$_1$, and (c) Fe$_{oct1}$.

For instance, in the Fe$_{tet1}$ termination, the surface Fe$_{tet1}$ atom shows a strong inward relaxation and the interlayer spacing between the surface layer and subsurface layer decreases by 37%. Further reduction in polarity is achieved by a 22% compression in the O$_1$-Fe$_{oct1}$ distance, which fully stabilizes the surface as there are progressively smaller changes in geometries for the rest of the slab. The overall relaxation of the Fe$_{tet1}$ termination is consistent with a previous LEED analysis.$^{45}$ For the ferryl and hydroxyl surfaces, the surface Fe$_{tet}$ atom retains its bulk-like tetrahedral coordination by binding to the oxygen adatom; also, the electrostatic attraction of a surface Fe ion to the oxygen atoms in the sublayer is balanced by the adatom. The newly formed Fe=O and Fe-O bonds are estimated to be 1.62 Å and 1.77 Å, which are consistent with the 1.5 Å-height adsorbates reported in STM studies.$^{52}$ Hence, the Fe$_{tet1}$-O$_1$ distance decreases by 17% and 9% for the ferryl and hydroxyl surfaces. However, the relaxations in the next layers in the slab are now in the opposite direction than in the Fe$_{tet1}$ slab, with the Fe$_{oct1}$-O$_2$ distance being compressed and the O$_2$-Fe$_{tet2}$ distance slightly expanding.
Figure 5. Surface relaxations in the outmost six layers given as a percentage ($\Delta_{ij}$) of the bulk interlayer distances between two adjacent surface planes (i) and (j) projected onto the c axis for the seven terminations of Fe$_3$O$_4$ (111). Terminations based on (a) Fe$_{tet1}$, (b) O$_1$, and (c) Fe$_{oct1}$ are shown in separate panels. The bulk interlayer distances between Fe$_{tet1}$-O, Fe$_{tet}$-O, and Fe$_{tet}$-Fe$_{oct2}$ are 1.19 Å, 0.64 Å, and 0.61 Å, respectively.

There are only moderate changes in interlayer spacing in the O$_1$ surface, and these are smaller in the V$_O$ surface due to the reduced surface charge. The introduction of a Fe vacancy in the Fe$_{oct1}$ termination (V$_{Fe}$), shown in 5c, changes the local geometries at the surface, although in a different fashion from the bulk Fe$_{oct1}$ termination. In the V$_{Fe}$ termination, one of the four oxygen atoms in the subsurface ends up migrating above
the iron surface due to the vacated volume and a strong electrostatic attraction between
two outermost layers. As a consequence, the surface Fe\textsubscript{oct1} atoms bind more strongly to
the surrounding oxygen atoms with a reduced bond length, 1.88 Å, compared to the
bond in the Fe\textsubscript{oct1} termination, 1.93 Å. Moreover, three remaining oxygen atoms in the
O\textsubscript{2} layer are allowed to fairly interact with the underneath iron layers, Fe\textsubscript{tet2} and Fe\textsubscript{oct2},
since the protruding oxygen exclusively forms ionic bonds with the two Fe atoms on the
surface.

c) Surface electronic structure and chemistry

Since the surfaces are created by cleaving ionic bonds, significant charge redistribution
occurs due to the appearance of dangling bonds. In this study, density of states (DOS),
Bader charge, net magnetic moment, and work function are calculated to describe the
related stabilization mechanism upon surface cleavage and change in surface chemistry.

Density of states

The DOS projected on Fe\textsubscript{oct}, Fe\textsubscript{tet}, and O atoms for near-surface (3-4) layers of each
termination are displayed in Figure 6. The electronic structure of bulk Fe\textsubscript{3}O\textsubscript{4} in a cubic
\textit{Fd\bar{3}m} phase, shown in the SI (Figure S2), can be described as a half-metallic oxide,
which is consistent with the crystal-field splitting explanation for the octahedral Fe ions
that the fivefold \textit{d} levels separate into three degenerate \textit{t\textsubscript{2g}} levels and two degenerate \textit{e\textsubscript{g}}
levels.\textsuperscript{60} The PDOS for the central layers of the slab retain this electronic structure.

