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Catalytic formation of oxalic acid on the partially oxidised greigite $Fe_3S_4(001)$ surface

David Santos-Carballal,^{1,*} and Nora H de Leeuw^{1,2}

4 Abstract:

Greigite (Fe₃S₄), with its ferredoxin-like 4Fe-4S redox centres, is a naturally occurring mineral 5 6 capable of acting as a catalyst in the conversion of carbon dioxide (CO₂) into low molecular-7 weight organic acids (LMWOAs), which are of paramount significance in several soil and plant 8 processes as well as in the chemical industry. In this paper, we report the reaction between CO_2 and water (H₂O) to form oxalic acid (H₂C₂O₄) on the partially oxidised greigite Fe₃S₄(001) 9 10 surface by means of spin-polarised density functional theory calculations with on-site 11 Coulomb corrections and long-range dispersion interactions (DFT+U-D2). We have calculated the bulk phase of Fe₃S₄ and the two reconstructed Tasker type 3 terminations of its (001) 12 surface, whose properties are in good agreement with available experimental data. We have 13 obtained the relevant phase diagram, showing that the Fe₃S₄(001) surface becomes 62.5% 14 partially oxidised, by replacing S by O atoms, in the presence of water at the typical conditions 15 of calcination [Mitchell et al. Faraday Discuss. 2021, 230, 30-51]. The adsorption and co-16 adsorption of the reactants on the partially oxidised Fe₃S₄(001) surface are exothermic 17 18 processes. We have considered three mechanistic pathways to explain the formation of $H_2C_2O_4$, showing that the coupling of the C–C bond and second protonation are the 19 20 elementary steps with the largest energy penalty. Our calculations suggest that the partially oxidised $Fe_3S_4(001)$ surface is a mineral phase that can catalyse the formation of $H_2C_2O_4$ under 21 22 favourable conditions, which has important implications for natural ecosystems and is a process that can be harnessed for the industrial manufacture of this organic acid. 23

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24 **1. Introduction**

Oxalic acid (H₂C₂O₄), which is the smallest dicarboxylic low molecular weight organic acid 25 (LMWOA),¹ has traditionally been regarded as a major player in the chemical industry^{2,3} and 26 natural systems.⁴ Oxalic acid is widely used for the treatment of metallic surfaces⁵⁻⁸ and 27 textiles,^{9–11} as a bleaching agent,^{12–15} for the preparation of esters,^{16–18} as a reagent in 28 chemical synthesis^{19–21} and for separation of rare-earth elements.^{22–25} The most sophisticated 29 methods currently used in the industrial production of oxalic acid involve the oxidation of 30 carbohydrates,^{26–29} ethylene glycol^{30,31} and carbon monoxide.³² Moreover, oxalic acid is the 31 metabolic product of plants,^{33–35} soil fungi^{36–38} and bacteria,^{39–41} and to a lesser extent of 32 animals,^{42–44} playing a key role in modulating the interactions between these organisms.^{4,45,46} 33 Three main pathways have been suggested for the biosynthesis of oxalic acid, including the 34 cytoplasmatic pathway,⁴⁷ the tricarboxylic acid pathway⁴⁸ and the glyoxylate pathway.^{36,49,50} 35 From a catalytic point of view, ferredoxins have gained renewed significance owing to their 36 biological role in electron transfers in several metabolic reactions.^{51,52} Ferredoxins are iron-37 sulfur proteins, with one of the most notable families containing cubane Fe₄-S₄ catalytic 38 centres.⁵³ The catalytic centre of ferredoxins can be either exposed or hidden to solvent, 39 leading to low reduction potentials with a minimum of -600 mV or high potentials with a 40 maximum of +350 mV, respectively.^{53,54} Structurally, ferredoxins contain Fe ions in the 41 oxidation states +2 and +3, which are forming tetrahedrally coordinated FeS₄ units, whereas 42 each S atom is bonded to three Fe ions.⁵⁴ Ferredoxin oxidoreductases are responsible for the 43 coupling of the C-C from CO₂ at cellular level in the reductive tricarboxylic acid cycle.⁵⁵ 44 45 Attempts to synthesise biomimetic Fe₄-S₄ clusters have been successful in controlling the binding of substrate through remote steric effects, but their preparation requires carefully 46 controlled conditions, toxic chemicals and complex steps.⁵⁶ 47

Greigite (Fe₃S₄) is a naturally occurring mineral that contains Fe₄–S₄ clusters with the cubane structure,⁵⁷ very similar to the catalytic centre in ferredoxins. Unsurprisingly, this mineral phase has been found to be catalytically active towards the activation and conversion of CO₂ into small organic molecules, including formic acid, acetic acid, pyruvic acid and methanol.^{58–} Recent studies have suggested that the surfaces of Fe₃S₄ become partially oxidised easily in air or wet conditions, forming core-shell iron sulfide-iron oxide nanoparticles,^{61,62} which enhances the catalytic activity.

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The important biocatalytic role of ferredoxins and the structural similarities with Fe₃S₄ have 55 led us to study the catalytic conversion of CO₂ and H₂O into H₂C₂O₄ over the partially oxidised 56 Fe₃S₄(001) surface by means of spin polarised density functional theory (DFT) calculations. 57 58 We have modelled the bulk phase of Fe₃S₄, as well as the two reconstructed Tasker type 3 59 terminations of its (001) surface, and discuss the changes in several structural, electronic and 60 magnetic properties. We have also calculated the effect of the ratio of the partial pressures of H₂O, *i.e.* the reactant, and H₂S, *i.e.* the product of the oxidation, and temperature on the 61 oxygen coverage of the Fe₃S₄(001) surface. Next, we have considered the adsorption of the 62 63 single CO_2 and H_2O molecules at different sites on the partially oxidised $Fe_3S_4(001)$ surface 64 and assessed a number of descriptors of the chemical activation. We have calculated the 65 structures and energies of possible surface-bound intermediates and saddle points and constructed the reaction profiles for three mechanisms, *i.e. via* (*i*) oxalate, (*ii*) carboxylate and 66 67 (*iii*) bicarbonate pathways.

68 **2. Computational Methods**

69 2.1. Calculation Details

The bulk phase, pristine and partially oxidised (001) surfaces of Fe₃S₄, as well as the reaction 70 mechanisms were calculated using unrestricted DFT calculations as implemented in the 71 Vienna *ab initio* simulation package (VASP).^{63–66} The strongly constrained and appropriately 72 73 normed (SCAN) functional, which meets all 17 known exact constraints within the metageneralised gradient approximation (meta-GGA), was used for the exchange-correlation 74 energy functional.⁶⁷ *Meta*-GGA functionals, which comprise up to the second order derivative 75 of the electron density, *i.e.* the non-interacting kinetic energy density term, provide energies 76 and structures that are in excellent agreement with experiments.^{68–72} The projector 77 augmented wave (PAW) formalism was used to treat the core states of Fe: [Ar], O: [He], C: 78 [He] and S: [Ne], their kinetic energies and the interactions with the valence orbitals.^{73,74} The 79 80 aspherical components of the density gradient were also considered within the PAW onecentre terms, which is required by *meta*-GGA functionals to describe properly the strongly 81 directional Fe 3d states. The D2 semiempirical method of Grimme was included in our 82 calculations to correct the long-range dispersion interactions,⁷⁵ which is particularly 83 important for an appropriate description of materials and interface properties. 59,68,76-85 A 84

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periodic plane-wave basis set with an upper kinetic energy threshold fixed at 400 eV was 85 employed to expand the Kohn-Sham valence states. The electronic ground state was 86 determined using a preconditioned conjugate gradients minimisation algorithm, which 87 88 optimises completely and self-consistently the total (free) energy, which is the variational quantity within this methodology.^{86–88} This stable and robust electronic minimisation 89 technique is preferred for modelling surface slabs with vacuum gaps when using meta-GGA 90 91 functionals, as it applies an all band simultaneous update of orbitals. The time step needed to carry out the line optimisation of the energy along the preconditioned conjugate gradient 92 93 was scaled automatically. The optimisation of the electronic density was terminated when 94 the energy difference between two consecutive self-consistent loop steps was smaller than 10⁻⁵ eV. The internal coordinates were relaxed to their equilibrium positions using a Newton 95 line optimiser, an efficient conjugate gradients technique,^{89,90} which was stopped when the 96 Hellmann-Feynman forces were below 0.01 eV $Å^{-1}$. 97

98 We have also used a mean-field Hubbard model, *i.e.* the so-called DFT + U method, to enhance the calculation of the delocalised Fe 3d states.^{91–93} We have applied a simplified rotationally 99 invariant strong intra-atomic Coulomb interaction⁹⁴ and developed the value for the effective 100 parameter U_{eff} = 5.26 eV following the linear response approach of Cococcioni *et al.*,⁹⁵ which 101 has been used successfully for the simulation of (Ni,Fe)oxyhydroxides (Ni_{1-x}Fe_xOOH)⁹⁶ and 102 defective ceria CeO₂.⁹⁷ We have carried out a series of static calculations to determine the 103 $U_{\rm eff}$ value, where we used the conventional cubic unit cell of Fe₃S₄ with ferrimagnetic ordering 104 with the experimental lattice parameter and internal positions.⁵⁷ We broke the symmetry of 105 the tetrahedral (Fe_A) and octahedral Fe (Fe_B) sublattices one at a time, by treating one of these 106 atoms differently to the rest. We calculated the number of d electrons (N_d) for (i) a standard 107 108 DFT, (ii) a non-self-consistent response function (NSCRF) and (iii) a self-consistent response 109 function (SCRF) to various spherical potentials (V) acting on both channels of the spin of that cation site. The NSCRF and SCRF were found by fitting linearly N_d vs. V, see Figure SI1 110 (Supporting Information). The optimal $U_{\rm eff}$ value was obtained by calculating the weighted 111 arithmetic mean of the difference between the reciprocals of the gradients of the linear 112 fittings. 113

114 **2.2 Surface and Molecular Models**

The (001) surface was investigated using the periodic $p(1 \times 1)$ supercell, which was created 115 from the bulk Fe₃S₄ using the METADISE code.⁹⁸ A vacuum of 18 Å was added in the direction 116 perpendicular to the surface to prevent interaction with the periodically repeated images and 117 to ensure that the surface can accommodate, in any conformation, the $H_2C_2O_4$ molecule, 118 which has a length of 4.7 Å. We considered four formula unit (f.u.) layers for the simulation 119 120 of the (001) surface slab, with an area of 102.517 Å². The atoms in the two topmost layers 121 were allowed to relax during geometry optimisations, to mimic the surface, while those atoms in the remaining two bottom layers were kept at their optimised bulk positions. The Brillouin 122 zone of the surface was sampled using a Γ -centred Monkhorst-Pack (MP) mesh of 5 × 5 × 1 k-123 points.⁹⁹ For the geometry optimisations, the partial occupancies were determined using the 124 finite temperature Gaussian smearing method⁸⁷ with 0.01 eV for the width of the smearing, 125 which was tested to ensure that the electronic entropy (TS) vanished. The free energy (F), 126 127 which is the variational quantity in the finite temperature Gaussian smearing approach differs from $E(\sigma)$ by $TS.^{87,100}$ However, we employed the tetrahedron method with Blöchl 128 corrections¹⁰¹ for the calculation of accurate energies as well as the electronic and magnetic 129 properties in static simulations of the optimised structures. Convergence of energy 130 differences within 1 meV per atom was further tested for the simulation supercell using 131 different vacuum thicknesses, cut off energy values as well as various numbers of total and 132 relaxed atomic layers. Dipole corrections were also included perpendicular to the surface 133 plane to compensate for any dipole created by the chemical species added onto the relaxed 134 side of the slab and to enhance the electronic convergence.^{102,103} 135

The isolated CO₂ and H₂O molecules were modelled in a periodic box of $14 \times 15 \times 16$ Å³ to ensure negligible interaction with their images in the neighbouring cells. The Gaussian scheme was used to determine the electronic partial occupancies during geometry optimisations and energy calculations, sampling only the Γ point of the Brillouin zone.

