Chemical Science

EDGE ARTICLE

Check for updates

Cite this: Chem. Sci., 2020, 11, 2448

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 4th December 2019 Accepted 23rd January 2020

DOI: 10.1039/c9sc06149e

rsc.li/chemical-science

Introduction

The termination of metal oxide surfaces is critical for their functionality and tuning their chemical behaviour. For example, when ceria is reduced by CO, the (110) termination has the lowest oxygen vacancy formation energy of the three low Miller index terminations.^{1,2} Another example is that only {100} terminated Cu₂O nanocrystals exhibit activity for the photodegradation of methyl orange, but not {111} or {110} surfaces.³ These termination-dependent phenomena are related to the local geometry, the bonding, and the chemical identity on the catalyst surface at an atomic level.

Magnetite (Fe₃O₄), which is low-cost and abundant, is a promising material for a wide range of applications,⁴⁻⁶ including catalytic processes for mitigating environmental toxins.⁷ The reducible material contains a spinel structure containing both Fe^{2+} (octahedral coordinated) and Fe^{3+} (tetrahedral and octahedral coordinated). $Fe_3O_4(111)$ is the most thermodynamically favoured termination of magnetite,⁸

Dual Lewis site creation for activation of methanol on $Fe_3O_4(111)$ thin films⁺

Fang Xu, ^(b)^a Wei Chen, ^(b)^b Constantin A. Walenta, ^(b)^a Christopher R. O'Connor ^(b)^a and Cynthia M. Friend ^(b)*^{ac}

Despite a wide application in heterogeneous catalysis, the surface termination of $Fe_3O_4(111)$ remains controversial. Herein, a surface with both Lewis acid and base sites is created through formation of an $Fe_3O_4(111)$ film on α - $Fe_2O_3(0001)$. The dual functionality is generated from a locally nonuniform surface layer of O adatoms and Fe_{tet1} sites. This reactive layer is reproducibly formed even in oxygen-free environments because of the high mobility of ions in the underlying α - $Fe_2O_3(0001)$. The atomic structure of the $Fe_3O_4(111)$ surface was identified by scanning tunneling microscopy (STM) and density functional theory (DFT) using the registry of the overlayers with the surface and the distinct electronic structure of oxygen adatom (O_{ad}) and uncovered lattice Fe_{tet1} . The surface is dominated by the interface of O_{ad} and Fe_{tet1} . a Lewis acid–base pair, which favors methanol dissociation at room temperature to form methoxy. Methoxy is further oxidized to yield formaldehyde at 700 K in temperature programmed reaction spectra, corresponding to an approximate activation barrier of 179 kJ mol⁻¹. The surface termination of $Fe_3O_4(111)$ is fully recovered by rapid heating to 720 K in vacuum, demonstrating the high mobility of ions in this material. The work establishes a clear fundamental understanding of a unique magnetite surface and provides insights into the origin of selective oxidation of alcohols on magnetite-terminated catalysts.

forming repeating layers of $-Fe_{tet1}-O_1-Fe_{oct1}-O_2-Fe_{tet2}-Fe_{oct2}-Fe_{tet1}-$ (subscripts refer to a tetrahedral, octahedral coordination, or an oxygen layer).⁹ All iron-containing layers share a similar hexagonal unit cell with a spacing of ~0.6 nm.

The surface termination of Fe_3O_4 is important because it affects local electronic structure and therefore also local chemical reactivity. Depending on the surface termination, either Lewis acid sites (electron acceptors) or Lewis base sites (electron donors) may be present. Coordinatively unsaturated Fe cation sites, for example, Fe_B sites on $Fe_3O_4(001)$, will act as a strong Lewis acid that leads to water dissociation.¹⁰ Anions on metal oxide surfaces are electron donors, *i.e.* Lewis bases, which can play a role in bond dissociation and recombination, facilitating elementary steps in a reaction on the surface.

The determination of the structure of Fe₃O₄(111), and therefore, understanding the chemical functionality of the surface, is challenging and remains highly debated.¹¹⁻¹⁷ Very similar unit cells with various possible terminations and multiple domains may coexist.^{12,18} There are three predominant preparation methods for Fe₃O₄(111) surfaces: (1) preparation of bulk Fe₃O₄(111);¹⁹⁻²² (2) growth of thin films of Fe₃O₄(111) on zero-valent^{8,23-25} and nonreducible surfaces,^{26,27} especially Pt;²⁸⁻³⁰ and (3) formation of thin films *via* reduction of α -Fe₂O₃(0001).^{17,31} The surface termination is different for the various preparations. Of these, bulk Fe₃O₄(111) prepared in vacuum possesses the most varieties of surface terminations.



