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Theoretical Study of the Adsorption/Dissociation Reactions of Formic Acid on the α -Al₂O₃(0001) Surface

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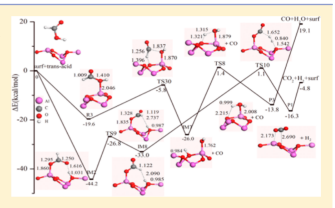
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5 Supporting Information

ABSTRACT: Formic acid was used as the model of lauric acid to investigate the microscopic mechanism of the anti-icing behavior and was checked to find out if it can be catalyzed to produce H_2 for fuel cells by the α -Al₂O₃(0001) 2 × 2 supercell slab model. The density functional theory with the all-electron double numerical polarized basis sets was employed. The results show that when it involves the carboxyl O and hydroxyl H atom the 1,2-dissociated adsorbate is the most stable intermediate on the dry Al₂O₃(0001) surface and is energetic barrier free to form the fairly stable ROCO- and HO-covered surface with the binding energy of 59.5 kcal/mol, and this dissociation mode has the lowest energy barrier of 14.9 kcal/mol to form the HOCO-and H_2 O-covered surface after the surface is fully hydroxylated. The energetic barrier of the HCOOH dehydrogenation and



dehydration reactions on the alumina surface decreased considerably from 65.3 to 30.6 kcal/mol and from 62.1 to 26.8 kcal/mol, respectively, in comparison with the gaseous decomposition. The dissociated configuration of lauric acid was tested, and it was found that it dissociated with free energy barrier through 1,2-hydrogen migration into the $C_{11}H_{23}OCO$ - and HO-covered surface with a binding energy of 60.7 kcal/mol. The present theoretical work is useful to gain some new insights on the microscopic interaction mechanism of the superhydrophobic alumina surface fabrication procedure and on the heterogeneous catalysis reactions of the H_2 production.

1. INTRODUCTION

28 The production of H_2 is a great challenge in heterogeneous 29 catalysis. The decompositions of several carbon-containing 30 compounds such as methanol, ethanol, and dimethyl ether are 31 used as sources of H_2 .^{1–4} As the simplest and the most 32 abundant fatty acid, 5 formic acid is the potential source of H_2 33 for fuel cells.

Metal oxide is currently used for several applications of technological interest as a consequence of its electrical behavior and catalytic activity. A focus of such exploration is the interfacial interaction of the highly valuable surface. Alumina is used extensively as a substrate for heterogeneous catalysis. Among the many polymorphs of alumina, the termodynamically stable α -Al₂O₃, or corundum, has been studied most extensively, both experimentally and theoretically. Below the substrate for heterogeneous of the substrate for heterogeneous are substrated for heterogeneous and the substrate for heterogeneous of the substrate for heterogeneous are substrated for heterogeneous are substrated for heterogeneous and the substrate for heterogeneous are substrated for heterogeneous

43 Ojeda et al. ²² found that formic acid can form H_2 at high 44 chemical potentials with the catalysis of Au on the Al_2O_3 45 surface. The adsorption of acetate species (CH_3COO^-) on the 46 Ag/Al_2O_3 catalysis was studied by Gao et al. ²³ Solymosi et al. ¹ 47 studied the dehydration of formic acid which was favored on 48 the Au-deposited Al_2O_3 surface, and pure CO-free H_2 was 49 obtained on Au/SiO_2 and Au/CeO_2 surfaces at below 473 K.

However, the role of supporter Al_2O_3 in this reaction is 50 unknown. Carlos-Cuellar et al.²⁴ investigated the heteroge-51 neous uptake kinetics of formaldehyde on α -Al $_2O_3$ with a 52 Knudsen cell reactor at 295 K. Tong et al.²⁵ studied the 53 atmospheric heterogeneous reactions of formic acid on dust 54 particles (Al $_2O_3$) at ambient conditions by using a DRIFTS 55 reactor and found that the carboxylates had formed during the 56 uptake process.

Additionally, we have obtained the superhydrophobic 58 alumina surface with anti-icing property by etching with 59 chemicals and modifying with low-surface energetic subject 60 fatty acid, 26,27 but the reaction mechanism between the surface 61 and fatty acid is unknown. In order to understand more about 62 the anti-icing behavior of the superhydrophobic surface, the 63 microscopic adsorption/dissociation mechanisms of fatty acid 64 on the alumina surface should be investigated. Formic acid is 65 used as the model of fatty acid to reveal the reaction 66 mechanisms on the α -Al₂O₃(0001) surface. Furthermore, the 67 catalytic mechanisms of dehydrogenation/dehydration reac-

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69 tions of formic acid on the α -Al₂O₃(0001) surface are studied 70 using the first-principles calculation based on density functional 71 theory.

2. COMPUTATIONAL METHOD

72 For the slab calculation, we applied periodic, density-functional 73 theoretical (DFT) calculations implemented in the Dmol³ 4 program. To accurately account for the van der Waals (vdW) 75 interactions, self-consistent field (SCF) energies of the systems 76 were corrected for dispersion forces using the DFT-D3 method 77 developed by Grimme et al. The Kohn—Sham equation was 78 solved in a self-consistent manner under the generalized 79 gradient approximation (GGA). Electron exchange and 80 correlation energies were calculated using Becke—Lee—Yang—81 Parr (BLYP) with the all-electron double numerical (DND) 82 and triple numerical polarized (TNP) basis sets. DND basis 83 sets include a d-type polarized function on heavy atoms, while 84 the TNP basis sets include additional sets of d-type polarized 85 functions on all atoms. The numerical basis set solving the 86 atomic DFT equation is capable of minimizing or even 87 eliminating basis set superposition error (BSSE). 33

The atom-centered grids were used for the numerical 89 integration with a real-space cutoff of 4.2 Å. The self-90 consistent-field (SCF) convergence criterion was set to the 91 root-mean-square change in the electronic density to be less 92 than 10^{-5} eV without thermal smearing. The equilibrium 93 geometries of reactants, intermediates, and products were fully 94 optimized using the Broyden Flecher Goldfarb Shanno (BFGS) 95 method within the delocalized coordinates. Transition states 96 were obtained tentatively using the synchronous transit 97 methods and then refined through the eigenvector following 98 optimization. The convergence criteria applied for geometry 99 optimization were enforced to 2×10^{-5} au for energy, 0.004 100 au/Å for force, and 0.005 Å for maximum displacement. 101 Harmonic vibrational frequencies of the optimized structures 102 were computed by diagonalizing the mass-weighted second-103 derivative matrix which was built using two-sided finite 104 differences with a displacement step of 0.01 Å and with 105 HessianAtoms sets of relaxed atoms.