140 The effective Bader atomic charges were obtained using an improved grid of charge density 141 values without lattice bias,^{104–106} whereas the magnetic moments were integrated within the 142 same atomic basins. The work function (Φ), which is the minimum thermodynamic work 143 required for an electron to leave the Fe₃S₄(001) surface, was calculated as $\Phi = E_{vac} - E_{F}$, where

5

 E_{vac} is the potential at the vacuum level and E_F is the Fermi level of the slab. The STM images 144 were calculated using the Tersoff-Hamann method¹⁰⁷ in the implementation of Vanpoucke 145 and Brocks.¹⁰⁸ The wavenumbers of the fundamental harmonic vibrational modes were 146 147 determined using the central finite differences approach, by allowing each atom to move by a small displacement in the three Cartesian directions. The atomic displacements are used to 148 calculate the elements of the Hessian matrix, which are computed by numerical 149 150 differentiation of the analytical gradients of the energy with respect to the energy. The saddle points and minimum energy pathways (MEP) between initial and final states were calculated 151 using the climbing image nudged elastic band (cNEB) method.^{109,110} The cNEB algorithm used 152 153 for modelling the MEPs has been employed successfully before to obtain the saddle point structures and energies of several processes in heterogeneous catalysis.^{59,61,68,78,111} We have 154 155 used five images to model the MEP, which were optimised globally by means of the limitedmemory Broyden-Fletcher-Goldfarb-Shannon (LBFGS) method.^{89,112} The saddle points were 156 157 characterised by a single imaginary frequency along the reaction coordinate, which were 158 determined using the central finite differences method.

159 2.3 Surface Energy Diagrams

160 The surface energies for the slab before (γ_u) and after relaxation (γ_r) were evaluated as,

161
$$\gamma_{\rm u} = \frac{E_{\rm u} - n_{\rm b} E_{\rm b}}{2A} \tag{1}$$

162
$$\gamma_{\rm r} = \frac{E_{\rm r} - n_{\rm b} E_{\rm b}}{A} - \gamma_{\rm u}$$
(2)

where $E_{\rm u}$, $E_{\rm r}$ and $E_{\rm b}$ are the energies of the unrelaxed slab, the half-relaxed slab and one Fe₃S₄ f.u. in the bulk, respectively, whereas $n_{\rm b}$ and A are the number of Fe₃S₄ f.u. in the supercell and the surface area, respectively. The degree of relaxation (*R*) was calculated as $R = 100(\gamma_{\rm u} - \gamma_{\rm r})/\gamma_{\rm u}$.

167 The temperature-corrected energy $[E_{M}(T, p^{\circ})]$ for the adsorbate molecule (*M*) was 168 calculated according to,

169
$$E_{M}(T, p^{\circ}) = E_{M} - TS(T, p^{\circ})$$
 (3)

where $E_{\rm M}$ is the calculated DFT energy for the isolated adsorbate, T is the temperature and $S(T, p^{\circ})$ is the experimental entropy in the standard state, extracted from thermodynamic tables.¹¹³ The average adsorption energy for the adsorbates $[E_{\rm ads}(T, p^{\circ})]$ was obtained via,

173
$$E_{ads}(T, p^{\circ}) = \frac{E_{c} - (E_{r} + \sum n_{M} E_{M}(T, p^{\circ}))}{\sum n_{M}}$$
(4)

where E_c is the energy of the slab with the adsorbed species and n_M is the number of surfacebound adsorbates.

The process of partial oxidation of the Fe₃S₄(001) surface can be represented by the following
 chemical reaction,

178 $\operatorname{Fe}_{24}S_{32}(001) + n\operatorname{H}_2O(g) = \operatorname{Fe}_{24}S_{32-n}O_n(001) + n\operatorname{H}_2S(g)$ (5)

179 where *n* is the number of O atoms oxidising the surface.

For the partially oxidised surfaces, the effect of temperature on the surface free energy $[\sigma(T,p)]$ was introduced as follows,^{85,114–116}

182
$$\sigma(T,p) = \gamma_{\rm r} + C \left(E_{\rm ads}(T,p^{\circ}) - RT \ln \frac{p_{\rm H_2O}}{p_{\rm H_2S}} \right)$$
(6)

where *C* is the coverage of oxygen and *R* is the ideal gas constant. The last term represents the change in free energy of the H₂O and H₂S gases (assuming ideal gas behaviour) in equilibrium with the surface at constant temperature when the ratio of their partial pressures is allowed to change from an arbitrarily small to an arbitrarily large value.

The activation free energy (F_{ai}) was calculated as the difference between the free energy of 187 the saddle point and the free energy of the reactants, where the index *i* refers to each of the 188 three elemental steps that we calculated. The free energy of the elemental step (ΔF_i) was 189 obtained as the difference of the free energy of the products and the free energy of the 190 191 reactants. The energy diagrams were prepared using the most stable binding modes of the 192 reactants, intermediates and final products, which were connected by saddle points and MEPs. The Helmholtz free energy (ΔF) was calculated as $\Delta F = \Delta E - T \Delta S_{vib}$ where S_{vib} is the 193 vibrational entropy, which was obtained as follows for the adsorbed species, 194

195
$$S_{\text{vib}} = R\left(\sum_{i} \frac{\hbar\omega_i/k_{\text{B}}T}{exp(\hbar\omega_i/k_{\text{B}}T)-1} - \sum_{i} \ln(1 - exp(-\hbar\omega_i/k_{\text{B}}T))\right)$$
(7)

where \hbar is the reduced Planck constant, ω_i are the vibrational frequencies and $k_{\rm B}$ is the Boltzmann constant. We have used the entropies reported in thermodynamic tables for the isolated CO₂, H₂O, H₂C₂O₄ and O₂, to avoid calculating the significant translational and rotational components of the entropy for these gas phase molecules.¹¹⁷

200 **3. Results and Discussion**

3.1. Bulk Phase of Greigite (Fe₃S₄)

We first investigated the optimised bulk structure of Fe₃S₄, which is later used to create the 202 (001) surface for the calculation of the phase diagrams, molecular adsorptions and reaction 203 mechanisms. Our starting point for Fe₃S₄ is the bulk structure characterised by Li and 204 collaborators using room temperature powder X-ray diffraction (XRD).⁵⁷ Fe₃S₄ has the typical 205 206 face-centred cubic (fcc) lattice of the spinel structure and crystallises in the space group $Fd\overline{3}m$ (no. 227). Figure 1 depicts the conventional cubic unit cell of Fe₃S₄ containing eight 207 formula units (f.u.) and the periodically repeated images of those ions located at the edges 208 and corners of the cell. The sulfide anions are formally in the 2- oxidation state and exhibit a 209 nearly regular cubic close-packed (ccp) distribution along the [111] direction. The S atoms 210 occupy the tetrahedrally distorted Wyckoff 32e positions with coordinates (u, u, u) and are 211 surrounded by 12 other equidistant nearest-neighbour anions. The difference of the sulfur 212 213 parameter u from $\frac{1}{4}$, in direct coordinates, represents the shift of the anions in the [111] direction to accommodate cations of different coordination number and radii in the 214 215 interstitial sites. Fe₃S₄ is an inverse 2–3 type spinel, as it contains two 3+ and one 2+ cations per formula unit. The tetrahedral Fe_A atoms are in the highest 3+ oxidation state and occupy 216 217 the 8*a* crystallographic positions at (1/8, 1/8, 1/8). The octahedral Fe_B cations, which have two equal populations in the 2+ and 3+ valence states each, fill the 16d octahedral holes with 218 coordinates (1/2, 1/2, 1/2). However, only 1/8 of the total number of tetrahedral positions 219 220 and half of the total number of octahedral holes are filled by the cations. The cations form 221 two different types of alternate layers in the [111] direction, one composed only by Fe_B ions and the other by both Fe_A and Fe_B . 222



223

Figure 1. Polyhedral model of the face-centred cubic (*fcc*) conventional unit cell containing 8
formula units (f.u.) of Fe₃S₄. Crystallographic directions are indicated. Fe_A atoms are in dark
yellow, Fe_B atoms are in magenta and S atoms are in light yellow.