View Article Online

View Journal | View Issue

^aDepartment of Chemistry and Chemical Biology, Harvard University, Cambridge, MA 02138, USA. E-mail: friend@fas.harvard.edu

^bDepartment of Physics, Harvard University, Cambridge, Massachusetts 02138, USA John A. Paulson School of Engineering and Applied Sciences, Harvard University, Cambridge, Massachusetts 02138, USA

[†] Electronic supplementary information (ESI) available: Supporting results and movie M-1 are available for this paper. See DOI: 10.1039/c9sc06149e

Tetrahedrally coordinated iron, Fe_{tet1}, with a hexagonal pattern, co-exists with octahedrally coordinated iron, Fe_{oct2},^{20,32} signified by the honeycomb structure which is often related to oxygenpoor conditions. Lattice oxygen, O, terminated bulk $Fe_3O_4(111)$ is also observed.³³ Additionally, oxygen adatoms have been proposed^{14,32,34} based on STM and chemical activity. The terminations of $Fe_3O_4(111)$ thin films on Pt are predominantly assigned as Fe_{tet1}^{23,28,35} based on scanning tunnelling microscopy (STM),23,28,35 dynamical low energy electron diffraction (LEED) intensity analysis,23,35 infrared reflection-absorption spectroscopy (IRRAS),²⁸ and density functional theory (DFT).²⁸ Thin films of $Fe_3O_4(111)$ formed from reduction of $Fe_2O_3(0001)$ are generally thought to be terminated by Fe_{tet1};^{9,31,36,37} however, the surface structures often co-exist with O terminated patches with a unique hexagonal periodicity of 0.3 nm,^{31,36,37} presumably due to the specific preparation conditions. Alternative preparation conditions lead to a surface terminated in the "biphase" structure in all of the three preparation methods,9,19,24,38,39 identified by STM, LEED and IRRAS. The "biphase" is a mixed oxide variously ascribed to FeO/Fe₃O₄ or Fe₃O₄/Fe₂O₃.^{13,40}

Herein, evidence is presented for a surface termination of $Fe_3O_4(111)$ films that has dual functionality, containing both Lewis acid and base sites. The surface studied here is terminated by a mixture of tetrahedral Fe_{tet1} and O adatoms, based on the registry of the overlayers with the surface and the differentiation of empty and filled states on different regions of the surface using STM and DFT. The surface is prepared by sputtering and annealing an α -Fe₂O₃(0001) single crystal in vacuum. The oxygen adatom layer was robust, even after 40+ preparation cycles in an oxygen-free environment, due to the high mobility of ions in the iron oxides. We further show that O adatoms and nearby uncovered Fe_{tet1} are Lewis acid–base pairs that actively dissociate methanol at room temperature. The results present insight into understanding the surface chemistry of magnetite films and reduced hematite catalysts.

Methods

Sample preparation

A natural α -Fe₂O₃(0001) single crystal was purchased from SurfaceNet GmbH. The sample was prepared using consecutive cycles of 15 min Ar⁺ sputtering and 20 min annealing at 980 K in UHV until a clear LEED pattern of Fe₃O₄(111) was detected.

Scanning tunnelling microscopy experiments

The STM experiments were carried out in a commercial Omicron VT Beam Deflection AFM/STM housed in an ultrahigh vacuum chamber equipped with LEED and direct dosing tube described in previous work.⁴¹ All images were scanned at room temperature using a commercially-cut Pt/Ir tip. Pressure of methanol vapor was read by an ion gauge placed far away from the sample. The actual methanol pressure above the sample surface during *in situ* experiments was estimated to be similar to the reading from the ion gauge, as a result of tip shading effect and direct dosing effect.⁴¹ All STM images were processed using SPIP 6.0.2, Scanning Probe Image Processor Software by Image Metrology.

X-ray photoelectron spectroscopy experiments

XPS experiments were conducted using a commercial SPECS instrument at the Center for Functional Nanomaterials at Brookhaven National Laboratory. The UHV XPS system consisted of an analysis chamber (base pressure $\sim \!\! 1 \times 10^{-9}$ Torr) and a preparation chamber (base pressure $\sim \!\! 5 \times 10^{-10}$ Torr). The X-ray source uses monochromatized Al K α radiation. The sample was irradiated under an angle of 43° and the photoelectrons were measured with normal emission using a hemispherical analyser at a pass-energy of 50 eV. Calibration was completed by setting the Ag 3d_{5/2} peak to 368.2 eV, the peak obtained using a reference Ag(111) crystal. Intensity was adjusted by normalizing the signal-noise ratio in baselines.

Temperature programmed reaction spectroscopy experiments

TPRS experiments were carried out in a UHV setup with a base pressure of $< 8 \times 10^{-11}$ Torr described before.⁴² Deuterated methanol (CD₃OH, Sigma Aldrich, 99.8 atom% D) was purified by freeze-pump-thaw cycles and was introduced to the surface with a needle doser at 130 K. The TPRS experiments were carried out with a heating rate of 1 K s⁻¹ and the resulting spectra were corrected for fragmentation pattern contributions.

Density functional theory calculations

DFT calculations were performed using VASP⁴³ with the projector-augmented wave potentials. The GGA-PBE⁴⁴ exchangecorrelation functional was used. The DFT-TS method⁴⁵ was used to include the van der Waals correction. The on-site Coulomb repulsion of Fe 3d electrons was treated by DFT + U⁴⁶ approach where $U_{\rm eff}$ is equal to 4.0 eV. The kinetic energy cut-off of the plane-wave basis sets was 550 eV. The optimized lattice constant of Fe₃O₄ was 8.440 Å, close to the experimental value of 8.396 Å. The Fe_{tet1}-terminated (111) surface was modelled by a slab structure, including 17 atomic layers and a more than 12 Å vacuum region along the *z* direction. A Gamma-centred $5 \times 5 \times 1$ *k*-point mesh⁴⁷ was utilized to sample the Brillouin zone of the $\sqrt{3} \times \sqrt{3}$ supercell (10.34 Å along the two in-plane lattice vectors). The ground-state ferrimagnetic phase⁴⁸ was used for the spin configuration.