The Al-terminated α -Al₂O₃(0001) basal surface was built by 107 cleaving the bulk corundum crystal whose lattice constants were a = 4.759 Å and c = 12.991 Å. It consisted of an 18-layer-109 thick slab with six complete AlO₃Al layers (Al₁₂O₁₈). To avoid 110 unphysical interlayer interactions, the slabs were separated by a 111 vacuum region of 10 Å. The BLYP/DND with Γ -point 112 calculation was performed with a 2 × 2 supercell model 113 $(Al_{48}O_{72})$ in this work. In all calculations, we allowed the top 114 five atomic layers of the slab and their adsorbates to relax, 115 whereas the other 13 atomic layers were held fixed at the bulk 116 positions. In this work, we calculated the adsorption energies according to the following equation, $E_{ads} = E(slab + adsorbate)$ 118 - E(slab) - E(adsorbate), in which E(slab + adsorbate), 119 E(slab), and E(adsorbate) were the calculated electronic 120 energies of species adsorbed on the surface, the bare surface, 121 and the gas-phase molecule, respectively.

3. RESULTS AND DISCUSSION

3.1. α -Al₂O₃(0001) 2 × 2 Supercell Surface Energy. The BLYP/DND with Γ -point calculations are performed with a 2 124 × 2 supercell bulk model (Al₄₈O₇₂) with the lattice constants of 125 a=9.518 Å and c=12.991 Å and the Al₄₈O₇₂(0001) surface 126 with the experimental lattice constants of a=9.518 Å and c=12.991 Å and c=1

22.506 Å. The surface Al atoms relaxed toward the surface O $_{127}$ atoms, decreasing the interlay spacing from 0.840 to 0.176 Å. $_{128}$ The changes of the $\rm Al_{48}O_{72}(0001)$ surface before and after $_{129}$ being optimized are shown as Figure 1. The relaxation ratio is $_{130~\rm fl}$

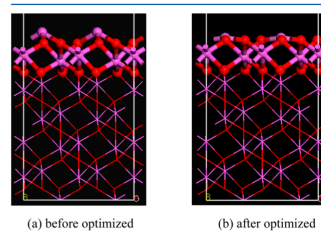


Figure 1. Changes of the α -Al $_2$ O $_3$ (0001) surface before and after being optimized. The pink and red balls represent Al and O atoms, respectively.

79% in comparison with the experimental data of 51-63%. 34,35 131 The geometric reconstruction of the surface can be better 132 characterized using the O–Al–O bond angle and the AlO₃ 133 tetrahedral. The O–Al–O bond angle increases from 101.1° in 134 the bulk to 118.9° in the surface, and the AlO₃ tetrahedral 135 increases from 103.8° to 159.6°. The relaxation ratio is 17.6% 136 and 53.8%, respectively. The surface energy is 1.70 J/m² (SE = 137 ($E_{\rm surface} - E_{\rm bulk}$)/2A, where A is the basal surface area) which is 138 nearly the same as the experimental value of 1.69 J/m². We 139 know that the four Al atoms on the surface have the same 140 chemical surrounding, which means that they have the same 141 reactive site as a Lewis acid from the figure.

3. 2. Free HCOOH Molecule. In the gas phase, the isolated 143 HCOOH molecule has cis and trans conformations, and it 144 prefers the trans position. 37 The trans is 4.7 kcal/mol more 145 stable than the cis at the theoretical level of BLYP/DND/1 \times 1 146 \times 1 in this study, as other studies 38,39 depicted. The bond 147 distances and bond angles of the optimized trans-HCOOH 148 structure are depicted in Table 1. Our results obtained from the 149 table agree well with experiment 38 and with previously 150 published DFT results. 39

Table 1. Bond Distances (Å) and Bond Angles (deg) for the Gas-Phase trans-HCOOH

	C-O	c=o	С-Н	О-Н	O-C-O	С-О-Н	
this paper	1.367	1.212	1.104	0.992	125.3	105.9	
experiment ^a	1.361	1.217	1.106	0.984	123.4	107.3	
${\it calculation}^b$	1.356	1.206	1.108	0.982	124.9	106.1	
^a From ref 38. ^b From ref 39.							

3.3. Theoretical Level. Table 2 shows the influences of 152 t2 basis sets, k-point mesh, atomic orbital cutoff value, and vacuum 153 thickness on the binding energy of adsorbed compound IM1 154 (optimized geometry is shown in Figure 2). The binding energy 155 f2 is 59.5 kcal/mol at the theoretical level of BLYP/DND/1 \times 1 \times 156 1 with the atomic orbital cutoff value of 4.2 Å and the vacuum 157 thickness of 10 Å. If the k-point is changed to be 3 \times 3 \times 3, the 158

Table 2. Adsorption Energies of IM1 at Different Theoretical Levels (kcal/mol)

effect factor		ac	lsorption energies at d	ifferent theoretical lev	els		
basis set	DND	DND	DND	DNP	TNP	DND	
k-point	$1 \times 1 \times 1$	$3 \times 3 \times 3$	$1 \times 1 \times 1$	$1 \times 1 \times 1$	$1 \times 1 \times 1$	$1 \times 1 \times 1$	
cutoff (Å)	4.2	4.2	4.8	4.2	4.2	4.2	
vacuum thickness (Å)	10	10	10	10	10	20	
$\Delta E^a(ext{kcal/mol})$	-59.5	-58.1	-56.2	-57.2	-59.6	-57.7	
^a Relative energies to the isolated formic acid and the free α -Al ₂ O ₃ (0001) surface.							