Table 1 summarises the calculated and previously reported structural, electronic and 227 magnetic properties for the Fe₃S₄ bulk material. Our calculations reveal that the lattice 228 parameter for the cubic unit cell of Fe₃S₄ was overestimated by 0.25 Å with respect to 229 experiment,⁵⁷ but is in excellent agreement with the value reported by Devey et al. when 230 using PW91 + U where U_{eff} = 5 eV.¹¹⁸ The atomic positions were allowed to relax fully, but the 231 232 cations remained in their ideal crystallographic sites. However, the value obtained for the sulfur *u* parameter is just slightly larger than in experiment, indicating that the anions suffer 233 a relatively larger displacement in the simulation cell. The anion-cation bond distances, which 234 are mathematically related to the sulfur parameter by $d(Fe_A - S) = \sqrt{3}a(u - 1/8)$ and 235 $d(\text{Fe}_B - \text{S}) = a(1/2 - u)$, were overestimated by 0.08 and 0.05 Å, respectively, compared 236 to a previous study¹¹⁹ but in agreement with a larger lattice parameter. Our computational 237 setup provides better atomic Bader charges (q) and magnetic moments (m_s) than calculations 238 using small values of U_{eff} ,^{118,120} and our results are in better agreement for comparable 239 Hubbard parameters to the one used in this study.¹¹⁸ Although the atomic charges of 1.371 240 e⁻ for Fe_A, 1.189 e⁻ for Fe_B and -0.937 e⁻ for S are still underestimated with regards to their 241 242 oxidation states, this is a common finding with Bader charges, and they do provide evidence that our Fe₃S₄ is indeed an inverse spinel as $q(Fe_A) > q(Fe_B)$, which has been observed 243 before.^{57,121} The larger magnetic moment of $-4.177 \ \mu_B$ atom⁻¹ for Fe_A refers to the high-spin 244

electronic distribution $e_{\downarrow}^2 t_{2\downarrow}^3 e_{\uparrow}^x$, whereas we can assign the configuration $t_{2q\uparrow}^3 e_{\uparrow}^2 t_{2q\downarrow}^{1-x}$ to the 245 antiparallel aligned Fe_B, implying that the conduction mechanism involves both cation 246 sublattices. Our total spin magnetisation of saturation (M_s) of 2.00 μ_B f.u.⁻¹ is in excellent 247 agreement with the early works of Morrish and collaborators using Mössbauer 248 spectroscopy.^{121,122} Note that a wide range of different values have been reported for Fe₃S₄, 249 corresponding to other electronic and magnetic properties.^{119,123–127} Figure 2 displays the 250 251 electronic density of states (DOS) for the bulk phase of Fe_3S_4 . The occupied *e* and t_2 states of the Fe_A ions appear strongly hybridised as a narrow single peak in the minority channel of 252 spins at around -8.5 eV. The Fe_A cations also have a small *e* band crossing the Fermi level in 253 the α channel, illustrative of the itinerant electron provided by the t_{2q} valence band of the Fe_B 254 sublattice in the minority channel of the spins. The t_{2g} and e_g valence bands of the Fe_B cations 255 are located between -7.0 and -3.0 eV in the majority channel of the spins. The virtual *e* and 256 t_2 states of Fe_A and the t_{2q} and e_q conduction bands of Fe_B appear above 1.5 eV. Importantly, 257 258 the hybridisation of the S p levels with the t_{2g} states of the Fe_B cations observed in the minority spin channel of the DOS of the bulk of Fe_3S_4 explains the small magnetic moment of -0.263259 $\mu_{\rm B}$ calculated for the anion. 260

Table 1. Unit cell lattice parameter (*a*), sulfur parameter (*u*), interatomic distances (*d*), atomic charges (*q*), atomic magnetic moments (m_s) and total spin magnetisation (M_s) for the thiospinel greigite Fe₃S₄ from this work and previous reports. Negative atomic magnetic moments indicate antiparallel alignment.

Property	This work	Previous reports	Reference
a (Å)	10.1251	9.8719	57
u	0.2586	0.2546	57
<i>d</i> (Fe _A −S) (Å)	2.29	2.21	119
<i>d</i> (Fe _B −S) (Å)	2.47	2.42	119
$q(Fe_A)$ (e ⁻ atom ⁻¹)	1.371	1.1	120
q(Fe _B) (e⁻ atom⁻¹)	1.189	1.0	120
<i>q</i> (S) (e ⁻ atom ⁻¹)	-0.937	-0.8	120
m_s (Fe _A) (μ_B atom ⁻¹)	-4.177	-2.8	120
$m_s(Fe_B)$ (μ_B atom ⁻¹)	3.614	3.0	120
$m_s(S)$ (μ_B atom ⁻¹)	-0.263	0.1	120
<i>M</i> _S (Fe ₃ S ₄) (μ _B f.u. ^{−1})	2.000	2.00	121



265

Figure 2. Atomic projections of the spin decomposed total density of states (PDOS) for the bulk phase of Fe₃S₄. α and β stand for the majority and minority channel of the spins, respectively. Fe_A and Fe_B contributions are from the 3*d* bands, whereas S contributions are from the 3*p* orbitals. Fe_A projections are in black, Fe_B projections are in red and S projections are in green.

271 **3.2. Pristine Fe₃S₄(001) Surface**

We have also simulated the two non-polar stoichiometric and symmetric terminations of the 272 pristine Fe₃S₄(001) surface, where we have kept the cell borders fixed at the parameters 273 calculated for the bulk, represented in Figure 3. The stacking of the atomic planes is 274 275 $(Fe_A)-(Fe_{B4}-S_8)-(Fe_A)$ and $(Fe_{B2}-S_4)-(Fe_{A2})-(Fe_{B2}-S_4)$ for the terminations Fe_A and Fe_B-S_A respectively, where the atoms within parenthesis are located approximately within the same 276 277 layers. We have employed single surface unit cells to simulate terminations Fe_A and Fe_B -S of the pristine $Fe_3S_4(001)$ surface, which are the smallest periodically repeating surface unit cells 278 that we can use to model the stacking sequences of these systems. Both terminations Fe_A and 279 Fe_B-S are reconstructed Tasker type 3 surfaces,¹²⁸ where the dipole moment was removed 280 before geometry optimisation by displacing half of the ions from the exposed stoichiometric 281 stacking sequence at the top of the slab to the bottom of the slab. Termination Fe_A has a bulk-282 like structure comprising a full monolayer (ML) of S atoms, where 25% are 4-fold and the 283 284 remaining 75% have one dangling bond. The topmost plane of termination Fe_A also contains 285 0.5 ML of 5-fold Fe_B cations, which form alternating rows in the [110] direction, on top of which are 0.125 ML of 2-fold Fe_A ions with the $(\sqrt{2} \times \sqrt{2})R45^\circ$ symmetry, according to the 286 vectorial notation of Wood.¹²⁹ Termination Fe_B-S presents 0.5 ML of 2-fold S atoms, 287 unsurprisingly also forming alternating rows in the [110] direction, alongside 0.25 ML of Fe_B 288 289 with 3 dangling bonds, whereas the layer beneath contains 0.25 ML of 2-fold Fe_A ions.



290

Figure 3. Side (top panels) and top (bottom panels) views of terminations (a) Fe_A and (b) Fe_B -S of the $Fe_3S_4(001)$ surface after geometry optimisation. Layers containing atoms with dangling bonds are highlighted. Crystallographic directions and stacking sequence of the atomic layers before geometry optimisation are indicated. Fe_A atoms are in dark yellow, Fe_B atoms are in magenta and S atoms are in light yellow.

Figure 4 shows the relaxation of the interplanar distances, which was calculated as Δ_{ii} = 296 $100 (d_{ij} - d_{ij}^{\circ})/d_{ij}^{\circ}$, where d_{ij} is the separation between the relaxed *i* and j = i + 1 layers 297 and d_{ij}° is the separation for the equivalent planes in the bulk.¹³⁰ Following the geometry 298 299 optimisation of both terminations, we found a relaxation behaviour that was increasingly 300 attenuated towards the bulk. Each pair of layers Fe_A -i and Fe_B -S-j that were allowed to relax showed a preference to move together in alternating directions. For termination FeA, the 301 protruding twofold Fe_A-1 ions suffered a strong inward relaxation of 141%, recovering their 302 303 bulk tetrahedral coordination, when they merged with the subsurface layer Fe_A -3 where 0.25 ML of fourfold Fe_A are located, see Figures 3 (a) and 4. The negatively charged subsurface 304 plane Fe_B-S-2 experienced only a moderate 13% shift towards the bulk, as it was pulled 305 306 electrostatically by the merged layer beneath containing a larger number of Fe_A ions and 307 therefore more positive charge carriers than in the bulk. The outward displacement of 16% of the subsurface layer Fe_A-3 can also be rationalised in terms of the Coulomb attraction 308 exerted by the plane Fe_B-S-2 above, which lies closer as a result of its inward relaxation. The 309 310 subsurface layer Fe_B–S-4 shifted by 9% towards the surface, to compensate for the expansion of the plane Fe_A-3 above, whereas Fe_A-5 preferred to reduce its interplanar distance to the 311

closer Fe_B–S-6 layer that was not allowed to relax. The relaxation pattern of termination Fe_A for the Fe₃S₄(001) surface is in excellent agreement with the relaxation reported for the same termination of the magnetite Fe₃O₄(001) surface.¹¹⁶ Moreover, the contractions and expansions of the interplanar distances Fe_B–S-2 and Fe_A-3 are considerably larger for termination Fe_B–S that than for termination Fe_A, see Figure 4. However, the inward shift of the topmost layer Fe_B–S-2 of termination Fe_B–S is only half of the contraction of the exposed layer Fe_A-1 of termination Fe_A.



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Figure 4. Relaxation (Δ_{ij}) of the interplanar distance for the terminations (a) Fe_A and (b) Fe_B–S of the Fe₃S₄(001) surface in black and red, respectively.

From our calculations of the surface energies before and after geometry optimisation, we 322 have found that termination Fe_A is the most stable of the $Fe_3S_4(001)$ planes, see Table 2. The 323 largest relaxation of R = 50.68% obtained for termination Fe_A is a testament of the strong 324 325 inward displacement of the protruding Fe_A ion to the subsurface layer. We have calculated 326 the atomic Bader charges, which are lower for the exposed Fe_A and S ions than for their bulk 327 counterparts. The (001) surface cations, Fe_A and Fe_B , are more ionic at termination Fe_A than 328 at termination Fe_B -S, whereas we observed the opposite trend for the S anion. We also found an increment by 0.068 e^- atom⁻¹ in the positive charge of the Fe_B ions in termination Fe_A 329 compared to their bulk values. The magnetic moments calculated for all the under-330 331 coordinated atoms are larger in termination Fe_A than in termination Fe_B -S. Our calculations

332 only predict larger magnetic moments than in the bulk for the Fe_B ions with dangling bonds in both terminations, Fe_A and Fe_B -S, and for the exposed S atoms in the termination Fe_A . In 333 contrast to the bulk, we found that the charges and magnetic moments are larger for the 334 335 exposed Fe_B cations than for the Fe_A ions for the two terminations considered in this study. 336 We also noted that the small magnetic moments of the S atoms, which are aligned parallel to 337 the Fe_A ions in the bulk, prefer to align parallel to the Fe_B cations in both terminations of the $Fe_3S_4(001)$ surface. The work function indicates that termination Fe_A is marginally less reactive, 338 by 0.055 eV, than termination Fe_B -S, which can provide more easily the loosest held electron 339 340 required for the catalytic formation of $H_2C_2O_4$. From here, we do not consider the Fe_B-S 341 termination for further analysis due to its large relaxed surface energy, which makes it 342 unlikely to be expressed in the crystal morphology of Fe₃S₄.

Table 2. Surface energies before (γ_u) and after relaxation (γ_r) , percentage of relaxation (R), average atomic charges (q), average magnetic moments (m_s) and work function (Φ) for terminations Fe_A and Fe_B–S of the pristine Fe₃S₄(001) surface. Negative atomic magnetic moments indicate antiparallel alignment.