Different adsorption sites of O on the surface were calculated to find the most stable configuration. During structural relaxation, the central 5 layers of the slab were constrained to their bulk positions, and the remaining atoms were fully relaxed to a force threshold of 0.01 eV $Å^{-1}$. The adsorption an oxygen adatom is calculated energy of as $E_{\text{ads}} = \left(E[nO/\text{Fe}_3O_4] - E[\text{Fe}_3O_4] - \frac{n}{2}E[O_2] \right) / n$, where *n* is the number of O adatoms in a unit cell. The STM images at \pm 1.5 eV tunnelling bias were simulated using the electronic states with energies ranging from the Fermi level $(E_{\rm F})$ to ± 1.5 eV. The atomic structures were visualized by QuteMol, and the simulated STM data were analysed using p4vasp with further smooth and colouring in SPIP 6.0.2.

Results and discussion

Identity of oxygen adatoms on Fe₃O₄(111)/α-Fe₂O₃(0001)

A Fe₃O₄(111) film forms on top of α -Fe₂O₃(0001) after repeated cycles of sputtering and annealing in UHV and is evident by the transformation of the LEED patterns from $(\sqrt{3} \times \sqrt{3})R30^\circ$, characteristic of α -Fe₂O₃(0001), to a Fe₃O₄(111) $p(2 \times 2)$ pattern (Fig. S1†).^{49,50} The top layers of the surface are reduced by sputtering but subsequently re-oxidized by annealing in vacuum to 980 K for 20 minutes according to X-ray photoelectron spectroscopy results (Fig. S2†). The increase in O : Fe ratio during annealing is attributed to diffusion of iron cations and interstitials into the bulk, as the cation diffusion in the inverse spinel structure is rapid and the oxygen lattice is rigid.⁵¹⁻⁵⁴

The surface termination of $Fe_3O_4(111)/\alpha$ - $Fe_2O_3(0001)$ contains an unsaturated close-packed hexagonal pattern, which is assigned as an oxygen adatom layer (Fig. 1). The hexagonal pattern, which is reproducible and covers most of terraces across the surface, has a spacing of ~0.6 nm along the closepacked directions and occupies a-top positions, templated by the substrate (Fig. 1A). The apparent height between the same feature on the two adjacent terraces is ~0.48 nm, corresponding to a repeating distance of layers within $Fe_3O_4(111)$. The a-top stacked position indicates that a new layer, other than those from bulk, is formed. The lateral periodicity of 0.6 nm rules out the termination of lattice layer O_1 , and the absence of honeycomb structure under all scanning conditions rules out the Fe_{oct2} termination.



Fig. 1 The atomic-scale structure of $Fe_3O_4(111)/\alpha$ - $Fe_2O_3(0001)$ is determined using: (A) an STM image of an oxygen adatom terminated $Fe_3O_4(111)$ film, and (B) the corresponding DFT slab models with O adsorption coverage of 0.67 ML. The insert in (A) is a magnified image with close-packed directions of substrate indicated in black dashed lines. Scanning conditions: -1.5 V, 0.02 nA.

The surface in this work has a different topography and electronic structure compared to previous studies of $Fe_3O_4(111)$.^{28,36,55} Although the surface prepared herein (Fig. 1) may appear qualitatively similar to images in the literature assigned to a termination by Fe_{tet1} co-existing with lattice O that has a 0.3 nm spacing,^{36,56} it is clearly different, as discussed in more detail below. In this work, the top layer occupies an a-top position and is only visible in filled states in STM, different from the Fe_{tet1} layer that contains both filled and empty states³⁶ (Fig. 2A and S3[†]). The top layer in this work is not related to adsorbates from background,²⁰ as the coverage does not increase with longer time exposure in UHV. Thus, the surface termination of the $Fe_3O_4(111)$ film formed from reduction of oxygen-rich bulk $Fe_2O_3(0001)$ is intrinsically different from the



Fig. 2 Identification of oxygen adatoms and uncovered Fe_{tet1} by their distinct electronic states. (A) Experimental STM images showing a biasdependent contrast change. (B) Simulated STM images based on DFT calculations. (C) DFT calculated site-projected density of states (PDOS) of oxygen adatoms (O_{ad}) and uncovered Fe_{tet1} atoms on the top layer. The PDOS includes both spin up and spin down components (*i.e.* not spin-resolved). The Fermi level is shifted to 0.0 eV. Experimental tunnelling current was 0.25 nA; probing bias was indicated in each panel. The regions encompassed by the dashed lines on the DFT model are a visual guide.

 $Fe_3O_4(111)$ terminations described in previous literature, most likely due to different preparation conditions.