2.005 1.7371.888 1.721 1.874 R2 -33.3 R1 -32.2 R3 -28.4 R4 -25.7 2.188 1.894 1.870 IM8 -39.9 IM3 -51.8 IM1 -59.5 IM2 -54.7 IM12 -15.8 IM13 -7.4 IM14 -8.8 IM17 3.1 0.989 2.240 1.888 IM19 -54.2 IM20 -41.2 IM18 6.3

Figure 2. Geometries of the adsorption structures of formic acid on the α -Al₂O₃(0001) surface. Bond distances are in angstroms, and angles are in degrees.

 $_{160}$ value is changed to be 4.8 Å, the binding energy is 56.2 kcal/ $_{161}$ mol. The value is 57.2 and 59.6 kcal/mol with the basis sets $_{162}$ being changed to be DNP and TNP, respectively. The binding $_{163}$ energy is 57.7 kcal/mol with changing vacuum thickness of 20 $_{164}$ Å. From the above results, it can be concluded that the $_{165}$ influences of basis sets, k-point mesh, vacuum region, and $_{166}$ atomic orbital cutoff value have little significance on the $_{167}$ research system. All calculations are performed at the $_{168}$ theoretical level of BLYP/DND/1 \times 1 \times 1 with the atomic $_{169}$ orbital cutoff of 4.2 Å and vacuum thickness of 10 Å.

Additionally, in order to certificate the model of the relaxed $_{170}$ top five layers is proper in this work, we compare the energetic $_{171}$ barrier of the reaction from R1 to IM12, TS16 (optimized $_{172}$ geometries are shown in Figure S1, Supporting Information), $_{173}$ with the relaxed top five-layer model and the relaxed top nine- $_{174}$ layer model. They are calculated to be 24.3 and 26.7 kcal/mol, $_{175}$ respectively. So the relaxation of the top five layers works well $_{176}$ in this study.

3.4. Single HCOOH Adsorption/Dissociation at the 2 $_{178}$ × 2 Supercell. A single isolated HCOOH molecule in all $_{179}$ models with no restrictions can approach the α -Al₂O₃(0001) $_{180}$

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181 surface to form two types of physical adsorbates, namely, 182 carboxyl adsorption and hydroxyl adsorption, including R1, R2 183 and R3, R4, respectively, and to form one type of chemical 184 dissociated adsorbate, including IM1 and IM2, two kinds of 185 dissociated compounds. Their optimized geometries are shown 186 in Figure 2. Their binding energies are 32.2, 33.3, 28.4, 25.7, 187 59.5, and 54.7 kcal/mol, respectively, as shown in Table S (Supporting Information) ($E_{\rm ads} = E({\rm slab} + {\rm adsorbate}) - 189 E({\rm slab}) - E({\rm adsorbate})$, $E({\rm slab} + {\rm adsorbate})$, $E({\rm slab})$, and 190 $E({\rm adsorbate})$ are the energies of the adsorption compounds on 191 the surface, the bare ${\rm Al_2O_3}(0001)$ surface, and the free formic 192 acid molecule, respectively).

In the molecular adsorption compounds R1 and R2 of the 193 carbonyl type, formic acid interacts with the surface through the carboxyl oxygen and the CH hydrogen. If the CH hydrogen atom locates at the 2-O_s (the nearest crystal oxygen to the Al atom) to form a five-membered ring, the binding energy of R1 is 32.2 kcal/mol. If the H atom locates at the 4-O_s (the second nearest crystal oxygen to the Al atom) to form a seven-200 membered ring, the binding energy of R2 is 33.3 kcal/mol. For the hydroxyl type adsorption compounds R3 and R4, the main 202 interactions involve the hydroxyl O atom and H atom of formic 203 acid. It forms an intermolecular four-membered ring of the 204 hydroxyl H with the 2-O_s in R3, and it is a six-membered ring 205 of the hydroxyl H with the 4-O_s in R4. The binding energy of 206 R3 and R4 is 28.4 and 25.7 kcal/mol, respectively. Among the molecular adsorbates, R2 is the most stable complex.

If the formic acid approaches the surface through the carboxyl O atom and hydroxyl H atom, it can dissociate into HCOO and H without overcoming any energetic barrier. If the lissociated H atom interacts with the 2-O_s atom, the intermediate is IM1 with a six-membered ring. If the dissociated H atom interacts with the 4-O_s atom, the intermediate is IM2 H atom interacts with the 4-O_s atom, the intermediate is IM2 lissociated with an eight-membered ring. The binding energy of IM1 and liss IM2 is 59.5 and 54.7 kcal/mol, respectively. To certificate the energetic barrier free process of this kind of dissociation, we lift fixed the C=O and O-H bonds of formic acid, scanned the liss Al_s-O_f bond length, and optimized the geometries of the molecular adsorption compounds. The adsorption energies of the compounds along the Al_s-O_f bond lengths are shown in Figure 3. We can see that the binding energies are increasing

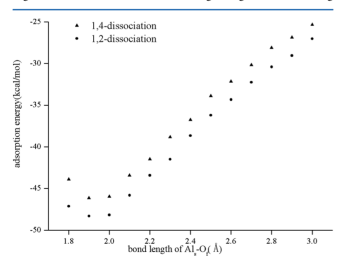


Figure 3. Adsorption energetic changes with formic acid approaching the α -Al₂O₃(0001) surface through the carboxyl O atom and hydroxyl H atom. Bond distances are in angstroms, and angles are in degrees.

along with formic acid approaching the surface until it reaches 222 1.9 Å, the bond length of the Al–O bond. This confirms the 223 above result that it is free-barrier energetic dissociation when 224 interactions involve carboxyl O and hydroxyl H. 225