The PDOS for the surface layers in the Fe\textsubscript{tet1} termination, plotted in Figure 6a, show that
the minority (spin-down) Fe \textit{t\textsubscript{2g}} states from the octahedral sites are no longer present at
or near the Fermi level and there appears a surface state related to tetrahedral-site iron in the middle of the majority spin (spin-up) gap. The partial charge density for this state is shown in Figure 6d, and demonstrates that the 3d orbitals in the direction of reduced coordination, i.e., the [111] direction, are more stabilized and now lie below the Fermi level. In general, surface truncation of iron-terminated surfaces promotes a further splitting within Fe\textsubscript{tet1} e\textsubscript{g} and t\textsubscript{2g} orbitals at the surface as the symmetry is lowered because of missing apical oxygen.\textsuperscript{83} Since the Fe\textsubscript{tet1} termination has a positively charged surface, the top Fe\textsubscript{tet1} layer obtains more electrons through electronic redistribution to reduce the polarity, with the bottom of the conduction band becoming occupied. The ferryl termination, where the Fe\textsubscript{tet1} surface has an additional oxygen atom bound to the exposed iron atom, shows a different DOS with respect to the Fe\textsubscript{tet1} surface; the ferryl group comprises a set of states near the Fermi level and the surface state related to tetrahedral-site iron no longer appears in the spin-up gap (the PDOS of adsorbate oxygen is colored in magenta in Figure 6). In the hydroxyl termination these hybridized states are stabilized and now located in the same region as the surface oxygen atoms. In the O\textsubscript{1} subsurface, oxygen 2p-derived states still have a strong dispersion between -1 eV to -6 eV. Compared to bulk states, they are shifted to higher binding energy by \approx 1 eV in response to the lowered coordination upon surface cleavage.
Figure 6. Electronic densities of states of near-surface (3-4) layers for the surfaces: (a) Fe\textsubscript{tet1}, (b) ferryl, and (c) hydroxyl terminations from PBE+U calculations from -8 eV below to +2 eV above the Fermi level. The Fermi level (= zero of energy, see text) is indicated with a dashed line. Partial charge densities of the surface states marked with short arrows for (d) Fe\textsubscript{tet1} and (e) ferryl terminations. PDOS from Fe\textsubscript{tet1} atoms are blue, Fe\textsubscript{oct} atoms are green and lattice oxygen atoms are red.

In the Fe\textsubscript{oct1} termination with a vacancy, V\textsubscript{Fe}, there are no longer octahedral states observed around the Fermi energy in contrast to Fe\textsubscript{oct2} (see Figure S2 in SI) although the two models have the same Fe coverage over the oxygen sublayer, \textit{i.e.}, \(\frac{1}{2}\) ML. Instead, the electrons are redistributed over the surface in a fashion similar to the Fe\textsubscript{oct1} termination. In both cases, the Fe 3\textit{d} states on octahedral sites are shifted to lower energy, leading to a reduction of these Fe atoms to stabilize the positively charged outmost layer. The occupied Fe 3\textit{d} minority spin states (between -1 eV and -0.5 eV) are more populated in the bulk-terminated Fe\textsubscript{oct1} surface. Despite overall similarity in the PDOS between Fe\textsubscript{oct1} and V\textsubscript{Fe}, the states just below the Fermi-level (marked with arrows c and d in Figure 7) are qualitatively different. In the Fe\textsubscript{oct1} surface the charge is
delocalized over the first layer of the surface, whereas the iron vacancy leads to the state becoming more localized on each Fe atom on the surface.

Figure 7. Electronic densities of states of near-surface (3-4) layers for the surfaces: (a) Fe_{oct1}, (b) and (b) V_{Fe} terminations from PBE+U calculations from -8 eV below to +2 eV above the Fermi level. The Fermi level (= zero of energy, see text) is indicated with a dashed line. Partial charge densities of the surface states marked with short arrows for (c) Fe_{oct1} and (d) V_{Fe} terminations. PDOS from Fe_{tet1} atoms are blue, Fe_{oct} atoms are green and lattice oxygen atoms are red.

For the O_{1} termination, the surface also tends to be oxidized due to the highly negatively charged surface layer. Hence, the O 2p states which comprise the valence band in the bulk shift above the Fermi level, leading some of these states to become unoccupied upon cleavage. The V_{O} surface shows the similar results. The DOS for these surfaces are shown in the section S5 of SI.