Considering the re-oxidation during annealing, the surface structure is ascribed to a mixture of oxygen adatoms on the Fettet1 substrate and exposed Fettet1 sites. Under the preparation conditions used here, the coverage of the oxygen adatoms measured by STM is 0.5–0.7 ML, typically at \sim 0.64 ML (1 monolayer, ML, is defined as an equivalent number of Fe_{tet1} atoms in a primitive unit cell). The O_{ad} atoms are templated by Fe_{tet1}, forming patches that strictly follow the Fe_{tet1} geometry at high coverages. Different annealing temperatures (720-1010 K) in UHV do not lead to a change in the Oad structure although the population of O adatoms may vary. Oxidation in 5×10^{-7} mbar in O2 during annealing followed by a cleaning cycle results in formation of clusters, possibly oxide islands, and a slightly higher O_{ad} coverage. Some features at the edge of close-packed O adatom patches are mobile at room temperature (Movie, M-1, in the ESI[†]), possibly due to surface hydroxyls or reversible dissociative adsorption of trace background water vapor.57,58

The proposed structure is validated by DFT calculations on 0.67 ML oxygen adatoms covered Fettet1 terminated surface which show that the a-top position is the most stable adsorption site among others (Fig. 1B). Other oxygen adsorption sites considered have adsorption energies weaker by at least 1.48 eV per O adatom (Fig. S4[†]). The adsorption energy of the a-top oxygen (Fig. 1B) is 0.24 eV per O adatom. This value becomes negative when using gas-phase O2 as a reference and considering its correction in the DFT-calculated energy (there are two different binding energies often used in the literature for O₂ (0.4 48 or 0.68 59 eV) leading to a binding energy of O of either -0.16 or -0.44 eV, respectively). The calculational results are consistent with prior theoretical work that both exposed Fe_{tet1} and O-adsorbed Fe_{tet1} are stable terminations.⁴⁸ The calculated bond length of O_{ad}-Fe_{tet1} is 1.62 Å (Fig. 1B), which is substantially shorter than the lattice Fe_{tet1}-O bond length of 1.90 Å, and is comparable with the length of a Fe=O bond on ferryl oxygen terminated α -Fe₂O₃(0001) of 1.58 Å.⁶⁰ Here, the O_{ad} is not bound to a local α-Fe₂O₃(0001), as measured periodicity suggests a Fe₃O₄(111) surface. Thus, the O_{ad} is strongly bonded to the lattice Fe_{tet1} on a $Fe_3O_4(111)$ surface.

The assignment of oxygen adatoms is further confirmed by agreement between STM measurements and DFT calculations, including the bias dependence of the images. Similar biasdependent features that are visible in filled states (negative bias) were reported previously; however, they are either ascribed as lattice oxygen layers with a 0.3 nm periodicity³⁶ or as isolated OH or H adsorbates.20 In this work, the filled state image of oxygen adatoms are bright under a scanning voltage of -1.5 V (sample biased), whereas the inverted contrast was observed when switching the scanning voltage to +1.5 V (Fig. 2A, and a larger scale comparison in Fig. S3[†]). The contrast corresponds to a change in density of states, which is calculated using DFT. Based on models in Fig. 1B, the projected density of states (PDOS) of oxygen adatoms (O_{ad}) and surface iron atoms (Fe_{tet1}) that were not covered by oxygen adatoms were calculated and found to be well isolated (Fig. 2C): Oad occupies two bands at about -0.5 V and -1.5 V, none of which was occupied by Fe_{tet1}; Fe_{tet1} occupies three bands at about +0.5 V, +1.0 V and +1.7 V, neither of which was occupied by O_{ad} . The distribution of PDOS can be used as fingerprint for element identity. The separated PDOS leads to a contrast change of O_{ad} and Fe_{tet1} in simulated STM images (Fig. 2B), confirming the generality for the observed surface property by STM (Fig. 2A). Further, the calculated PDOS of O_{ad} coverages of 0, 0.67, 0.75, and 1 ML suggests an O_{ad} coverage dependency, reinforcing the ability of oxygen adatoms to change the surface electronic properties (Fig. S5†).

The active O_{ad} and Fe_{tet1} interface for methanol dissociation at room temperature

Selective oxidation of methanol to formaldehyde occurs on the as-prepared Fe₃O₄(111)/ α -Fe₂O₃(0001) with methoxy as an intermediate, indicated by temperature programmed reaction (Fig. 3). No other product, including CO₂, is detected. Deuterated methanol, CD₃OH, was used to differentiate O–H and C–H bonds. Desorption of a surface-bound layer of CD₃OH was observed in a peak at 250 K, accompanied by H₂O evolution from disproportionation reaction of hydroxyls and/or original water impurities in deuterated methanol. Carbon–deuterium bond cleavage starts at 600 K and peaks at 700 K, leading to the formation of formaldehyde, CD₂O, methanol-d₃, and deuterated water (HDO and D₂O). This elementary step corresponds to an approximate activation barrier of 179 kJ mol⁻¹ calculated assuming first-order kinetics,⁶¹ and a pre-exponential factor of 10^{12} s^{-1} .⁶² At this temperature, trace amounts of surface



Fig. 3 (A) Temperature programmed reaction data for methanol- d_3 (CD₃OH) show that methanol is dissociated and subsequently oxidized on as-prepared Fe₃O₄(111)/ α -Fe₂O₃(0001). Traces of water, formal-dehyde and methanol were tracked by mass-ion ratio indicated in the panel and corrected for fragmentation patterns. The heating rate is 1 K s⁻¹. (B) The reaction mechanism is illustrated in a schematic.