Termination	Fe _A	Fe _B –S
γ _u (meV Å ⁻²)	66.3	118.7
γ _r (meV Å ⁻²)	32.7	75.2
R (%)	50.68	36.69
$q(Fe_A)$ (e ⁻ atom ⁻¹)	1.187	1.004
$q(Fe_B)$ (e ⁻ atom ⁻¹)	1.257	1.022
<i>q</i> (S) (e ⁻ atom ⁻¹)	-0.845	-0.919
$m_s(Fe_A)$ (μ_B atom ⁻¹)	-3.858	-3.679
$m_s(Fe_B)$ (μ_B atom ⁻¹)	4.040	3.713
$m_s(S)$ (μ_B atom ⁻¹)	0.318	0.139
Φ (eV)	5.613	5.558

347 **3.3. Partially Oxidised Fe₃S₄(001) Surface**

We next analysed the relative stabilities of termination Fe_A of the $Fe_3S_4(001)$ surface containing different coverages of oxygen as a function of the ratio of the partial pressures of H₂O and H₂S and temperature. We have successfully used these ab initio thermodynamic techniques previously to model the oxidation¹¹⁵ and hydrogenation⁷⁸ of cobalt, the hydration of platinum⁸⁵ and the redox behaviour of Fe_3O_4 .¹¹⁶ We have calculated several coverages, by replacing in turn each of the S atoms with dangling bonds by O atoms. There is strong

experimental evidence, backed up by calculations, that different iron and nickel sulfide phases, 354 including Fe₃S₄,^{61,62} troilite (hexagonal FeS),¹³¹ pyrrhotite (hexagonal Fe_{1-x}S), ^{68,132} polydymite 355 (Ni₃S₄)⁶⁸ and violarite (FeNi₂S₄)⁶⁸ oxidise partially and easily when they are in contact with 356 357 water, forming catalytically active core-shell sulfide-oxide nanoparticles. The dimension of our computational slab allows the substitution of up to eight S atoms from termination Fe_A of 358 the Fe₃S₄(001) surface. We have considered several distributions of the S and O atoms for 359 each O coverage and used equation 6 to calculate their surface free energies. The atomic 360 configurations with the lowest surface free energies for each ratio of the two anions were 361 362 selected to build the surface phase diagram. The surface free energies for each O coverage 363 are represented by non-planar potential energy surfaces that intersect each other at specific 364 temperatures and ratios of partial pressures of H₂O and H₂S, as shown in a bird's eye view in Figure SI2. The curves formed where two potential energy surfaces cross correspond to the 365 366 conditions required to modify the extent of the partial oxidation of the $Fe_3S_4(001)$ surface. 367 However, the most convenient way to depict and analyse this type of diagram is by making a 368 bi-dimensional projection along the surface free energy axis onto the plane formed by the temperature and ratio of partial pressures of H₂O and H₂S, see Figure 5 (a). The surface phase 369 370 diagram shows that termination Fe_A of the Fe₃S₄(001) surface requires temperatures above 230 K and a ratio of partial pressures of H₂O and H₂S below 100 to remain pristine. The lowest 371 possible oxygen coverage of C = 0.98 O nm⁻² is achieved for a maximum temperature of 300 372 K and a minimum ratio of partial pressures of H₂O and H₂S of 10^{2.2}. The coverages 2.93 and 373 3.90 O nm⁻², with very limited ranges of stabilities and equivalent to three and four O atoms, 374 respectively, appear at higher temperatures and higher ratio of partial pressures of H₂O and 375 H_2S than the 0.98 O nm⁻² coverage. The surface is capable of accommodating five O atoms, 376 *i.e.* a coverage of 4.88 O nm^{-2} if the temperature is increased up to 530 K for a ratio of partial 377 pressures of H₂O and H₂S smaller than 10^{5.5}. Figure 5 (b) displays this coverage of O atoms, 378 which is consistent with a 62.5% partial oxidation, and is in good agreement with our previous 379 work on FeNi₂S₄, where we found that 75% of the surface S could be replaced by O.⁶⁸ Our 380 results for the Fe₃S₄(001) surface suggest that a coverage of C = 4.88 O nm⁻² has a relatively 381 wide range of stabilities between 300 and 530 K when H₂O is the predominant gas. For 382 comparison, FeNi₂S₄ suffers spontaneous surface oxidation when it is calcined at 473 K in air 383 or moisture.⁶⁸ The coverage is expanded to 6.83 O nm⁻², or seven O atoms, if the ratio of 384 partial pressures of H₂O and H₂S is increased by a further $10^{0.5}$ with respect to C = 4.88 O nm⁻² 385

and the temperature does not exceed 600 K. The full coverage of C = 7.80 O nm⁻² only becomes thermodynamically favourable for a temperature and ratio of partial pressures of H₂O and H₂S larger than the values required for the incorporation of seven O atoms in our surface. Coverages C = 1.95 and 5.85 O nm⁻², corresponding to two and six O atoms, respectively, are not accessible, as they are the least energetically stable partially oxidised Fe₃S₄(001) surface slab models within the conditions of pressure and temperature of our phase diagram.





Figure 5. (a) Phase diagram for the partially oxidised termination Fe_A of the $Fe_3S_4(001)$ surface as a function of the ratio of the partial pressures of H_2O and H_2S and temperature. The O coverages (*C*) are indicated for each region. (b) Top view of the $Fe_3S_4(001)$ surface with an oxygen coverage of C = 4.88 O nm⁻². Layers containing atoms with dangling bonds are highlighted. Crystallographic directions and the adsorption sites Fe_A (\blacksquare), Fe_B (\blacktriangle), O (×) and S (\bullet) are indicated. Fe_A atoms are in dark yellow, Fe_B atoms are in magenta, S atoms are in light yellow and O atoms are in red.

The STM images of both the pristine termination Fe_A of the $Fe_3S_4(001)$ surface and with an oxygen coverage of C = 4.88 O nm⁻² are represented in Figure 6. The negative sample bias of -2.5 eV used to generate these STM images indicates that the electrons hop from the valence band to the probe tip, as expected for the half-metal properties predicted in the DOS for the bulk of Fe₃S₄. The S atoms appear as the brightest spots in the pristine and oxidised surfaces, whereas the cations are not well resolved. The O atoms can be identified in the oxidised surface, but they are less bright and smaller than the S anions, in agreement with their smaller ionic radius. The pristine surface displays a pattern of well-ordered arrays of S atoms along the $[1\overline{10}]$ direction, which is lost after oxidation.





Figure 6. Scanning tunnelling microscopy (STM) images of termination Fe_A of the Fe₃S₄(001) surface, where (a) represents the pristine surface and (b) the surface with an oxygen coverage of $C = 4.88 \text{ O nm}^{-2}$. The images were generated using a bias of V = -2.5 eV, a tip distance of d= 2.00 Å, and a density of (a) $\rho = 0.0101 \text{ e} \text{ Å}^{-3}$ and (b) $\rho = 0.0120 \text{ e} \text{ Å}^{-3}$. Crystallographic directions are indicated. Fe_A atoms are in dark yellow, Fe_B atoms are in magenta, S atoms are in light yellow and O atoms are in red.

417 To provide additional insight into the Fe₃S₄(001) surface, we have also analysed its 418 thermodynamic stability, electronic and magnetic properties, as well as the surface reactivity 419 as a function of the oxygen coverage. The surface free energy (σ) at 0 K, which quantifies the 420 work needed to create the surface of a material and indicates its stability, increases gradually

with the oxygen coverage, indicating that partially oxidising the Fe₃S₄(001) facet is an 421 422 energetically unfavourable process, see Figure 7 (a). Our calculations suggest that the surface free energy increases relatively rapidly for coverages from 0.00 O $nm^{-2} < C < 1.95$ O nm^{-2} and 423 especially for 4.88 O nm⁻² < C < 5.85 O nm⁻² and 6.83 O nm⁻² < C < 7.80 O nm⁻². The surface 424 free energy barely changes for the coverages 2.93 and 3.90 O nm⁻², explaining the narrow 425 range of conditions which need to be controlled precisely to develop these particular partial 426 degrees of oxidation. Interestingly, the partially oxidised surface with a coverage of 6.83 O 427 nm^{-2} is approximately 13 meV Å⁻² more stable than the facet with C = 5.85 O nm^{-2} . The atomic 428 charges and magnetic moments do not change noticeably and remain fairly constant for any 429 oxygen coverage, with the exception of the smaller negative charge of the S atom for C = 6.83430 O nm⁻² as shown in Figure 7 (b) and (c). The work function shows two minima at the coverages 431 C = 1.95 and 5.85 O nm⁻², suggesting that these partial degrees of oxidation make the 432 Fe₃S₄(001) surface considerably more reactive, see Figure 7 (d). 433





Figure 7. (a) Surface free energies (σ) at 0 K, (b) average atomic charges (q), (c) average magnetic moments (m_s) and (d) work function (Φ) for different oxygen coverages of the Fe₃S₄(001) surface. Negative atomic magnetic moments indicate antiparallel alignment.

438 **3.4. Molecular Adsorptions on the Partially Oxidised Fe₃S₄(001) Surface**

We have investigated the interaction of the single molecules CO_2 and H_2O on the $Fe_3S_4(001)$ 439 surface with a coverage 4.88 O nm^{-2} , which from hereon is the surface model of our catalyst. 440 In order to simplify the notation used in this paper, we will refer to the Fe₃S₄(001) surface 441 442 with a 62.5% degree of partial oxidation as "62.50-Fe₃S₄(001)." We have assessed the effect 443 of these molecular adsorptions on the surface free energy, charge transfer, atomic magnetic moment and work function of the 62.50-Fe₃S₄(001) surface. We also discuss the geometries 444 of the most favourable adsorption modes and the shifts of the fundamental scaled vibrational 445 modes of the adsorbates. Figure 5 (b) shows the four adsorption sites investigated in this 446 study, including atop the (i) Fe_A , (ii) Fe_B , (iii) S and (iv) O atoms. 447

Table 3 lists the calculated bond distances (*d*), bond angles (\angle) as well as the harmonic (unscaled) and scaled asymmetric stretching (v_{asym}), symmetric stretching (v_{sym}) and bending (δ) fundamental vibrational modes for the CO₂ and H₂O adsorbates, which we have used to benchmark our computational SCAN-D2 setup. Our calculations indicate that we have

obtained, up to the second decimal, the correct equilibrium bond distances for the CO₂ and 452 H₂O molecules compared to the experimental values.^{133,134} The bond angle of 180° predicted 453 for the CO2 molecule is in perfect agreement with the value inferred from electronic 454 spectroscopy,¹³³ whereas the bond angle calculated for the H₂O molecule is only slightly 455 overestimated with respect to experiment.¹³⁴ Despite their different point groups, *i.e.* C_{2v} for 456 H₂O and $D_{\infty h}$ for CO₂, both molecules are characterised by only three fundamental vibrational 457 modes. Compared to previous reports, our calculated asymmetric stretching, symmetric 458 stretching and bending harmonic vibrational modes are red-shifted for CO₂¹³³ and H₂O^{135,136}, 459 460 although the computational description is better for the former adsorbate than for the latter. 461 The smallest deviation from the experimental value was calculated for the bending mode of 462 CO₂ and the largest difference was calculated for the asymmetric stretching mode of H₂O, which were overestimated by 2 and 167 cm⁻¹, respectively. The discrepancies with 463 464 experiment are larger for H₂O than for CO₂ due to the larger anharmonicity of the vibrations 465 for the hydrogen-containing bonds with respect to the C=O bonds. Our harmonic vibrational 466 frequencies are calculated as the second derivative of the potential energy with respect to the atomic positions in the vicinity of the minimum of the well. This section of the potential 467 energy surface shows quadratic behaviour, leading to harmonic vibrational energy levels that 468 469 are equally separated. However, experimental fundamental modes represent the transition 470 between the ground state and the first state vibrational energy levels of an anharmonic potential energy surface. We have also determined the empirical scaling factors needed to 471 472 convert our harmonic frequencies calculated for the isolated molecules into anharmonic vibrational modes, which can be compared directly with experiment.^{137–141} The scaling factor 473 (c) was estimated as 474