**Bader charge and net magnetic moment**

The ionic and electronic relaxations described above can also be understood through changes in Bader charge and magnetic moment on each of the atoms when going from bulk to the surface, as presented in Table 1. The atomic charges for the bulk cubic
system show that all the equivalent octahedral ions with a formal oxidation state of +2.5
have a Bader charge of +1.72 |e| and the tetrahedral ferric sites, a charge of +1.86 |e|.
The oxygen ions each have a charge of -1.33 |e|. In general, the charges of both Fe and
O tend to decrease in absolute values compared to the bulk due to reduced
coordination of surface atoms and electronic redistribution. We present the results of
four terminations in Table 1. The hydroxyl surface has a similar charge and spin
distribution as the ferryl surface. As with the surface relaxations and DOS, there is little
change between the O_1 and V_O terminations.

| Table 1. Average change in atomic charge and magnetization (per atom) in the top layers for four terminations. All the values listed represent the changes from the values for the equivalent atoms in the bulk system. Net magnetic moment for bulk Fe_{oct}, Fe_{tet}, and O atom are 3.96-3.98, 4.09, and 0.03 µ_B, respectively. The experimental magnetic moment for bulk Fe_{tet} is 3.82 µ_B.\(^{84}\) Values from the PW91+U (U_{eff} = 3.61 eV) calculations of ref. 50. |
|-----------------|-----------------|-----------------|-----------------|
| layer           | Fe_{tet1}       | Fe_{oct1}       | ferryl          |
| Fe_{tet1}       | +0.41 (0.54\(^{a}\)) | +0.60 (-0.46) | O               |
| O_1             | -0.07 +0.22     |                 | Fe_{tet1}       |
| Fe_{oct1}       | -0.19 +0.22     |                 | O_1             |
| O_2             | -0.04 +0.04     |                 | Fe_{oct1}       |
| layer           | V_{Fe}          |                 |                 |
| Fe_{oct1}       | +0.60 -0.46     |                 | Fe_{oct1}       |
| O_2             | +0.03 -0.01     |                 | O_2             |
| Fe_{tet2}       | +0.02 -0.04     |                 | Fe_{tet2}       |
| Fe_{oct2}       | +0.24 -0.30     |                 | Fe_{oct2}       |

Due to the dangling bonds, the top layer in the Fe_{tet1} termination gains 0.41 electrons, as anticipated from the change in the DOS that the originally unoccupied conduction band of the spin-up channel is slightly shifted below the Fermi level. The oxygen ions in the second layer also help reduction of the surface dipole by losing 0.07 electrons to be
less negative, which contributes to further stabilization. Meanwhile, each of the three iron ions in the third layer becomes more positive by 0.19 $|e|$, which alleviates the overall negatively charged nature of the surface and subsurface. In the ferryl termination, i.e., when adding an atop oxygen to the exposed Fe$_{tet1}$ atom, the charge on the Fe$_{tet1}$ atoms remain close to the bulk value. Just as in the Fe$_{tet1}$ surface, there is an increase in positive charge character for Fe$_{oct1}$ in the third layer.

In the Fe$_{oct1}$ and V$_{Fe}$ terminations, there are essentially more electrons residing on the topmost layer than in the bulk system. These electrons reduce the positive charge on the iron atoms on the surface, although the O$_2$ sublayer preserves its bulk charge state. Notably, in the V$_{Fe}$ termination, the cationic vacancy brings about a lesser electron gain to the remaining two surface Fe$_{oct1}$ atoms.

The variations in magnetic moment of the surface atoms are consistent with the changes in atomic charge, especially for the metal-terminated surfaces. For the ferryl group, there is a decrease in magnetic moment of -0.83 $\mu_B$ for the Fe$_{tet1}$ atom despite a negligible variation in atomic charge, 0.04 e; this is the result of combined effects due to depopulation of the originally occupied spin-up Fe$_{tet}$ 3$d$ orbitals in the valence band and population of the previously unoccupied spin-down Fe$_{tet}$ 3$d$ orbitals, which is related to bonding with the atop oxygen atom.