hydroxyl OH contribute to a desorption peak of H₂O in the high temperature oxidation feature. These data show that methoxy (CD₃O) is formed at lower temperature in competition with methanol desorption; ultimately yielding CD₂O and CD₃OH at high temperature. Similar reactivity was previously reported on $Fe_3O_4(111)$.²²

The formation of methoxy on $Fe_3O_4(111)$, which is rich in Lewis acid-base pairs, was captured by in situ STM at room temperature (Fig. 4). The surface covered by O_{ad} is shown in Fig. 4A. The atomic structure of the substrate (Fig. 4B) is not resolved during methanol exposure to the surface at room temperature due to the presence of weakly-bound methanol which is mobile on the surface, thus, producing a fuzzy image. After 21 min under 4 \times 10⁻¹⁰ mbar methanol, the immobile bright features are attributed to methoxy formed from methanol dissociation. These features, as a single methoxy or a few in a group, have an apparent width from 1.0 to 3.0 nm and an apparent height of \sim 0.3 nm (Fig. 4B). These methoxy species are stable on the surface after heating to 400 K (Fig. 4C), consistent with temperature programmed reaction results. The oxidation of methanol consumes oxygen from magnetite to form water through the Mars-van Krevelen mechanism. However, the oxygen adatoms reappear after heating the surface to 720 K (Fig. 4D), due to quick ion exchange with bulk and replenishment of O_{ad} at the elevated temperature.

The as-prepared $Fe_3O_4(111)$ surface contains O_{ad} - Fe_{tet} interfaces, *i.e.* Lewis acid–base pairs, where methanol oxidation occurs readily. The active site for the first elementary step for methanol oxidation can be tracked by methoxy formation. These sites include both oxygen adatoms and nearby uncovered lattice iron measured by *in situ* STM (white circles in Fig. 4). In contrast, small domains with an alternative compact structure



Fig. 4 In situ STM of methanol dissociation on Fe₃O₄(111) films at room temperature tracks the location of active sites. The same surface was imaged (A) before and (B) after 21 min exposure of methanol. $P_{MeOH} = 4 \times 10^{-10}$ mbar (after background subtraction); scanning conditions: -1.5 V, 0.015 nA; scanning from bottom to top. White circles highlight the same positions of methoxy species and interfaces of Fe_{tet} (dark in panel A) and O_{ad} (bright in panel A). (C) and (D) are *ex situ* STM of the post methanol surface after being annealed to 400 K and 720 K, respectively.

do not contain the O_{ad} -Fe_{tet} interfaces is inert to O–H bond cleavage, as no methoxy was observed in these areas (Fig. S6†). The alternative compact structures are commonly assigned as O-terminated FeO islands,^{9,37} oxygen-terminated oxide surfaces,⁶³ and Fe_{1-x}O islands.^{19,40}

Additionally, the $Fe_3O_4(111)$ surface is active for other reactions, including water dissociation and formic acid decomposition.⁶⁴ Fe₃O₄(111) films and bulk are known to dissociate water based on experimental and theoretical studies that dissociated OH⁻ and H⁺ bound to surface Fe cations and O anions respectively.^{14,29,31,65} The dissociated OH⁻ can further react with non-dissociated water to form complexes on a Fe_{tet1} terminated films at low temperatures.57,66 The ring-like waterhydroxyl complexes, or (2×2) super structure, also forms but less stably⁶⁷ on $(\sqrt{2} \times \sqrt{2})R45^{\circ}$ -reconstructed surfaces of bulk Fe₃O₄(001) single crystals.⁶⁸ At room temperature, the Fe₃O₄(001) surface is only covered by hydroxyls, which desorb as water at 520 K by abstraction of lattice oxygen.69 The reaction with water becomes more extensive on both (111) and (001) terminations with an increase of vapor pressure,21 and can lead to a lifting of the subsurface cation vacancy reconstruction on Fe₃O₄(001) surfaces.⁷⁰

Conclusions

A well-ordered $Fe_3O_4(111)$ film-terminated hematite is formed through reduction and heating. The surface termination is a mixture of O adatoms (O_{ad}) and uncovered lattice Fe_{tet} , yielding Lewis base and acid sites, respectively. Reduced top layers are oxidized by annealing to 980 K *in vacuo*, an oxygen deficient environment. Rapid ion exchange occurs at these temperatures,⁵² especially cation diffusion, and oxygen-rich bulk leads to re-oxidation of surface layers. Further investigations on electronic structure by STM and DFT show a well separated density of states on O_{ad} and Fe_{tet} , allowing identification of the two species.

The mixture of O_{ad} and Fe_{tet} sites impacts reactivity of chemical reactions that require both acid and base sites, illustrated by CD_3OH oxidation. The surface is active for dissociation of the O–H bond near room temperature, to yield methoxy. In this process, the H can be transferred to the O adatoms to yield adsorbed OH and methoxy is bound to the exposed Fe_{tet} sites. The resulting methoxy is relatively immobile on the surface at room temperature, based on STM imaging. The O_{ad} act as proton acceptors in the dissociative adsorption of methanol. The Lewis acidity of Fe_{tet1} is theoretically predicted to accommodate methoxy species.⁷¹ Further oxidation with C–D cleavage occurs at 700 K, leading to the formation of formal-dehyde and water.