The geometric changes after formic acid adsorption/ 226 dissociation at the 2 \times 2 supercell α -Al₂O₃(0001) surface are 227 listed in Table 3. As depicted, the bond length of Al_s-O_f is 228 t3 between 1.997 and 2.115 Å for the physical adsorption 229 compounds R1, R2, R3, and R4. For the dissociated 230 compounds, the bond length of Al_s-O_f is 1.836 and 1.860 Å 231 in IM1 and IM2, respectively, and there is a new bond O_s-H_f 232 with the length of 1.005 and 1.031 Å in IM1 and IM2, 233 respectively. The adsorption/dissociation of formic acid can 234 draw up the surface Al_s atom, as shown by d_{Al-O} which is the 235 distance of the outmost Al_s to the next nearest O_s . The d_{Al-O} 236 increases from 0.176 to 0.470, 0.456, 0.375, 0.421, 0.554, and 237 0.593 Å for R1, R2, R3, R4, IM1, and IM2 complexes, 238 respectively. Obviously, the dissociation intermediates can draw 239 up the Al atom more than the adsorption compounds, and R1 240 and R2 can draw up a little more than R3 and R4. We note that 241 this bond length of Al_s-O_s in IM1 is 1.869 Å which is even 242 0.113 Å longer than that in the bulk of 1.856 Å. The O₅Al₅O₅ 243 bond angle and the Al_sO_{s3} tetrahedral of the outmost layer of 244 the surface decrease as the Al_s atom is drawn up. They decrease 245 about 6° and 28° for the carbonyl adsorbates, about 4° and 21° 246 for the hydroxyl adsorbates, and about 14° and 40° for the 247 dissociated intermediates, respectively. In the dissociated 248 intermediate IM1, the angle and tetrahedral is close to that in 249 the bulk, and they are 104.8° and 119.2° in IM1 and 101.1° and 250 103.8° in the bulk.

There are four dissociation pathways for the four molecular 252 adsorption compounds. The adsorption energies are shown in 253 Table S (Supporting Information), and the optimized geo- 254 metries are shown in Figure S1 (Supporting Information). The 255 dissociated intermediations are COOH and H for the carbonyl 256 adsorbates. The first route, starting from the initial molecular 257 adduct of R1, possesses a five-membered ring transition state, 258 TS16. The hydrogen atom migrates from formic acid to the 2- 259 O_s to form the 1,2-dissociated intermediate IM12 with the ²⁶⁰ binding energy of 15.8 kcal/mol. The barrier height for TS16 is 261 24.3 kcal/mol. In the second dissociation path, the hydrogen 262 atom migrates from formic acid to the 4-O_s through TS25 to 263 form the 1,4-dissociated intermediate IM17 with the 264 exothermicity of 3.1 kcal/mol, and the barrier height for 265 TS25 is as high as 54.5 kcal/mol. Obviously, for the carbonyl 266 adsorbates, the 1,2-dissociation is easier than the 1,4-267 dissociation, and the 1,2-dissociated intermediate IM12 is 268 18.9 kcal/mol more stable than the 1,4-dissociated intermediate 269 IM17. The dissociated intermediations are HCOO and H for 270 the hydroxyl adsorbates in the later two pathways. If the route 271 starts from R3, it possesses a four-membered ring transition 272 state, TS31, with the energetic barrier of 15.0 kcal/mol to form 273 the 1,2-dissociated intermediate IM19 with the binding energy 274 of 54.2 kcal/mol. If the route starts from R4, it possesses a six- 275 membered ring transition state, TS36, with the energetic barrier 276 of 7.2 kcal/mol to form the 1,4-dissociation intermediate IM20 277 with the binding energy of 41.2 kcal/mol. The 1,2-dissociated 278 intermediate IM19 is 13.0 kcal/mol more stable than the 1,4- 279 dissociated intermediate IM20. The dissociated intermediate 280 HCOO of the hydroxyl adsorbates is about 56.2 kcal/mol more 281 stable than that of the carbonyl adsorbates when they adsorb on 282 the α -Al₂O₃(0001) surface. Through overcoming 2.4 and 1.7 283 kcal/mol energetic barriers, the HCOO group in IM19 and 284

Table 3. Geometric Changes after Formic Acid Adsorption/Dissociation at the α-Al₂O₃(0001) Surface

model	O_s – H_f^a	$Al_s - O_f^b$	$d_{\mathrm{Al-O}}^{c}$	$\delta(d_{\mathrm{Al-O}})$	$l_{\rm Al-O}^{d}$	$\delta(l_{\mathrm{Al-O}})$	A_{OAlO}^{e}	$\delta(A_{\rm OAlO})$	$\varphi(AlO_3)^f$	$\delta(arphi_{ ext{AlO3}})$
bulk			0.840		1.856		101.1		103.8	
surf			0.176		1.703		118.9		159.6	
R1	2.392	1.997	0.470	0.294	1.737	0.034	112.6	-6.3	131.2	-28.4
R2	2.101	2.001	0.456	0.280	1.733	0.030	112.1	-6.8	131.9	-27.7
R3	2.650	2.046	0.375	0.199	1.719	0.016	114.5	-4.4	139.3	-20.3
R4	1.983	2.103	0.421	0.245	1.721	0.018	115.2	-3.7	137.2	-22.4
IM1	1.005	1.836	0.554	0.378	1.869	0.166	104.8	-14.1	119.2	-40.4
IM2	1.031	1.860	0.593	0.417	1.766	0.063	105.9	-13.0	119.5	-40.1

^aThe bond distance of the outmost O_s atom of the surface and the H_f atom of formic acid, in angstroms. ^bThe bond distance of the outmost Al_s atom of the surface and the O_f atom of formic acid, in angstroms. ^cInterlayer spacing between the outmost Al_s and O_s layers, in angstroms, and δ represents the changes before and after formic acid adsorption on the clean surface. ^c $Al_s - O_s$ bond distance of the outmost layer on the surface, in angstroms. ^cBond angle of the $O_sAl_sO_s$ of the outmost layer on the surface, in angstroms. ^fDihedral angle of the Al_sO_{s3} tetrahedral structure, in degrees.