In the top two iron layers of Fe$_{oct2}$ termination, see Table S3, the Fe$_{oct}$ atoms gain electron density in the bulk unoccupied minority spin (spin-down) bands, which leads to a magnetic moment decrease. This is also the case for the Fe$_{tet1}$, Fe$_{oct1}$, and V$_{Fe}$
terminations; this result is reasonable as the Fe 3d orbitals in the minority spin states are now more partially occupied while the majority bands stay fully occupied.

4 Work function

The work function is one of the critical surface parameters and is illustrative of the charge distribution over the surface of a material. The work function, \( \Phi \), of a surface in the DFT-slab framework is defined as:

\[
\Phi = E_{\text{vac}} - E_f
\]  

(7)

where \( E_{\text{vac}} \) is the plane-averaged electrostatic potential energy of an electron in the vacuum region away from the slab surface at the distance where the potential energy has reached its asymptotic value, and \( E_f \) denotes the Fermi energy of the total system.

Table 2 collects the work functions for each surface. The calculated work function of 5.76 eV for the model Fe\text{tet1} termination is 0.24 eV higher than the experimentally measured value, 5.52 eV, for the same surface.\textsuperscript{23} The value that we calculate for the Fe\text{tet1} surface is 0.28 eV higher than the values from PW91+\( U \) (\( U_{\text{eff}} = 3.61 \) eV) calculations in ref. 50 and 0.15 eV and 0.75 eV lower than the O\textsubscript{1} and Fe\text{oct2} value, respectively. The other three iron-terminated surfaces have lower work functions by up to 2.61 eV whereas the oxygen-terminated surfaces, ferryl and O\textsubscript{1}, show a narrow range (0.33 eV) of work functions. Thus, there is a large difference between the calculated work functions of the iron- and oxygen-terminated surfaces, which can be explained on the basis of simple electrostatics: Magnetite has a dipole moment in the [111] direction.
generated by alternating cation layers and anion layers; when the termination ends with
a negatively charged layer, the corresponding potential increases at the surface, which
gives rise to a relatively high vacuum level, while positively top charged layers give rise
to the opposite effect.

Table 2. Work function is calculated for the eight surface terminations. *Value from the
PW91+U (U_{eff} = 3.61 eV) calculations of ref. 50.

<table>
<thead>
<tr>
<th>Surface</th>
<th>Termination</th>
<th>Work function (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe_{oct1}</td>
<td>3/4ML Fe_{oct}</td>
<td>3.91</td>
</tr>
<tr>
<td>V_{Fe}</td>
<td>2/4ML Fe_{oct}</td>
<td>4.03</td>
</tr>
<tr>
<td>Fe_{oct2}</td>
<td>2/4ML (Fe_{oct}+Fe_{tet})</td>
<td>3.15 (3.90)^a</td>
</tr>
<tr>
<td>Fe_{tet1}</td>
<td>1/4ML Fe_{tet}</td>
<td>5.76 (5.48)^a</td>
</tr>
<tr>
<td>hydroxyl</td>
<td>1/4ML Fe_{tet}+OH</td>
<td>5.80</td>
</tr>
<tr>
<td>ferryl</td>
<td>1/4ML Fe_{tet}+O</td>
<td>7.61</td>
</tr>
<tr>
<td>V_{O}</td>
<td>3/4ML O</td>
<td>7.33</td>
</tr>
<tr>
<td>O_{1}</td>
<td>1ML O</td>
<td>7.94 (8.09)^a</td>
</tr>
<tr>
<td>Exp.</td>
<td>23/4ML Fe_{tet}</td>
<td>5.52</td>
</tr>
</tbody>
</table>

The work function decreases by 2.6 eV upon an increase in iron density over the
oxygen layer from 1/4 ML (Fe_{tet1}) to 2/4 ML (Fe_{oct2}). When introducing a Fe cation
vacancy on the Fe_{oct1} termination, the work function increases by +0.11 eV compared to
the bulk-terminated Fe_{oct1} surface (3.91 eV). This small change in work function
between Fe_{oct1} and V_{Fe} can be attributed to the charge states of the Fe_{oct} atoms
becoming more positive, from +1.12 |e| to +1.37 |e|, in the presence of the defect. The
work function for Fe_{oct2} is also smaller than the work function of the V_{Fe} termination with
exactly the same iron coverage at the surface. Given that the exposed surface cations
of Fe_{oct2} (+1.33 |e|) are as positive as in the V_{Fe} surface, the oxygen atom above the V_{Fe}
surface plays a substantial role in determining the work function difference of 0.88 eV between the two surfaces.