The fundamental understanding of $Fe_3O_4(111)/\alpha$ -Fe₂O₃(0001) from in this work contributes to the mechanistic study of magnetite as a material for applications at surfaces and interfaces, especially for heterogeneous catalysis. Fe₃O₄(111), the most thermally stable facet, will dominate the termination of magnetite under equilibrium conditions. As surface termination can affect the present of Lewis acid–base pairs,⁷² work herein with a bulk of iron oxide is closely related to the magnetite catalysts or hematite catalysts under reducing conditions.

The dual Lewis site-meaning the structure contains an adjacent Lewis acid and base pair-is a promising feature in designing efficient, low-cost, and robust heterogeneous catalysts. The synergistic interactions between intermediates adsorbed on acid and base sites enhance catalytic efficiency in redox reactions, such as the industrially-important production of formaldehyde from methanol on iron oxide-based materials.73,74 In this regard, the Lewis base facilitates the formation of the oxidation intermediate at room temperature by O-H bond splitting, while the Lewis acid is the reaction of the final oxidation step. Designing catalyst materials with both Lewis sites in close proximity is a promising principle for multistep oxidation reactions of alcohols. Additionally, compared with catalysts that need a second metal component to form dual Lewis sites,75 the catalyst herein is free of sintering concerns at elevated temperatures.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was primarily supported by the U.S. Army Research Office under Grant No. W911NF1820143. The calculations were performed at the Oak Ridge Leadership Computing Facility (OLCF) and the National Energy Research Scientific Computing Center (NERSC), while XPS instrumentation support was provided by the Center for Functional Nanomaterials, Brookhaven National Laboratory, of the U.S. Department of Energy. Work by W. C. was supported by the Integrated Mesoscale Architectures for Sustainable Catalysis (IMASC), an Energy Frontier Research Center funded by the U.S. Department of Energy, Office of Science, Basic Energy Sciences under Award # DE-SC0012573. We would like to acknowledge Dr Robert J. Madix for his contributions in discussions.

References

- 1 Z. Wu, M. Li and S. H. Overbury, J. Catal., 2012, 285, 61-73.
- 2 M. Nolan, S. C. Parker and G. W. Watson, *Surf. Sci.*, 2005, **595**, 223–232.
- 3 W.-C. Huang, L.-M. Lyu, Y.-C. Yang and M. H. Huang, J. Am. Chem. Soc., 2012, 134, 1261–1267.
- 4 L. M. Rossi, A. D. Quach and Z. Rosenzweig, *Anal. Bioanal. Chem.*, 2004, **380**, 606–613.
- 5 W.-M. Zhang, X.-L. Wu, J.-S. Hu, Y.-G. Guo and L.-J. Wan, *Adv. Funct. Mater.*, 2008, **18**, 3941–3946.
- 6 M. Bibes and A. Barthelemy, *IEEE Trans. Electron Devices*, 2007, 54, 1003–1023.
- 7 M. Hua, S. Zhang, B. Pan, W. Zhang, L. Lv and Q. Zhang, J. Hazard. Mater., 2012, 211–212, 317–331.
- 8 Y. S. Dedkov, M. Fonin, D. V. Vyalikh, J. O. Hauch, S. L. Molodtsov, U. Rudiger and G. Guntherodt, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2004, 70, 073405.