475

$$c = \frac{\sum \omega_{\exp} \omega_{\text{calc}}}{\sum \omega_{\text{calc}}^2}$$
(8)

476 where ω_{exp} are the experimental wavenumbers and ω_{calc} are the calculated harmonic 477 vibrational wavenumbers. The uncertainty (*u*) of the scaling factors is calculated as

478
$$u = \frac{\sum \left[\omega_{calc}^2 (c - \omega_{exp}/\omega_{calc})^2\right]}{\sum \omega_{calc}^2}$$
(9)

We obtained the values of $c = 0.9655 \pm 0.0001$ for CO₂ and 0.96089 \pm 0.00004 for H₂O, which supports the excellent performance of the meta-GGA SCAN functional for the calculation of 481 vibrational frequencies. After scaling the wavenumbers to account for anharmonicity effects, we found that the asymmetric stretching for both molecules and the symmetric stretching of 482 483 H₂O were overestimated, whereas the remaining vibrational frequencies were 484 underestimated with respect to the experimental values. Unsurprisingly, the largest difference of 35 cm⁻¹ between scaled and experimental wavenumbers was obtained for the 485 bending mode of H₂O and the smallest shift of only 1 cm⁻¹ was calculated for the symmetric 486 487 stretching mode of H₂O. In sections 3.4.1 and 3.4.2 only the scaled vibrational numbers are discussed to facilitate comparison with future experiments. 488

Table 3. Calculated and previously reported intramolecular bond distances [d(C=O) and d(H–O)], bond angles [\angle (O=C=O) and \angle (H–O–H)] and wavenumbers for the fundamental vibrational modes for the isolated CO₂ and H₂O molecules, respectively. The presented harmonic vibrational modes, both unscaled and scaled, are the asymmetric stretching (v_{asym}), symmetric stretching (v_{sym}) and bending (δ) modes for the O=C=O and H–O–H groups in the CO₂ and H₂O molecules, respectively.

		CO ₂	H ₂ O
d (Å)	Calculated	1.165	0.964
	Experimental	1.162 (Ref ¹³³)	0.958 (Ref ¹³⁴)
∠ (°)	Calculated	180.00	105.09
	Experimental	180.00 (Ref ¹³³)	104.48 (Ref ¹³⁴)
v _{asym} (cm⁻¹)	Unscaled	2447	3923
	Scaled	2363	3770
	Experimental	2349 (Ref ¹³³)	3756 (Ref ¹³⁵)
ν _{sym} (cm⁻¹)	Unscaled	1366	3807
	Scaled	1319	3658
	Experimental	1333 (Ref ¹³³)	3657 (Ref ¹³⁶)
δ (cm⁻¹)	Unscaled	669	1623
	Scaled	646	1560
	Experimental	667 (Ref ¹³³)	1595 (Ref ¹³⁶)

495 3.4.1. CO₂ Interaction with the Partially Oxidised Fe₃S₄(001) Surface

We have probed the adsorption of a single CO₂ molecule in a pre-activated configuration, where we reduced its point group from $D_{\infty h}$ to $C_{2\nu}$, by bending the apex angle to \angle (O=C=O) = 130°.^{58,59,76,111,142–146} To characterise the adsorption configurations of our CO₂ molecule, we have borrowed the notation widely used in organometallics to describe the bonding and structural types of metal–CO₂ complexes, which is based on the hapticity of the ligand (μ_n)

and the number of surface atoms (m) it can bridge (η^m) .^{147–149} We approached the bent CO₂ 501 molecule in the $\mu_3 - \eta^3$ configuration, with both the C_2 rotational axis and the vertical mirror 502 503 plane σ_v perpendicular to the surface, *i.e.* with the C atom coordinating the exposed O or S 504 atoms of the catalyst and the molecular O atoms bonding the Fe cations with dangling bonds. We have also considered the initial η^1 –O adsorption modes, where the symmetry elements 505 506 C_2 and σ_v of the adsorbate lie parallel and perpendicular, respectively, to the surface and the molecular O atom is interacting with the under-coordinated Fe_A or Fe_B ions. We placed the 507 CO₂ molecule at 1.8 Å from the partially oxidised surface of the thiospinel before allowing the 508 509 atomic positions of the interface to relax.

510 Our calculations indicate that the CO_2 molecule energetically prefers to interact molecularly with the O ion of the 62.5O-Fe₃S₄(001) surface in the $\mu_3 - \eta^3$ configuration, where it releases 511 0.746 eV, see Table 4 and Figure 8 (a). We also found evidence that the CO₂ molecule binds 512 exothermically to the Fe_A ion in the end-on η^1 –O configuration, with an adsorption energy 513 $E_{ads} = -0.188$ eV, which is less favourable than at the O site. Despite starting in the $\mu_3 - \eta^3$ 514 configuration, the optimised structure of the adsorption of CO₂ on the S site is the side-on 515 $\mu_2 - \eta^2 - (C, O)$ mode, which is a slightly endothermic process requiring 0.016 eV. The $\eta^1 - O$ 516 517 interaction configuration with the exposed Fe_B atoms of the 62.5O-Fe₃S₄(001) surface displays the largest positive energy at E_{ads} = 0.589 eV. The trend of binding energies on the anion sites 518 can be rationalised in terms of the number of interactions that the CO₂ molecule forms with 519 the surface, *i.e.* three in the $\mu_3 - \eta^3$ configuration on the most exothermic O position, and two 520 in the $\mu_2 - \eta^2 - (C, O)$ mode on the S atom. However, the trend of the adsorption strength on 521 the cation sites depends on their total number of dangling bonds, *i.e.* two and one for the Fe_A 522 and Fe_B atoms, respectively. Quesne and collaborators reported that chemisorption of the 523 activated CO₂ molecule on the surfaces of transition metal carbides occurs either directly, 524 525 spontaneously and barrierless, or via a linearly physisorbed intermediate with a small energy barrier.¹⁵⁰ Both the chemisorption and physisorption processes of CO₂ on the surfaces of 526 527 transition metal carbides are exothermic, with the former releasing more energy than the latter.¹⁵⁰ Thus, regardless of the adsorption mechanism of CO₂ on the partially oxidised 62.5O-528 Fe₃S₄(001) surface, we expect this process to take place on the O site, given its relatively large 529 exothermic adsorption energy of -0.746 eV. 530

The interactions have a noticeable impact on the geometry of the interface, the vibrational
 properties of the CO₂ molecule, as well as on the electronic properties and thermodynamic

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stability of the partially oxidised surface of our catalyst. The largest interfacial O-Fe distances, 533 with the values d = 2.308 and 2.157 Å were calculated for the η^1 –O structures on the Fe_A and 534 Fe^B sites, respectively, indicating that the only coordination bond for these adsorption modes 535 536 is weak. The CO₂ molecule is more strongly attached to the anion sites, and particularly to the surface O atom, since the distances for both the interfacial secondary O-Fe and main C-S 537 538 (C–O) coordination bonds are the smallest overall. The intramolecular C=O bond distances, which experience minor elongations for the η^1 –O interactions with the two types of Fe cation 539 sites, are stretched by 0.090 and 0.112 Å for the $\mu_3 - \eta^3$ adsorption on the O and $\mu_2 - \eta^2 - (C, O)$ 540 541 configuration on the S positions, respectively. Despite introducing a pre-activated bent CO₂ 542 molecule, which remained in this geometry on the anion sites during geometry optimisation, 543 we observed that the adsorbed molecule became approximately linear on the cation positions, similar to the molecule in its isolated state. This has also been found in other iron sulfide and 544 545 oxide catalytic materials, where a pre-activated molecule is introduced but allowed to relax unconstrainedly during geometry optimisation.^{58,60,111,146} 546

547 The changes observed in the intramolecular C=O bond distances are directly proportional to the red-shift of the asymmetric stretching vibrational modes compared to the isolated 548 adsorbate, with the smallest value of 1520 cm⁻¹ obtained for the ground state $\mu_3 - \eta^3$ binding 549 configuration on the surface O site. Our results suggest that the variations in the symmetric 550 stretching and average bending vibrational modes cannot be rationalised in terms of the 551 reduction of the intramolecular C=O bond length. As expected, the symmetric stretching 552 553 vibrational modes for the CO₂ molecule adsorbed on the anion sites are smaller than on the cation positions, but the largest wavenumbers at each type of site were calculated for the 554 most stable adsorption modes on the O and Fe_A positions. The interaction of CO₂ with the 555 surface O and S atoms leads to a blue-shift of the average bending vibrational mode with 556 557 regard to the isolated adsorbate, whereas the η^1 –O coordination to the Fe sites causes a redshift. Table SI1 summarises both the unscaled and scaled vibrational frequencies calculated 558 559 for the adsorbed CO₂ molecule.