285 IM20 can rotate to form IM3 and IM8 with the adsorption 286 energies of 51.8 and 39.9 kcal/mol, respectively. The COOH 287 group in IM12 can rotate to IM13 and IM14 and in IM17 can 288 rotate to IM18. The COOH group in IM12 is the most stable 289 formation.

We study the vibrational properties of IM1 since it is the most stable intermediate. An IR spectrum of the isolated formic acid molecule is necessary. It is difficult to find the spectrum of an isolated formic acid molecule in the gas phase experimentally because it is mainly found as a dimer in the atmosphere. However, formic acid monomers have been studied in an argon matrix at low temperatures and low pressures. Therefore, theoretical calculations are useful to validate previous experimental work. As shown in Table 4, the calculated

Table 4. Calculated Vibrational Frequencies (in cm $^{-1}$) of the Gas-Phase Formic Acid and of the Adsorption Complexes at the 2 \times 2 Supercell Al $_2$ O $_3$ (0001) Surface at the BLYP/DND Level

modes ^a	formic acid	formic $acid^b$	R3	IM1	$IM1^c$
$\nu(O_fH_f)$	3512	3570	3497	-	-
$\nu(O_sH)$	-	-	-	3184	-
$ u(\mathrm{CH_f})$	2936	2944	3039	2885	2866
$\nu(C=O_f)$	1754	1776	1775	-	-
$ u_{\rm as}({\rm O_fCO_f})$	-	-	-	1608	1600
$\delta(\mathrm{CH_f})$	1356	-	1378	1373	1378
$\delta({ m O_f \! H})$	1268	1223	1205	-	-
$\delta(O_sH)$	-	-	-	1263	-
$\nu_{\rm s}({ m O_fCO_f})$	-	-	-	1266	1393
$\nu(C-O_f)$	1054	1105	976	-	-

"as = asymmetric mode, s = symmetric mode, ν = stretching, δ = bending in-plane. ^bIn the gas phase. ²⁸ **Experimental data in parentheses taken from ref 25.

 $_{209}$ frequencies for the isolated formic acid molecule are in good $_{300}$ agreement with the experimental values, an indication of the $_{301}$ good quality of the theoretical data. At the same time, we can $_{302}$ get the molecular frequencies of the adsorbed compounds as $_{303}$ R3 on the $\rm Al_2O_3$ surface that cannot be observed in $_{304}$ experiments.

The O_fH_f stretching frequencies, $\nu(O_fH_f)$, of the isolated 306 formic acid are calculated at 3512 cm⁻¹ which disappears in 307 IM1 because the H atom migrates from formic acid to the 2- O_s 308 atom. Instead, there is a new band at 3184 cm⁻¹, which 309 corresponds to the stretching vibrational mode of the O_s - H_f 310 bond. The IR spectrum of the physical adsorption compound

cannot be recorded experimentally, but we can see that from 311 the frequency calculations. There is a 15 cm⁻¹ low-frequency 312 shift of the $\nu(O_fH_f)$ in R3, owing to the direct interaction 313 between O_f and Al_s which weakens the O_f-H_f bond. The C-H_f 314 stretching vibration of IM1 has a red shift of 51 to 2885 cm⁻¹ 315 which is similar to the experiments of ref 25 of 2866 cm⁻¹ after 316 the dissociation at the surface; on the contrary, a high- 317 frequency shift of 113 cm⁻¹ is observed in the $\nu(CH_f)$ mode in 318 R3. The reaction of formic acid on the surface to give the 319 formate anion is suggested by the absence of the C=O_f stretch 320 mode at 1780 cm⁻¹ and the C-O_fH_f stretch mode at 1100 321 cm⁻¹. ⁴² The C=O_f stretch mode is at 1754 cm⁻¹ in formic acid 322 and at 1775 cm⁻¹ in R3, and the C-O_tH_f stretch mode is at 323 1054 cm⁻¹ in formic acid and at 976 cm⁻¹ in R3; however, they 324 disappear in IM1. Instead, we see the asymmetric and 325 symmetric stretch modes of O_fCO_f at 1608 and 1266 cm⁻¹ 326 in IM1, respectively. The wavenumber of the OfCOf 327 asymmetric stretch mode is very satisfied with the experimental 328 data 1600 cm⁻¹ which was studied by Tong et al., ²⁵ but that for 329 the symmetric mode is smaller than the experimental value of 330 1393 cm⁻¹. The bending in-plane $\delta(CH_f)$ mode has about a 25 331 cm⁻¹ high-frequency shift in both R3 and IM1 compared to the 332 free HCOOH molecule, and it is nearly the same as that in ref 333 25 of 1378 cm $^{-1}$. The bending in-plane $\delta(O_fH)$ disappears in 334 IM1, and it is replaced by $\delta(O_sH)$ with the vibrational 335 frequencies at 1263 cm⁻¹. The $\delta(O_fH)$ has a 63 cm⁻¹ low- 336 frequency shift to 1205 cm⁻¹ in R3 after being adsorbed on 337 surface. The IR spectra confirmed that the molecular 338 frequencies of the most stable intermediate IM1 are satisfied 339 with the experiments.

3.5. HCOOH Dehydrogenation/Dehydration at the 2 341 **x 2 Supercell.** Furthermore, we studied the dehydrogenation 342 and dehydration of HCOOH on the 2 \times 2 supercell α - 343 Al₂O₃(0001) surface. The optimized geometries and the 344 adsorption energies for all of the reactants, intermediates, 345 transition states, and products are shown in Figure S1 and 346 Table S (Supporting Information), respectively. All of the 347 energetic routes are shown in Figures S2-4(Supporting 348 Information). Figure 4 gives the predominant energetic routes. 349 f4 For the predominant dehydrogenation route, if the free formic 350 acid molecule approaches the α -Al₂O₃(0001) surface without 351 any constraints through the carboxyl O atom and hydroxyl H 352 atom, the O-H bond of formic acid dissociates through an 353 energetic barrier free process to form IM2 with binding energy 354 of 54.7 kcal/mol first. Then the formate(HCOO) overcomes 355 an energetic barrier of 19.1 kcal/mol to form IM8 with the 356

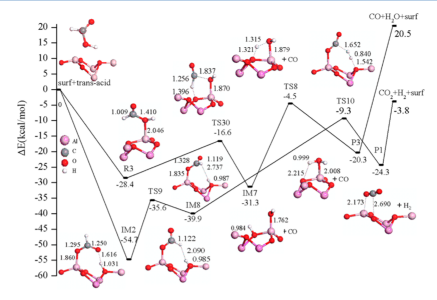


Figure 4. Predominant energetic routes for the dehydrogenation and dehydration reactions of formic acid on the α -Al₂O₃(0001) surface.