- 9 Y. Tang, H. Qin, K. Wu, Q. Guo and J. Guo, *Surf. Sci.*, 2013, **609**, 67–72.
- 10 N. Mulakaluri, R. Pentcheva and M. Scheffler, *J. Phys. Chem. C*, 2010, **114**, 11148–11156.
- 11 H. Sträter, H. Fedderwitz, B. Groß and N. Nilius, J. Phys. Chem. C, 2015, 119, 5975–5981.
- 12 G. S. Parkinson, Surf. Sci. Rep., 2016, 71, 272-365.
- 13 N. G. Condon, F. M. Leibsle, A. R. Lennie, P. W. Murray,
 D. J. Vaughan and G. Thornton, *Phys. Rev. Lett.*, 1995, 75, 1961–1964.
- 14 R. S. Cutting, C. A. Muryn, D. J. Vaughan and G. Thornton, *Surf. Sci.*, 2008, **602**, 1155–1165.
- 15 K. Adib, N. Camillone III, J. P. Fitts, K. T. Rim, G. W. Flynn, S. A. Joyce and R. M. Osgood Jr, *Surf. Sci.*, 2002, **497**, 127–138.
- 16 N. Camillone III, K. Adib, J. P. Fitts, K. T. Rim, G. W. Flynn, S. A. Joyce and R. M. Osgood, *Surf. Sci.*, 2002, **511**, 267–282.
- 17 K. Adib, G. G. Totir, J. P. Fitts, K. T. Rim, T. Mueller, G. W. Flynn, S. A. Joyce and R. M. Osgood Jr, *Surf. Sci.*, 2003, 537, 191–204.
- 18 H. Kuhlenbeck, S. Shaikhutdinov and H.-J. Freund, *Chem. Rev.*, 2013, **113**, 3986–4034.
- 19 M. Paul, M. Sing, R. Claessen, D. Schrupp and V. A. M. Brabers, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2007, **76**, 075412.
- 20 T. K. Shimizu, J. Jung, H. S. Kato, Y. Kim and M. Kawai, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2010, **81**, 235429.
- 21 T. Kendelewicz, P. Liu, C. S. Doyle, G. E. Brown Jr,
 E. J. Nelson and S. A. Chambers, *Surf. Sci.*, 2000, 453, 32–46.
- 22 Z. Li, D. V. Potapenko, K. T. Rim, M. Flytzani-Stephanopoulos, G. W. Flynn, R. M. Osgood, X.-D. Wen and E. R. Batista, *J. Phys. Chem. C*, 2015, **119**, 1113–1120.
- 23 M. Ritter and W. Weiss, Surf. Sci., 1999, 432, 81-94.
- 24 F. Genuzio, A. Sala, T. Schmidt, D. Menzel and H.-J. Freund, J. Phys. Chem. C, 2014, 118, 29068–29076.
- 25 H. Xiang, F. Shi, M. S. Rzchowski, P. M. Voyles and Y. A. Chang, *Appl. Phys. Lett.*, 2010, **97**, 092508.
- 26 J. B. Moussy, S. Gota, A. Bataille, M. J. Guittet, M. Gautier-Soyer, F. Delille, B. Dieny, F. Ott, T. Doan, P. Warin, P. Bayle-Guillemaud, C. Gatel and E. Snoeck, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2004, **70**, 174448.
- 27 H. Hong, J. Kim, X. Fang, S. Hong and T.-C. Chiang, *Appl. Phys. Lett.*, 2017, **110**, 021601.
- 28 X. Li, J. Paier, J. Sauer, F. Mirabella, E. Zaki, F. Ivars-Barceló, S. Shaikhutdinov and H. J. Freund, *J. Phys. Chem. B*, 2018, 122, 527–533.
- 29 Y. Joseph, W. Ranke and W. Weiss, *J. Phys. Chem. B*, 2000, **104**, 3224–3236.
- 30 U. Leist, W. Ranke and K. Al-Shamery, *Phys. Chem. Chem. Phys.*, 2003, **5**, 2435–2441.
- 31 K. T. Rim, D. Eom, S.-W. Chan, M. Flytzani-Stephanopoulos, G. W. Flynn, X.-D. Wen and E. R. Batista, *J. Am. Chem. Soc.*, 2012, **134**, 18979–18985.
- 32 A. R. Lennie, N. G. Condon, F. M. Leibsle, P. W. Murray, G. Thornton and D. J. Vaughan, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1996, 53, 10244–10253.
- 33 N. Berdunov, S. Murphy, G. Mariotto and I. V. Shvets, *Phys. Rev. Lett.*, 2004, **93**, 057201.

- 34 R. S. Cutting, C. A. Muryn, G. Thornton and D. J. Vaughan, Geochim. Cosmochim. Acta, 2006, **70**, 3593–3612.
- 35 S. K. Shaikhutdinov, M. Ritter, X. G. Wang, H. Over and W. Weiss, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1999, **60**, 11062–11069.
- 36 K. T. Rim, J. P. Fitts, T. Müller, K. Adib, N. Camillone III, R. M. Osgood, S. A. Joyce and G. W. Flynn, *Surf. Sci.*, 2003, 541, 59–75.
- 37 K. T. Rim, T. Müller, J. P. Fitts, K. Adib, N. Camillone,
 R. M. Osgood, E. R. Batista, R. A. Friesner, S. A. Joyce and
 G. W. Flynn, *J. Phys. Chem. B*, 2004, **108**, 16753–16760.
- 38 G. Ketteler, W. Weiss, W. Ranke and R. Schlögl, *Phys. Chem. Chem. Phys.*, 2001, 3, 1114–1122.
- 39 L. Schöttner, A. Nefedov, C. Yang, S. Heissler, Y. Wang and C. Wöll, *Front. Chem.*, 2019, 7, 451.
- 40 N. G. Condon, F. M. Leibsle, T. Parker, A. R. Lennie, D. J. Vaughan and G. Thornton, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1997, 55, 15885–15894.
- 41 F. Xu, I. Fampiou, C. R. O'Connor, S. Karakalos, F. Hiebel,
 E. Kaxiras, R. J. Madix and C. M. Friend, *Phys. Chem. Chem. Phys.*, 2018, 20, 2196–2204.
- 42 C. A. Walenta, A. S. Crampton, F. Xu, U. Heiz and C. M. Friend, *J. Phys. Chem. C*, 2018, **122**, 25404–25410.
- 43 G. Kresse and J. Furthmüller, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1996, **54**, 11169–11186.
- 44 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, 77, 3865–3868.
- 45 A. Tkatchenko and M. Scheffler, *Phys. Rev. Lett.*, 2009, **102**, 073005.
- 46 S. L. Dudarev, G. A. Botton, S. Y. Savrasov, C. J. Humphreys and A. P. Sutton, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1998, 57, 1505–1509.
- 47 M. Methfessel and A. T. Paxton, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1989, **40**, 3616–3621.
- 48 J. Noh, O. I. Osman, S. G. Aziz, P. Winget and J.-L. Brédas, *Chem. Mater.*, 2015, **27**, 5856–5867.
- 49 C. A. Walenta, A. S. Crampton, F. Xu, U. Heiz and C. M. Friend, *J. Phys. Chem. C*, 2018, **122**, 25404–25410.
- 50 W. Weiss and M. Ritter, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1999, **59**, 5201–5213.
- 51 S. Hallström, L. Höglund and J. Ågren, *Acta Mater.*, 2011, **59**, 53–60.
- 52 S. Nie, E. Starodub, M. Monti, D. A. Siegel, L. Vergara, F. El Gabaly, N. C. Bartelt, J. de la Figuera and K. F. McCarty, *J. Am. Chem. Soc.*, 2013, **135**, 10091–10098.
- 53 Z. D. Sharp, Geology, 1991, 19, 653-656.
- 54 R. Dieckmann and H. Schmalzried, *Berichte der Bunsengesellschaft für physikalische Chemie*, 1977, 81, pp. 414–419.