The charge transfers denote that the cation sites, which are deficient in electrons, receive a minor charge $\Delta q = 0.03 \text{ e}^-$ from the adsorbate. However, the electron-rich O and S positions donate -0.209 and -0.537 e^- , respectively, to the σ^* antibonding molecular orbital of the CO₂ molecule, which weakens the intramolecular C=O bond, forcing a bent configuration. The adsorption of the CO₂ molecule reduces the work function, which makes the 62.50-Fe₃S₄(001)

- surface of our catalyst more reactive, except for the $\mu_2 \eta^2 (C, O)$ interaction with the S site. The lowest work function values $\Phi = 4.662$ and 4.221 eV required to remove the loosest held electron were predicted when the adsorbate binds the Fe_A and Fe_B sites, respectively, as the surfaces have increased marginally their electron density after adsorption of CO₂. The exothermic interactions of a single CO₂ molecule enhance the thermodynamic stability of our $62.5O-Fe_3S_4(001)$ surface, with the lowest value of the surface free energy $\sigma = 65.087$ meV Å⁻² computed for the $\mu_3 - \eta^3$ adsorption on the O site.
- Previous studies have shown that the (001) and (111) surfaces of the thiospinels Fe₃S₄^{58,111} 572 and FeNi₂S₄⁷⁶ are unable to activate the CO₂ molecule, which only remains physisorbed. 573 574 Among the pure sulfides, the (111) surface of mackinawite (tetragonal FeS) is the only catalyst able to dissociate a chemically activated CO₂ molecule into stable fragments.^{111,146} The 575 576 Fe₃O₄(001) and (111) surfaces are also capable of strongly chemisorbing the adsorbate, but the dissociation reaction has large activation energies, while the CO and O fragments are at 577 least 1.5 eV less stable than the CO₂ molecule.¹¹¹ To date, the catalytic activity towards the 578 579 reduction of bicarbonate has been reported for the major surfaces of the partially oxidised sulfides Fe_3S_4 , 58,62 hexagonal $Fe_{1-x}S$, $^{68,132}Ni_3S_4$ and $FeNi_2S_4$, 68 with the latter described as the 580 581 most efficacious system due to the synergism between Fe and Ni and the O content. Our DFT calculations show that the activation of the adsorbed CO₂ molecule can be enhanced via 582 583 partial oxidation of the least reactive Fe₃S₄(001) surface.

584	Table 4. Structural type, adsorption energies (E_{ads}) at 0 K, average interatomic distances (d),
585	bond angle (\angle), scaled wavenumbers for the fundamental vibrational modes and charge
586	transfers (Δq) calculated for a single CO ₂ molecule interacting with the different adsorption
587	sites considered for the partially oxidised 62.50 -Fe $_3S_4(001)$ surface. The presented vibrational
588	modes are the asymmetric stretching (v_{asym}), symmetric stretching (v_{sym}) and average bending
589	(δ) modes. The surface free energies (σ) at 0 K and work functions (Φ) are also reported.
590	Negative values of Δq indicate charge transfer from the surface to the adsorbate.

Adsorption site	0	S	Fe _A	Fe _B
Structural type	$\mu_3 - \eta^3$	μ ₂ –η²–(C, O)	η¹–0	η ¹ -0
E _{ads} (eV)	-0.746	0.016	-0.188	0.589
d(O _{mol} –Fe) (Å)	2.047	1.997	2.308	2.157
d(C–O _{surf}) (Å)	1.353			
d(C–S _{surf}) (Å)		1.899		
d(C=O _{mol}) (Å)	1.277	1.255	1.171	1.177
∠(O=C=O) (°)	127.98	130.47	177.46	172.67
<i>v</i> _{asym} (O=C=O) (cm ⁻¹)	1520	1690	2329	2272
<i>v</i> _{sym} (O=C=O) (cm ⁻¹)	1225	1088	1297	1278
δ (O=C=O) (cm ⁻¹)	795	671	581	594
<i>∆q</i> (e⁻)	-0.209	-0.537	0.003	0.003
arPhi (eV)	5.172	5.687	4.662	4.221
σ (meV Å⁻²)	65.087	72.524	70.534	78.110



Figure 8. Molecular adsorption of (a) CO_2 and (b) H_2O on the O and Fe_A , Fe_A sites, respectively, of the partially oxidised 62.5O-Fe₃S₄(001) surface. Side (top panels) and top (bottom panels) views are displayed. Layers containing atoms with dangling bonds are highlighted. Crystallographic directions are indicated. Fe_A atoms are in dark yellow, Fe_B atoms are in magenta, S atoms are in light yellow, O atoms are in red, H atoms are in white and C atoms are in black.

598 3.4.2. H₂O Interaction with the Partially Oxidised Fe₃S₄(001) Surface

599 In this section, we focus on discussing the molecular and dissociative adsorption of a single H_2O molecule on the partially oxidised 62.50-Fe₃S₄(001) surface, which are competing 600 processes that have been reported on the surfaces of oxides,^{151–159} sulfides^{160–164} and 601 metals.⁸⁵ For the molecular binding modes, we initially placed the adsorbate in the μ_3 - η^3 602 603 configuration, with both symmetry elements C_2 and σ_v perpendicular to the surface of our 604 catalyst, *i.e.* with the molecular O atom coordinating the under-coordinated cations and the 605 H atoms interacting with the exposed anions. We also explored the possibility that the C_2 axis and σ_v plane of the H₂O molecule are oriented parallel and perpendicular to the surface, 606 respectively, with the H atoms forming hydrogen-bonds with the surface O anions in the η^1 –O 607 structure. For the dissociative adsorption configurations, we introduced one OH group 608 609 coordinating to the exposed Fe_A and Fe_B ions and forming hydrogen-bonds with the surface anions in the $\mu_2 - \eta^2 - (0, H)$ configuration and with the dissociated H binding a nearby O atom. 610 611 The O (H) interacting atoms from the adsorbate were located at 1.8 (1.00) Å from the surface of our catalyst before carrying out a full geometry optimisation. However, we have not 612 investigated the η^1 -O adsorption configurations via H-bonds, as the H₂O rotated during 613 geometry optimisation leading to the same binding modes considered for the surface Fe ions. 614 615 Table 5 lists the exothermic adsorption energies calculated for H_2O , which are clearly more favourable than for the interaction of the CO₂ molecule with the 62.5O-Fe₃S₄(001) surface. 616 The dissociative adsorption modes release 0.507 and 0.219 eV less than the molecular binding 617 configurations at the (Fe_A, Fe_A) and Fe_B sites, respectively, suggesting that H₂O prefers to 618 remain undissociated at the surface of our catalyst. The order of decreasing stabilities are E_{ads} 619 $(Fe_A, Fe_A) < E_{ads}$ $(Fe_A) \ll E_{ads}$ (Fe_B) and E_{ads} $(Fe_A, Fe_A) \ll E_{ads}$ (Fe_B) for the molecular and 620 dissociative binding modes, respectively, showing that the adsorption energies have a strong 621 dependence on the type of Fe site. The trend of adsorption energies can be explained in terms 622

623 of the total number of dangling bonds of the cation sites. The most stable Fe_A, Fe_A binding site 624 with the $\mu_4 - \eta^4$ configuration has a total of three dangling bonds, see Figure 8 (b), whereas the 625 Fe_A and Fe_B sites with the $\mu_3 - \eta^3$ structure have two and one dangling bonds, respectively.

626 The average interfacial O–Fe binding distances for H₂O, which are marginally smaller than in the interaction of CO₂, are inversely proportional to the adsorption energies. The smallest 627 distances $d(O_{mol}-Fe) = 1.909$ and 2.060 Å were calculated for the thermodynamically least 628 stable dissociative adsorptions on the Fe_B and (Fe_A, Fe_A) sites, respectively, as the OH group 629 is a better ligand than H₂O. Our calculations indicate that the molecular and dissociated H₂O 630 631 molecules are able to form hydrogen-bonds with the surface O and S atoms. The interfacial 632 hydrogen-bonds with S are stronger than with O for the interactions with the Fe_A and (Fe_A) , Fe_A) sites, as evidenced by the H–S distances, which are smaller than the H–O_{surf} lengths. We 633 also found that the hydrogen-bond lengths with the surface O atoms are directly proportional 634 635 to the adsorption energies for the molecularly adsorbed H₂O molecule, with the largest value d(H-O) = 2.800 Å calculated for the ground state $\mu_4 - \eta^4$ adsorption mode. However, the 636 637 opposite effect of the binding energies was observed for the hydrogen-bond lengths with the S atoms, with the smallest value d(H-S) = 1.963 Å obtained for the most stable $\mu_4 - \eta^4$ binding 638 configuration. The H–O_{surf} distance of 2.704 Å suggests that the OH group adsorbed on the 639 (Fe_A, Fe_A) site in the $\mu_3 - \eta^3$ structure is able to form a hydrogen-bond with the surface, which 640 is not observed for the dissociative adsorption mode on the Fe_B site. Furthermore, the 641 dissociated H atom prefers to sit slightly closer at 0.043 Å to the surface O atom for the 642 interaction with the (Fe_A , Fe_A) site than for the Fe_B site, in agreement with their relative 643 adsorption energies. The adsorption of H₂O induces minor changes on the intramolecular H–O 644 distances, which are between 0.009 and 0.105 Å larger for the two types of adsorption modes 645 than in the isolated adsorbate. Our calculations show that the dissociated H atom diffuses 646 647 1.089 Å further away from the O of the OH group for the interaction with the most stable (Fe_A, Fe_A) site than with the least favourable Fe_B position. Note that we did not find evidence of a 648 649 dissociated H atom binding to a nearby surface S atom. The intramolecular bond angle is less sensitive to adsorption of H₂O than CO₂, with the largest value \angle = 107.73° calculated for the 650 molecular $\mu_3 - \eta^3$ adsorption on the Fe_A site. However, this geometrical descriptor can be used 651 to discriminate the dissociative from the molecular adsorption modes of H₂O, as the 652 intramolecular bond angle values are typically larger for the former than for the latter. 653

The three fundamental vibrational modes become smaller as a result of the molecular and 654 dissociative adsorption of H₂O. The asymmetric stretching vibrational mode is below 3549 655 cm⁻¹ for the molecular adsorption configurations, whereas it lies above 3578 cm⁻¹ for the 656 657 dissociated OH groups. The symmetric stretching vibrational modes, which appear in a wide range of values between 3356 and 2003 cm⁻¹, show no clear correlation with any of the 658 659 properties discussed. The bending vibrational modes were estimated between 1514 and 1468 cm⁻¹ for the molecular adsorption modes and the calculated values were smaller than 904 660 cm⁻¹ for the dissociative binding configurations. Table SI2 displays both the harmonic and 661 662 scaled vibrational modes observed for the adsorbed H₂O molecule. We calculated only minor 663 charge transfers, with the H₂O molecule receiving no more than 0.032 e^- for the most stable 664 molecular adsorptions on the Fe_A and (Fe_A, Fe_A) sites. Our results indicate that, unlike the linear CO₂ molecule, the activation and adsorption of the angular H₂O species does not 665 require charge density from the surface. The work functions are $\phi \sim 5.5$ eV for all the 666 adsorption modes, except for the molecular $\mu_4 - \eta^4$ binding configuration on the (Fe_A, Fe_A) site, 667 668 which is around 0.5 eV smaller than the value for the 62.50-Fe₃S₄(001) surface. The large reduction of the surface free energy after adsorption of the H₂O molecule demonstrates that 669 the 62.50-Fe₃S₄(001) surface becomes more stable, even compared to the catalyst interacting 670 with CO₂. In line with our results, previous calculations have also shown that the molecular 671 adsorption of water is energetically preferred over the dissociative mode for CaO,¹⁵⁶ 672 $CaF_2^{156,165}$ and the Fe₃S₄(001) surface.¹⁶¹ Moreover, the partially oxidised 62.5O-Fe₃S₄(001) 673 674 surface also displays an improved activity towards the adsorption of H₂O than the (001) facet of pure Fe₃S₄. 675

676	Table 5. Structural type, adsorption energies (E_{ads}) at 0 K, average interatomic distances (d),
677	bond angle ($ ightarrow$), scaled wavenumbers for the fundamental vibrational modes and charge
678	transfers (Δq) calculated for a single H ₂ O molecule interacting with the different adsorption
679	sites considered for the partially oxidised 62.50 -Fe $_3S_4(001)$ surface. The presented vibrational
680	modes are the asymmetric stretching (v_{asym}), symmetric stretching (v_{sym}) and average bending
681	(δ) modes. The surface free energies (σ) at 0 K and work functions (Φ) are also reported.
682	Negative values of Δq indicate charge transfer from the surface to the adsorbate.