In the ferryl termination, the negatively charged oxygen atoms on the top of the Fe atoms bring about a substantial increase of 1.85 eV in work function compared to the bulk-terminated Fe\textsubscript{tet1} surface. Despite having a significantly less dense oxygen layer, the work function of the ferryl termination then is comparable to that of the oxygen-terminated surfaces, O\textsubscript{1} and V\textsubscript{O}. An explanation for this behavior is the bond dipole formed between iron and oxygen, which leads to a large dipole component in the direction perpendicular to the surface. In the bulk oxygen terminations, the Fe-O bonds are aligned at an angle to the surface, reducing the component perpendicular to the surface. Thus, the atop oxygen on Fe\textsubscript{tet1} plays a significant role in considerably increasing the work function despite having a smaller negative charge than the oxygen atoms on other terminations. This impact is diminished by the additional hydrogen in the hydroxyl group. Finally, we note that while there is some quantitative variation depending on the choice of methodology (shown in Figure S3), e.g., the work function varies up to 1.53 eV for the O\textsubscript{1} surface between PBE and HSE(15%), however, the overall trends upon terminations are similar compared to the results from PBE+\textit{U}.

d) Surface chemistry

As a consequence of atomic displacements and charge redistributions, the surfaces can have quite different chemistries compared to those predicted from the bulk properties; in addition, defect-modified surfaces can be different than bulk-terminated surfaces. An example is depicted in Figure 8, which illustrates the electrostatic potential profile measured at 1.3 Å, a typical bonding distance, above from the surface atoms for the
Fe$_{\text{oct1}}$ and V$_{\text{Fe}}$ terminations. Here, the octahedral Fe positions are distinguishable because of their lower potential, displayed in blue. Combined with the Bader charge analysis, it is clear that Fe$_{\text{oct1}}$ and V$_{\text{Fe}}$ are markedly different surfaces given the fact that in the latter the positive charge near the iron atoms is intensified and the oxygen protrusion contains a slight negative charge, in comparison to the bulk-truncated surface. Thus, it is important to emphasize that the mode of chemisorption of an organic layer, for instance, the binding of a phosphonic acid as was done on the (La, Sr)MnO$_3$ (LSMO) surface, could be significantly different on a vacancy-containing surface compared to a bulk termination. This change in mode of chemisorption would be significant in the performance of SAM-based magnetic tunnel junctions.

![Figure 8](image.png)

**Figure 8.** Planar electrostatic potential plot evaluated at 1.3 Å above the Fe$_{\text{oct1}}$ atoms for the Fe$_{\text{oct1}}$ (left) and V$_{\text{Fe}}$ (right) termination. The potential is illustrated in the reverse rainbow spectrum: lower potential in the blue region and higher potential in the red region.

e) Spin polarization near the Fermi level

The spin polarization at the surface is a major factor determining the spin-injection and collection efficiency of an electrode. The spin polarization ratio at the Fermi level can be
defined as \( P(E_f) = \frac{n_\uparrow - n_\downarrow}{n_\uparrow + n_\downarrow} \) where \( n_\uparrow \) and \( n_\downarrow \) are the densities at \( E_f \) of the majority and minority spins, respectively. Dedkov et al.\textsuperscript{86} measured \( P(E_f) \) to be -80±5% at room temperature near the surface of epitaxial \( \text{Fe}_3\text{O}_4 \) (111) films on Fe (110) using spin-resolved photoemission spectroscopy. They attributed the high spin polarization to the bulk \( \text{Fe}_3\text{O}_4 \) with a small reduction from the ideal value, -100%, due to the excitation of spin waves at the surface. The half-metallic state of \( \text{Fe}_3\text{O}_4 \) is retained even after high oxygen exposure as the spin-down electrons retain a high density-of-states near \( E_f \) contrary to spin-up electrons. In contrast to those results, Pratt et al.\textsuperscript{87} reported the opposite spin polarity using the naturally-grown single crystal annealed under <10\textsuperscript{-9} bar at 550 °C and proposed \( P(E_f) \) of >20% measured using a spin-polarized metastable helium beam [\( \text{He}(2^3\text{S}) \)] under magnetic fields up to 4 T at 298K. To reconcile these data, the authors assigned the difference to the penetration depth of their probe beams, which is much lower in the latter study than in UV photoemission where the octahedrally coordinated Fe layers below the surface can contribute to the cumulative spin polarization. This is also confirmed in the present study based on the quantitative evaluation of layer resolved spin polarity, as illustrated in Section S7. A comparison of the functional dependence of the cumulative layer polarization is also given in Figure S5.