- 55 C. Bamroongwongdee, M. Bowker, A. F. Carley, P. R. Davies, R. J. Davies and D. Edwards, *Faraday Discuss.*, 2013, **162**, 201–212.
- 56 K. T. Rim, D. Eom, L. Liu, E. Stolyarova, J. M. Raitano, S.-W. Chan, M. Flytzani-Stephanopoulos and G. W. Flynn, *J. Phys. Chem. C*, 2009, **113**, 10198–10205.
- 57 E. Zaki, F. Mirabella, F. Ivars-Barceló, J. Seifert, S. Carey, S. Shaikhutdinov, H.-J. Freund, X. Li, J. Paier and J. Sauer, *Phys. Chem. Chem. Phys.*, 2018, **20**, 15764–15774.
- 58 P. Dementyev, K. H. Dostert, F. Ivars-Barcelo, C. P. O'Brien, F. Mirabella, S. Schauermann, X. Li, J. Paier, J. Sauer and H. J. Freund, *Angew. Chem., Int. Ed. Engl.*, 2015, 54, 13942– 13946.
- 59 L. Wang, T. Maxisch and G. Ceder, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2006, 73, 195107.
- 60 C. Lemire, S. Bertarione, A. Zecchina, D. Scarano, A. Chaka, S. Shaikhutdinov and H. J. Freund, *Phys. Rev. Lett.*, 2005, 94, 166101.
- 61 P. A. Redhead, Vacuum, 1962, 12, 203-211.
- 62 R. J. Madix and S. G. Telford, Surf. Sci., 1995, 328, L576–L581.
- 63 A. Sala, H. Marchetto, Z. H. Qin, S. Shaikhutdinov, T. Schmidt and H. J. Freund, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2012, 86, 155430.
- 64 J. Balajka, M. A. Hines, W. J. I. DeBenedetti, M. Komora, J. Pavelec, M. Schmid and U. Diebold, *Science*, 2018, 361, 786–789.
- 65 Y. Joseph, C. Kuhrs, W. Ranke, M. Ritter and W. Weiss, *Chem. Phys. Lett.*, 1999, **314**, 195–202.
- 66 F. Mirabella, E. Zaki, F. Ivars-Barceló, X. Li, J. Paier, J. Sauer, S. Shaikhutdinov and H.-J. Freund, *Angew. Chem., Int. Ed.*, 2018, 57, 1409–1413.
- 67 E. Zaki, Z. Jakub, F. Mirabella, G. S. Parkinson, S. Shaikhutdinov and H.-J. Freund, *J. Phys. Chem. Lett.*, 2019, **10**, 2487–2492.
- 68 M. Meier, J. Hulva, Z. Jakub, J. Pavelec, M. Setvin, R. Bliem, M. Schmid, U. Diebold, C. Franchini and G. S. Parkinson, *Proc. Natl. Acad. Sci. U. S. A.*, 2018, **115**, E5642–E5650.
- 69 G. S. Parkinson, Z. Novotný, P. Jacobson, M. Schmid and U. Diebold, J. Am. Chem. Soc., 2011, 133, 12650–12655.
- 70 B. Arndt, M. Creutzburg, E. Grånäs, S. Volkov, K. Krausert, A. Vlad, H. Noei and A. Stierle, *J. Phys. Chem. C*, 2019, **123**, 26662–26672.
- 71 X. Li and J. Paier, J. Phys. Chem. C, 2019, 123, 8429-8438.
- 72 S. Zhang, Z.-Q. Huang, Y. Ma, W. Gao, J. Li, F. Cao, L. Li, C.-R. Chang and Y. Qu, *Nat. Commun.*, 2017, 8, 15266.
- 73 A. P. V. Soares, M. F. Portela and A. Kiennemann, *Catal. Rev.*, 2005, 47, 125–174.
- 74 C. Brookes, P. P. Wells, G. Cibin, N. Dimitratos, W. Jones, D. J. Morgan and M. Bowker, ACS Catal., 2014, 4, 243–250.
- 75 V. K. Díez, C. R. Apesteguía and J. I. Di Cosimo, *Catal. Today*, 2000, **63**, 53–62.