Adsorption site	Fe _A	Fe _A , Fe _A		Fe _B	
Туре	Molecular	Molecular	Dissociative	Molecular	Dissociative
Structural type	$\mu_3 - \eta^3$	$\mu_4-\eta^4$	$\mu_3 - \eta^3$	μ3-η ³	$\mu_3 - \eta^3$
E _{ads} (eV)	-1.094	-1.318	-0.811	-0.510	-0.291
d(O _{mol} –Fe) (Å)	2.151	2.301	2.060	2.089	1.909
<i>d</i> (H1–O _{surf}) (Å)	2.436	2.800	2.704		
<i>d</i> (H2–O _{surf}) (Å)			0.980	1.466	1.023
d(H₁−S) (Å)				2.599	2.753
d(H₂−S) (Å)	2.242	1.963	2.650		
<i>d</i> (H₁−O _{mol}) (Å)	0.981	0.977	0.973	0.979	0.974
<i>d</i> (H₂−O _{mol}) (Å)	1.001	1.044	2.728	1.069	1.639
∠(H−O−H) (°)	107.73	106.65	175.77	106.77	110.15
ν _{asym} (H−O−H) (cm ^{−1})	3461	3549	3602	3511	3578
<i>v</i> _{sym} (H−O−H) (cm ^{−1})	3053	2339	3356	2003	2585
δ(H−O−H) (cm⁻¹)	1514	1468	807	1514	904
<i>∆q</i> (e⁻)	-0.005	-0.032	0.020	0.010	0.041
arPhi (eV)	5.660	4.996	5.504	5.623	5.405
σ (meV Å ⁻²)	61.693	59.506	64.453	67.392	69.531

683 **3.5.** H₂C₂O₄ Formation on the Partially Oxidised Fe₃S₄(001) Surface

We have examined three pathways for the catalytic conversion of CO₂ and H₂O into H₂C₂O₄ 684 685 on the 62.5O-Fe₃S₄(001) surface, see Figure 9. Our selection of mechanisms, which describe 686 the combination of the reactants into key intermediates that are transformed into the desired H₂C₂O₄ and by-product O₂, are discussed in terms of their structural, local electronic and 687 688 vibrational properties. The Helmholtz free energies (ΔF), which are plotted at the representative temperature of 300 K, are referenced in our energy diagrams compared to the 689 690 isolated reactants and the heterogeneous catalyst, *i.e.* two CO₂ molecules, one H₂O molecule and the $62.5O-Fe_3S_4(001)$ surface slab. Note that the vibrational frequencies of the adsorbed 691 692 species were scaled by 0.9632, which is the average of the scaling factors calculated for the isolated CO₂ and H₂O molecules, before estimating the vibrational entropy of the adsorbed 693

694 species, see Tables SI3 and SI4. For the purpose of comparison, the Helmholtz free energy 695 profiles at 0 and 600 K are provided in Figures SI3 an SI4. The first step, which is common to 696 the three pathways, is the co-adsorption of the reactants to the surface of the catalyst. Note 697 that the adsorbed species are indicated with the symbol *.



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Figure 9. Minimum energy pathways (MEPs) for the conversion of CO₂ and H₂O into H₂C₂O₄ on the partially oxidised 62.5O-Fe₃S₄(001) surface at 300K. Minimum states are denoted by bold lines and saddle point are represented by narrow lines linked by dashed lines. Energies are referenced to the isolated $2CO_2(g) + H_2O(g)$ molecules, including the surface slab. Adsorbed species are presented followed by the symbol *.

704 3.5.1. Oxalate Pathway

In line with our observations for the $\mu_3 - \eta^3$ adsorption of the single molecules, the initial co-705 706 adsorption configuration comprises CO₂ interacting with two nearby surface O atoms and four Fe cations, whereas the H₂O species was introduced coordinating the (Fe_A, Fe_A) site and 707 forming two hydrogen-bonds in the $\mu_4 - \eta^4$ structure. We had assumed that the 708 (co-)adsorption of the reactants is barrierless, but found that this is actually an exothermic 709 process that releases $\Delta F = 417$ meV and is therefore 2.39 eV less favourable than the sum of 710 the adsorption energies of the single molecules, as shown in Figure 9. The co-adsorbed CO₂ 711 molecules interact with the O sites at 1.340 and 1.399 Å, and with the C_2 axis remaining 712 perpendicular to the surface only for the closest adsorbed species, see Figure 10 (a). Both, 713 the perpendicular and tilted CO₂ molecules are able to coordinate to the surface Fe_A and Fe_B 714

cations at an average distance of ~2.1 Å, but the former uses its two O atoms, while the latter 715 employs only one O atom, which explains their different relative orientations with respect to 716 717 the surface. Our calculations suggest that both CO₂ molecules become activated upon 718 adsorption, with the apex angle \angle at < 126.34°, which is already smaller than the value calculated for the most stable $\mu_3 - \eta^3$ adsorption mode of the single molecule. The 719 intramolecular bonds in the CO₂ molecules are elongated to an average value of $d(C=O) \approx 1.28$ 720 Å, in agreement with the single molecule adsorption. The H₂O species sits on the (Fe_A, Fe_A) 721 722 site at approximately the same distance calculated for the most stable single molecule $\mu_4 - \eta^4$ adsorption, with the intramolecular distances and bond angle remaining relatively 723 undisturbed. However, the surface S and O atoms forming the hydrogen-bonds are 0.120 Å 724 further away and 0.262 Å closer to the H₂O molecule, respectively, than in the ground state 725 726 single molecule adsorption configuration. The charge analysis of the co-adsorption state 727 indicates that the perpendicularly adsorbed CO₂ molecule received 0.027 e⁻ less charge than in the case of the single molecule adsorption, which was compensated for by the identical 728 electron density gained by the tilted species. On the other hand, the H₂O adsorbate received 729 only a negligible charge of $\Delta q = 0.002 \text{ e}^-$, which is considerably less than in the case of the 730 single molecule $\mu_4 - \eta^4$ adsorption. We attribute the smaller charges received by the H₂O and 731 perpendicular CO_2 molecules to the fact that they share the same surface Fe_A ion. 732



Figure 10. Side views of the adsorption configurations of (a) reactants $2CO_2^*$ and H_2O^* , intermediates (b) $C_2O_4^*$ and H_2O^* and (c) $HC_2O_4^*$ and OH^* , as well as (d) final products $H_2C_2O_4^*$ and O^* of the oxalate pathway on the partially oxidised 62.5O-Fe₃S₄(001) surface. Interatomic distances are provided in Ångstroms; crystallographic directions are indicated. Fe_A atoms are in dark yellow, Fe_B atoms are in magenta, S atoms are in light yellow, O atoms are in red, H atoms are in white and C atoms are in black.

733

740 The oxalate pathway leads to the combination of the two CO₂* molecules into the C₂O₄* 741 intermediate, which is an endothermic process with a reaction free energy of 2.100 eV and a saddle point A at 3.296 eV, as shown in Figure 9. The large free energy barrier of this process 742 743 is required to break the strong covalent bonds within the adsorbed carbonate-like groups, leading to the C atoms moving to at least 3.0 Å away from the nearest surface O atoms, see 744 Figure 10 (b). The product of this elementary step, which is the least stable intermediate, is a 745 staggered C₂O₄* group with the dihedral angle \angle (O–C–C–O) = 66° and the point group of 746 reduced symmetry D_{2d} . The C₂O₄* species is anchored to the exposed Fe cations of the surface 747 through three O atoms, with $d(O-Fe) \sim 2.0$ Å, which is slightly smaller than for the co-748 adsorption state. The C–C bond distance is 1.539 Å for the C₂O₄* intermediate, whereas the 749 750 intramolecular C=O distances and O=C=O angles do not change noticeably with respect to the 751 previous state. The calculated Bader charges show that the $C_2O_4^*$ species draws $-1.367 e^$ from the surface, which is considerably larger than for two co-adsorbed CO₂ molecules, 752 753 explaining the low stability of this intermediate.

The $C_2O_4^*$ species accepts one H atom from the H_2O molecule to form $HC_2O_4^*$ in state 4 of our energy diagram, see Figure 9. The free energy released during this elementary step is

0.846 eV, but it requires 1.679 eV to cross the saddle point B. The protonated O atom, with 756 the typical distance of 0.98 Å for the O–H bond, is the closest to the donor H₂O molecule, 757 belonging to the bidentate CO₂ moiety in the previous state, as shown in Figure 10 (c). The 758 759 dihedral angle becomes more orthogonal with \angle (O–C–C–O) = 81.82° after protonation, 760 whereas the largest intramolecular distances of $d(C-O) \sim 1.31$ Å were calculated in each molecular moiety for the protonated O and the O coordinating the surface Fe ion. The OH* 761 fragment moves 0.442 Å closer to the Fe_A ion that is shared with the $HC_2O_4^*$, as the basicity 762 of the former increases and the latter decreases, compared to H_2O^* and $C_2O_4^*$ respectively. 763 However, the shift of the OH* species happens at the cost of an increase in the H…S hydrogen 764 bond-distance by 0.383 Å. The protonation reduces almost by half the charge of the HC₂O₄* 765 intermediate to $q = -0.704 e^{-1}$ with respect to C₂O₄*, whereas the OH* is able to retain -0.650 766 767 e^{-} from the proton that it ceded.