Table 5. Energetic Barrier of Dehydration/Dehydrogenation of Formic Acid (kcal/mol)

method	dehydrogenation	dehydration
experiment ⁴³	65-68	62-65
experiment ⁴⁴	48.5	60.5
$PMP4/6-311++G(d,p)//UMP2/6-311G(d,p)^{45}$	65.2	63.0
$B3LYP/6-311+G(2d,p)//HF/3-21G(d,p)^{46}$	67.8	66.8
MP2/6-311G//HF/6-311G ⁴⁷	77.6	67.1
$B3LYP/6-311G(2d,p)//MP2/6-311G(2d,p)^{48}$	73.3	56.0
in the gas phase in this work	65.3	62.1
on the α -Al ₂ O ₃ (0001) surface in this work	30.6	26.8

357 binding energy of 39.9 kcal/mol. In IM8, the CH hydrogen and 358 hydroxyl H are in the distance of 2.090 Å.

Then, IM8 overcomes the 30.6 kcal/mol energetic barrier to 359 360 break the C-H bond to produce H₂ away from the surface and CO₂ adsorbing on the surface with the binding energy of 24.3 362 kcal/mol through TS10. In TS10, the C-H and O_s-H bond lengths increase from 1.122 and 0.985 Å to 1.652 and 1.542 Å, respectively, and the distance between the two H atoms decreases from 2.090 to 0.840 Å. The O and C atom of CO₂ is 365 366 in the distance of 2.173 and 2.690 Å with the surface Al and O atom, respectively. The energy of free CO2 and H2 molecules compared with that of the free HCOOH molecule is -3.8 kcal/ mol; that is, the dehydrogenation reaction of formic acid on the α -Al₂O₃(0001) surface is slightly exothermic by 3.8 kcal/mol. The C-H bond breaking step is the rate-determining step of the dehydrogenation reaction of formic acid on the α -373 Al₂O₃(0001) surface with the energetic barrier of 30.6 kcal/ 374 mol. Other experimental and theoretical energetic barriers for the dehydrogenation and dehydration reactions of formic acid are shown in Table 5. The energetic barrier of the dehydrogenation reaction of formic acid in the gas phase with our model is 65.3 kcal/mol in this work, and it is the same as the experiment⁴³ of 65-68 kcal/mol; however, it is different from the experimental⁴⁴ value of of 48.5 kcal/mol which is tested at the low voltage of 0.03-0.3 atm, and it is consistent 382 with the calculated value of 65.2 kcal/mol at the level of 383 PMP4/6-311++G(d,p)//UMP2/6-311G(d,p)⁴⁵ and of 67.8 384 kcal/mol at the level of B3LYP/6-311+G(2d,p)//HF/3-385 21G(d,p). The values at the level of MP2/6-311G//HF/6- $386 \ 311G^{47}$ and $B3LYP/6-311G(2d,p)//MP2/6-311G(2d,p)^{48}$ are

77.6 and 73.3 kcal/mol, respectively. It can be seen that the $_{387}$ dehydrogenation of formic acid on the $\alpha\text{-Al}_2\text{O}_3(0001)$ surface $_{388}$ can be enhanced considerably with the energetic barrier $_{389}$ decreasing from 65.3 to 30.6 kcal/mol. $_{390}$

The dehydration reaction path starts from R3 whose C-O 391 and C-H bonds break simultaneously through the five- 392 membered ring transition state TS30 with the energetic barrier 393 of 11.8 kcal/mol to form IM7 with binding energy of 31.3 kcal/ 394 mol. In TS30, the C-H and C-O bond lengths increase from 395 1.009 and 1.410 Å to 1.256 and 1.837 Å, respectively, and the 396 distance between CH hydrogen and the 2-O_s is 1.396 Å. In 397 IM17, the CO free molecule runs away from the α - 398 Al₂O₃(0001) surface, and the dissociated OH and H bond 399 with the surface Al atom and 2-O_s atom with the bond length of 400 1.762 and 0.984 Å, respectively. Then the H atom transfers 401 from the 2-O_s atom to the hydroxyl O atom through the four- 402 membered ring transition state TS8 with an energetic barrier of 403 26.8 kcal/mol to form the adsorbed H₂O molecule with the 404 binding energy of 20.3 kcal/mol. In TS8, the migrating H atom 405 is almost in the middle of the two O atoms with the distances 406 of O-H bonds 1.321 and 1.315 Å, respectively. The bond 407 length between the O atom of the adsorbed H₂O molecule and 408 the surface Al atom is 2.008 Å. The energy of free CO and H_2O 409 molecules compared with the free HCOOH molecule is 20.5 410 kcal/mol; that is, the dehydration reaction of formic acid on the 411 α -Al₂O₃(0001) surface is endothermic by 20.5 kcal/mol. The 412 rate-determining step of the dehydration reaction of formic acid 413 on the Al₂O₃ surface is the forming of the O-H bond in the 414 water molecule with the energetic barrier of 26.8 kcal/mol. The 415 energetic barrier of the dehydration reaction of formic acid in 416 417 the gas phase is 62.1 kcal/mol with our model in this work, and 418 it is nearly the same as the experimental values of 62–65 kcal/419 mol⁴³ and of 60.5 kcal/mol⁴⁴ and is consistent with other 420 calculations. 45–48 It can be seen that the dehydration of formic 421 acid on the α -Al₂O₃(0001) can be enhanced considerably with 422 the energetic barrier decreasing from 62.1 to 26.8 kcal/mol.