In Figure 9, the spin polarizations, \( P(E) \), calculated for the top six layers, \textit{i.e.} one repeat unit (corresponding to approximately 4.8 Å from the surface), are plotted for the eight terminations. As seen above, the DOS have distinctive features depending on the surface termination; these features are directly reflected in the surface spin polarization. The \( \text{Fe}_{\text{tet1}} \) termination, panel a, shows a positive spin polarization below and negative above the Fermi level. In contrast, the ferryl termination presents a negative spin
polarization on both sides of the Fermi energy; as seen in the DOS, the negative spin
polarity at $E_f$ originates from Fe 3d-O 2p hybridized orbitals of the newly formed Fe=O.
The negative spin polarization below the Fermi level is no longer evident in the hydroxyl
termination, as the spin-down states associated with the adatom are no longer in the
spin-up gap. The defect structure of the Fe$_{tet1}$ termination could significantly decrease
the positive value of $P(E_f)$, expected from the bulk electronic structure. As the
magnitude and polarization of the PDOS of the Fe$_{oct1}$ and V$_{Fe}$ terminations are similar,
P(E) presents a similar profile in the region of the Fermi level. In the Fe$_{oct1}$ surface, the
polarization ratio becomes positive slightly above the Fermi level, which is not seen in
the V$_{Fe}$ surface. The O$_1$ and V$_O$ terminations have nearly +100% spin polarization from -
0.5 eV to +0.5 eV and -0.5 eV to +0.15 eV, respectively.
Figure 9. Spin polarization as a function of energy, $P(E)$, calculated for eight terminations within $[E_f - 0.5 \text{ eV}, E_f + 0.5 \text{ eV}]$. The vertical dotted lines indicate the Fermi level. The blue regions correspond to positive spin polarization while the red ones indicate negative spin polarization.

IV. Conclusions

We have discussed the results of PBE+$U$ calculations on modified terminations of magnetite (111) in terms of their structural, electronic, and magnetic properties and
compared them to the bulk-terminated surfaces. The most probable surface terminations for Fe$_3$O$_4$ were compared using their relative stabilities for the relevant limiting regimes of oxygen chemical potentials. For the surfaces comprised of bulk-like terminations, the Fe$_{tet1}$-terminated surface has the lowest surface energy over a wide range of oxygen chemical potentials while Fe$_{oct2}$ becomes more stabilized in the oxygen poor limit. Importantly, the surfaces that contain either oxygen adatoms or cationic vacancies are calculated to be close in stability to the Fe$_{tet1}$ surface at oxygen-rich and oxygen-poor conditions, respectively. These modified surfaces with adsorbates or cationic vacancies have significantly different surface geometries and electronic/magnetic structures with respect to the previously considered structures without any defects. In particular, these defects have a major impact on the electron redistributions over surface layers, which lead to different spin polarizations near the Fermi level and distinctive surface chemistries. Current work is in progress in our laboratories to evaluate the impact of these surface chemistries on the nature of interfaces with organic layers.

Supporting Information

Electronic Supplementary Information (ESI) available: The methodological details for the range-separated hybrid functionals are given in Section S1. Convergence of electronic properties with respect to slab thickness is given in Section S2. Analyses of the surface stabilities and work functions using various DFT methodologies and exchange-correlation functionals are given in S3 and S6. Tabulated data for the ionic relaxations is given in Section 4. Section S5 shows the PDOS for bulk magnetite and all
seven terminations examined. Section S7 describes the layer-resolved spin polarization variation within PBE and hybrid functionals.

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