768 Figure 9 illustrates that the HC₂O₄* intermediate is capable of admitting a second proton from the OH* group, which is also an endothermic process with a reaction free energy of 2.202 eV 769 and an activation free energy C of 4.027 eV. The protonation increases the symmetry of 770 $H_2C_2O_4^*$ to point group $C_{2\nu}$, implying that the molecule becomes essentially flat with the 771 dihedral angle \angle (O-C-C-O) = 164°, see Figure 10 (d). The intramolecular bond distances 772 d(C-C) = 1.548 Å, $d(C=O) \sim 1.22$ Å, $d(H-O) \sim 1.00$ Å and $d(C-OH) \sim 1.341$ Å and angle $\angle (O-C-O)$ 773 = 125.8° of $H_2C_2O_4^*$ are very similar to the values calculated for the $C_2O_4^*$. Our calculations 774 suggest that H₂C₂O₄* is still coordinating one Fe_A and two Fe_B, with the smallest Fe–O distance 775 of 2.248 Å for the unprotonated O and the largest distances of 2.92 Å for the protonated O 776 atoms. The O* species moves approximately 0.2 Å towards the bulk after losing its H atom. 777 The transfer of the second proton quenches the negative charge of $H_2C_2O_4^*$ to $-0.037 e^-$, 778 whereas O* has -1.108 e⁻, almost duplicating the electron density of OH* in the previous 779 780 state

In the final step, the surface releases one $H_2C_2O_4$ and 0.5 O_2 molecules, leaving the system ready for the next catalytic cycle. These desorptions are endothermic processes that require 0.477 eV. The final state is the overall highest in free energy in our energy diagram, lying 3.518 eV above the reactants. To calculate these energies, we modelled the isolated O_2 molecule in the triplet state and the isolated $H_2C_2O_4$ in the point group C_{2h} , which are the electronic and structural ground states, respectively, of these species.

787 3.5.2. Carboxylate Pathway

The protonation of one of the CO₂* species in the carboxylate pathway leads to the most 788 789 stable intermediate in this study, which is 1.157 eV below the reactants in our free energy diagram, see Figure 9. This exothermic process has a reaction free energy of -0.740 eV and a 790 transition state D at 1.324 eV. The O–H bond distance is 1.01 Å for the bicarbonate-like group, 791 792 which also forms a hydrogen-bond of 1.727 Å to the neighbouring carbonate-like species, as 793 shown in Figure 11 (a). The intermolecular hydrogen-bond is enabled by the rotation of the two intermediates around the C_2 axis perpendicular to the surface, which also reduces their 794 interfacial C–O distance by an average of 0.05 Å and their apex O–C–O angles by at least 5°. 795 The protonation weakens the C–OH bond, as its distance increases by 0.1 Å, which implies 796 that part of the electron density is shared with the newly added H atom. We found that the S 797 798 atom that formed the hydrogen-bond to the H₂O molecule, moved to 1.751 Å away from the CO_2^* group during the proton transfer. The Bader analysis assigns 0.445 e⁻ to the CO_2H^* , 799 800 which receives a large Coulomb attraction from the negatively charged surface O atom, explaining the stability of these intermediates. Despite the CO_2^* species remaining bent, it 801 donates electron density back to the surface S atom and is only able to keep -0.032 e⁻. Our 802 calculations show that the OH* species displays very similar Fe–O and O–H bond distances 803 and electron density charge as the OH* group that coexists with HC₂O₄* in the oxalate 804 pathway. 805



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Figure 11. Side views of the adsorption configurations of the intermediates (a) CO_2H^* , CO_2^* and OH* and (b) $2CO_2H^*$ and O* of the carboxylate pathway, as well as intermediates (c) CO_3H^* , CO_2^* and H* and (d) CO_3H^* and CO_2H^* of the bicarbonate pathway on the partially oxidised 62.5O-Fe₃S₄(001) surface. Interatomic distances are provided in Ångströms; crystallographic directions are indicated. Fe_A atoms are in dark yellow, Fe_B atoms are in magenta, S atoms are in light yellow, O atoms are in red, H atoms are in white and C atoms are in black.

814 Figure 9 depicts the protonation of the second CO_2^* species, which is an endothermic process 815 with a reaction free energy of 1.547 eV and a saddle point *E* at 3.066 eV. We found that the 816 structural properties of the two CO₂H* species, such as the H–O and C–O bond distances and the O–C–O angle are very similar to the protonated group of the previous state, as shown in 817 818 Figure 11 (b). However, our calculations show that each CO_2H^* group can only coordinate a single Fe_B cation via the unprotonated O atom at the slightly different distances of 2.058 and 819 2.245 Å. The two CO₂H* species have a positive charge of ~0.5 e^- , whereas the O* gains 820 approximately half of one electron. 821

The combination of the two carboxylate groups to form the adsorbed $H_2C_2O_4^*$ molecule is an endothermic process with a reaction free energy of 2.650 eV, which is 0.448 eV larger than the free energy required to transfer the second proton to the $HC_2O_4^*$ in the oxalate pathway, see Figure 9. The free energy of the saddle point *F* is 3.998 eV for the formation of the C–C bond in the carboxylate pathway, whereas it is ~0.7 eV smaller for the saddle point *A* in the oxalate pathway.

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828 3.5.3. Bicarbonate Pathway

In the bicarbonate pathway, the H₂O* molecule dissociates and the H* atom migrates to a 829 nearby surface O atom, whereas the OH* group diffuses to one of the CO₂* species. This is an 830 endothermic process with a reaction free energy of 0.699 eV, *i.e.* the lowest in this study, and 831 832 a transition state G at 2.318 eV, shown in Figure 9. The CO₃H* species moves 1.921 Å away from the surface upon hydroxylation, remaining only bound to the undercoordinated Fe ions 833 at the average distance of 2.0 Å in the $\mu_2 - \eta^2 - (0, 0)$ configuration, see Figure 11 (c). The C–OH 834 distance is 1.330 Å in the CO₃H* species, which tilts to allow the formation of a short 835 hydrogen-bond of 1.764 Å with the CO_2^* molecule. The interfacial Fe_B–O and C–O bonds 836 between the CO₂* molecule, which prefers to stay in the side-on $\mu_2 - \eta^2 - (C, O)$ structure, and 837 the surface are 2.000 and 1.413 Å, respectively. The intramolecular C=O bond distance is 1.28 838 Å for the CO₃H* species and 1.300 and 1.233 Å for the O end coordinating the surface Fe_B and 839 the free O end, respectively, of the CO_2^* molecule. The dissociated H* atom, which stays 840 841 coordinated to the S atom that was part of the hydrogen-bond, diffuses to the subsurface 842 layer to increase its stability. The incorporation of the OH group into the CO₃H* species increases its negative charge to -0.672 e⁻ with respect to the value of -0.213 e⁻ for the CO₂* 843 molecule, whereas the subsurface H* atom becomes almost neutral with 0.053 e⁻. 844

Figure 9 depicts the protonation of the second CO₂* molecule in our free energy profile. This 845 elementary step releases -0.384 eV, which is the smallest exothermic free energy value 846 reported in this study, and has a saddle point of 2.120 eV. The newly formed CO₂H* species 847 remains bound to a surface O anion at 1.295 Å and to a Fe^B ion at 1.755 Å through the 848 849 molecular C and OH group, respectively, as shown in Figure 11 (d). Our calculations suggest that the intramolecular OH distance is 0.976 Å. However, we found that the proton diffusion 850 only leads to negligible changes in the structure of the CO₃H* species, including the length of 851 the Fe–O and hydrogen-bonds with the surface and CO₂H* intermediate, respectively. The 852 protonation provides the CO_2^*H intermediate with a positive charge of 0.382 e⁻, whereas 853 CO_3H^* increases its negative charge by 0.038 e⁻. 854

Finally, the coupling of the CO₃H* and CO₂H* species is an endothermic elementary step with a reaction free energy F_R = 3.143 eV and the largest activation energy value F_{SP} = 5.177 eV calculated in this work, see Figure 9.

4. Conclusions

We have used DFT methods to model the catalytic conversion of CO₂ and H₂O into H₂C₂O₄ on 859 the partially oxidised Fe₃S₄(001) surface. First, we have modelled the bulk phase of Fe₃S₄ and 860 found that the structural, electronic and magnetic properties are in good agreement with 861 previous reports. We have elucidated the relaxation patterns of the interplanar distances, as 862 863 well as the values for the surface energies, atomic charges, atomic magnetic moments and work functions for the two reconstructed Tasker type 3 terminations of the $Fe_3S_4(001)$ surface. 864 865 The thermodynamically most stable facet, which terminates in two-fold Fe_A ions with $(\sqrt{2} \times \sqrt{2})R45^{\circ}$ symmetry, has the largest relaxation, work function, ionic character and 866 magnetisation. We have predicted the phase diagram as a function of the ratio of partial 867 868 pressures of H_2O and H_2S and temperature and found that the processes of replacing 869 sequentially each S atom with dangling bonds by an O atom are endothermic. Only selected 870 coverages of O are allowed in the Fe₃S₄(001) surface, with 62.5% of partial oxidation becoming prominent at the typical experimental conditions in which samples of the catalyst are calcined. 871 We have also reported the interaction between single molecules of CO₂ and H₂O and the 872 Fe₃S₄(001) surface partially oxidised by 62.5%. Adsorption at the O site, which is energetically 873 preferred, bends and activates the CO₂ molecule and the σ^* antibonding molecular orbital 874 875 receives electronic density from the partially oxidised $Fe_3S_4(001)$ surface. The H₂O molecule 876 releases the largest adsorption energy when it coordinates two surface Fe_A cations and forms hydrogen-bonds with the exposed S and O anions, but no charge transfers were identified. 877 878 The co-adsorption of two CO₂ and one H₂O molecule at nearby surface sites is also an 879 exothermic process, although 239 meV less favourable than the interaction of the single 880 species.

881 We have calculated three minimum energy pathways for the formation of $H_2C_2O_4$, where we have considered the formation of an oxalate, carboxylate or bicarbonate intermediate in the 882 883 first elementary step. The energy profiles show that $C_2O_4^*$ and $HC_2O_4^*$ are particularly unstable intermediates, whose formation requires crossing saddle points of large energy 884 885 barriers. However, the carboxylate CO₂H* intermediate is the most stable species when it is co-adsorbed with CO₂*, and to a lesser extent, with the bicarbonate CO₃H*. The saddle points 886 887 with the largest energies were observed for the coupling of the C-C bond, whereas the 888 transfer of protons are the elementary steps with the smallest activation barriers. Our

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reaction mechanisms suggest that the energy released during the co-adsorption of the reactants is not enough to cross all the saddle points or to reach the final state. We would therefore argue that the feasibility of the conversion of CO_2 into $H_2C_2O_4$ depends strongly on the source of hydrogen and the reactivity of the surface. The high energy saddle points and intermediate species would become accessible if the catalytic process over the partially oxidised Fe₃S₄(001) surface is carried out within a continuous electrochemical cell with an appropriate voltage.

Future work will be focused on calculating the catalytic formation of pyruvic acid, which is an important intermediate in several metabolic pathways, on the partially oxidised surfaces of Fe₃S₄, including not only the stable (001) surface, but also more reactive surfaces such as the (111) plane. The reaction profiles for the conversion of CO₂ into pyruvic acid will allow us to compare this process and interconnect it into the catalytic formation of acetic acid, formic acid and oxalic acid.

902 Conflicts of interest

903 The authors have no conflicts of interest to declare.

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