3.6. Two HCOOH Molecule Adsorption at the 2 \times 2 Supercell. Since the surface Al atom can bond with more than 425 one O atom, it is interesting to investigate how the second 426 HCOOH or more HCOOH molecules react with the α -427 Al₂O₃(0001) surface. Starting from the most stable structure of 428 the molecular adsorbed compound R1, one formic acid 429 molecule attacks the surface Al atom to form compound R5 430 with a weak bond, as shown in Figure 5. The Al₈-O₆ bond is

Figure 5. Adsorption intermediates formed by addition of one/two HCOOH molecules to the α -Al₂O₃(0001) surface.

431 relatively long, 2.186 and 2.227 Å, compared with the bond 432 length of 1.997 Å in R1. The C–O bond of formic acid 433 increases from 1.338 to 1.352 Å and 1.362 Å, respectively. The 434 second formic acid can draw up the Al atom since the distance 435 between the first layer Al atoms and the first layer O atoms is 436 0.7 Å in R5 compared with that of 0.4 Å in R1. Meanwhile, the 437 O_s –Al $_s$ – O_f angle decreased from 97.1° to 90.2° and 84.6°, 438 respectively, and the O_f –Al $_s$ – O_f bond angle between the two 439 HCOOH molecules is 68.6°. The binding energy of R5 is 42.7 440 kcal/mol, which is slightly larger than that of R1, 32.2 kcal/mol. 441 The two formic acid molecules can absorb stably on the same 442 surface Al atom through carboxyl O atoms, but they cannot 443 bond with the same Al atom after dissociating into HCOO and 444 H. It concludes that each surface Al atom can just bond with 445 one dissociated HCOOH molecule.

446 **3.7. Fully Hydroxylated 2** \times **2 Supercell Surface.**447 Hydroxylation of the metal oxide surface is a general issue, and
448 it is important to analyze the influence of hydroxyl groups on
449 the reaction of formic acid at the α -Al₂O₃(0001) surface.

The coordinatively unsaturated surface Al ions provide 451 strong Lewis acid (electron-acceptor) sites for H₂O adsorp-452 tion. 49 It is believed that water dissociates on the Al-terminated 453 Al₂O₃(0001) surface, and the mechanism of water at the 454 Al₂O₃(0001) surface has been calculated extensively using 455 various cluster/lattice models and first-principles method-456 ologies. 50-56 Wittbrodt et al. 57 employed three cluster models 457 to investigate the interaction of water with the (0001) surface of α -Al₂O₃ at the B3LYP/6-31+G* level of theory. It was found that there are two types of dissociative reactions; namely, 1-2(proton transfer to the nearest crystal oxygen) and 1-4 (proton transfer to the second nearest oxygen) addition of 462 water take place. Wang et al. 58 studied a single water molecular 463 dissociation model on the α -Al₂O₃(0001) surface at the BLYP/ $464 \text{ TNP}/3 \times 3 \times 3 \text{ k-points level of theory and found that in the } 2$ 465 × 2 supercell model the hydroxylation prefers 1,2-dissociation 466 which is exothermic by 37.3 kcal/mol. The author also 467 calculated the multiple layer adsorption and confirmed that

the outmost Al atom of the (0001) surface is hydroxylated by 468 one H_2O molecule and inert to further hydroxylation by water. 469

Accordingly, the fully hydroxylated $Al_2O_3(0001)$ 2 \times 2 470 supercell surface with the outmost four Al atoms being 471 hydroxylated by $4H_2O$ molecules through the 1,2-migration 472 mode was studied in this work, and the geometry of the surface 473 is shown in Figure 6. The dissociated OH bonds with Al atom, 474 f6

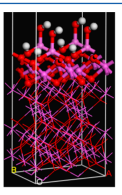


Figure 6. Optimized slab model of the hydroxylated α -Al₂O₃(0001) 2 \times 2 supercell surface.

and the H atom bonds with the surface O atom, forming a new 475 layer. The geometric and energetic changes from the clean to 476 the fully hydroxylated 2 \times 2 supercell $\rm Al_2O_3(0001)$ surface are 477 listed in Table 6. It is obvious that the dissociation adsorption 478 to of water draws up the surface Al atoms. The distance of the 479 surface Al to the first oxygen plane is increased by 0.408 Å. The 480 bond length of Al–O of the outmost layer increases to 1.862 Å 481 which is a little longer than that of the bulk. The bond angle of 482 OAlO and the dihedral angle of the AlO3 tetrahedral decrease 483 14.1° and 48 , respectively, after the surface is hydroxylated 484 fully by the water molecule.

The adsorption energy for $4H_2O$ molecules dissociation at 486 the α -Al $_2O_3(0001)$ 2 \times 2 supercell surface is -166.7 kcal/mol, 487 namely, -41.4 kcal/mol for each H_2O molecule dissociation on 488 the surface. The calculated energy in the work compares well to 489 the experimental values of 25–41 kcal/mol. 59

3.8. HCOOH Adsorption at the Fully Hydroxylated 2 \times 491 2 Supercell. The optimized geometries of the formic acid 492 molecule at the hydroxylated 2×2 supercell Al₂O₃(0001) 493 surface are shown in Figure 7. The O and H atoms of formic 494 f7 acid interact with the hydroxylated O-H hydrogen and Al- 495 OH oxygen of the surface, respectively, in the physical 496 adsorption compounds. When formic acid interacts with the 497 surface through hydroxyl O and H atoms, it forms R1 with the 498 binding energy of 22.7 kcal/mol. In the molecular adsorption 499 compounds R2 (O atom reacts with the 4-H of surface OH) 500 and R3 (O atom reacts with the 2-H of surface OH), formic 501 acid interacts with the surface through the carboxyl oxygen and 502 the OH hydrogen with the binding energy 21.4 and 25.3 kcal/ 503 mol, respectively. When the main interactions involve the 504 hydroxyl O atom and CH hydrogen atom, it forms R4 with the 505 binding energy of 11.1 kcal/mol. When it invovles the carboxyl 506 O atom and CH hydrogen atom, it forms R5 with the binding 507 energy of 19.6 kcal/mol.

Considering the O–H bond dissociation of formic acid into 509 P1 and P2 with the binding energy of 24.5 and 27.4 kcal/mol, 510 the product HCOO and H_2O molecule react with the same 511 surface Al atom through an oxygen atom.

Table 6. Geometric and Energetic Changes from the Clean to the Hydroxylated 2×2 Supercell α -Al₂O₃(0001) Surface

model	$d_{\mathrm{Al-O}}$	$\delta(d_{\mathrm{Al-O}})$	$l_{\rm Al-O}$	$\delta(l_{\mathrm{Al-O}})$	A_{OAlO}	$\delta(A_{ m OAlO})$	$\varphi(AlO_3)$	$\delta(arphi_{ ext{AlO3}})$	$E_{ m ads}$
bulk	0.840		1.856		101.1		103.8		
clean surface	0.176		1.703		118.9		159.6		
hydroxylation	0.584	0.408	1.862	0.159	104.8	-14.1	111.2	-48.4	-166.7

Figure 7. Optimized geometries and adsorption energies of the formic acid molecule at the fully hydroxylated α -Al₂O₃(0001) 2 × 2 supercell surface. Bond distances are in angstroms, and energies are in kcal/mol.

When it starts from R1, the product forms through the four-514 membered ring transition state TS1 with the energy barrier of 515 29.3 kcal/mol. If the reactors are R2 and R3, they overcome a 516 19.6 and 14.9 kcal/mol energy barrier through a six-membered 517 ring transition state, respectively.

This can conclude that the prominent route for HCOOH dissociation at the fully hydroxylated α -Al₂O₃(0001) 2 × 2 supercell surface is starting from R3 that through the 1,2-site adsorbs on the surface to form the HOCO–Al–OH₂ covered surface. The energy barrier is 14.9 kcal/mol, and the binding energy of the product is 24.5 kcal/mol. So it is easier for formic acid to dissociate at the dry α -Al₂O₃(0001) surface which is energy barrier free.

3.8. Lauric Acid Dissociation at the 2 \times 2 Supercell. In order to study the superhydrophobic mechanism of the α -528 Al_2O_3 surface modified with lauric acid, the absorption/529 dissociation of formic acid at the α -Al $_2\text{O}_3(0001)$ 2 \times 2 530 supercell surface was studied as the model. The optimized 531 geometries are shown in Figure 8. This confirms that the most 532 stable intermediate of the HCOOH molecule is the 1,2-533 dissociated intermediate involving the carboxyl O and hydroxyl 534 H with the binding energy of 59.5 kcal/mol. In order to testify 535 the dissociation configuration of lauric acid, the geometries of

the free $C_{11}H_{23}COOH$ molecule and the dissociated inter- 536 mediate were studied. It was found that the lauric acid molecule 537 dissociated into $C_{11}H_{23}OCO$ - and HO-covered surface, and it 538 was energy barrier free with the binding energy of 60.7 kcal/ 539 mol. It can be seen that the superhydrophobic alkylation group 540 spread upward which contributes the superhydrophobicity of 541 the alumina surface.

4. CONCLUSIONS

The formic acid molecule is adsorbed on the outer Al sites of 543 the α -Al $_2$ O $_3$ (0001) surface by two approaches, namely, 544 carbonyl adsorption and hydroxyl absorption. The molecular 545 adsorption compound involving the carboxyl O atom and CH 546 hydrogen with the surface 4-O $_s$ is the most stable one with the 547 binding energy of 33.3 kcal/mol. The chemical dissociation via 548 either 1,2- or 1,4-hydrogen migration is a free energetic barrier 549 when it involves the carboxyl O and hydroxyl H atoms, and the 550 1,2-dissociation produces the most stable intermediate with the 551 binding energy of 59.5 kcal/mol. When the modifier lauric acid 552 dissociates on the alumina surface, it forms the stable 553 $C_{11}H_{23}OCO$ - and HO-covered surfaces with the binding 554 energy of 60.7 kcal/mol.

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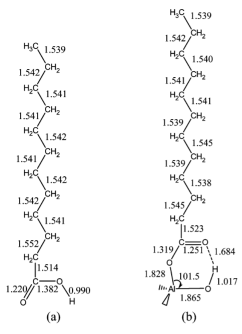


Figure 8. Optimized geometries of the free lauric acid molecule (a) and the 1,2-dissociated intermediate (b) at the α -Al₂O₃(0001) 2 × 2 supercell surface.

As for the vibrational frequencies, we can calculate those of the dissociated intermediates and the molecular adsorption compounds which cannot be observed in experiments on the α -ss9 Al₂O₃(0001) surface. The O_f—H_f and C=O_f stretching wibrations disappear in the dissociated intermediate compared to the free formic acid at 1754 cm $^{-1}$ and molecular adsorbed compound at 1775 cm $^{-1}$, but two new vibrational modes are attributed as the $\nu_{\rm as}({\rm OCO})$ and $\nu_{\rm s}({\rm OCO})$ at 1608 and 1266 cm $^{-1}$, respectively. The bending in-plane $\delta({\rm C-H_f})$ in the molecular compound at 1378 cm $^{-1}$ is the same as that in the dissociated intermediates at 1373 cm $^{-1}$. These values are consistent with experiments and confirm that the most stable intermediate is the 1,2-dissociated intermediate involving the carboxyl O and hydroxyl H.

Dehydrogenation and dehydration routes have been revealed theoretically on the α -Al₂O₃(0001) surfaces. The predominant reaction path to form CO₂ + H₂ proceeds via the stepwise mechanisms with the energetic barrier of 30.6 kcal/mol. In CO + H₂O involves the concerted CO and CH bond breaking with the energetic barrier of 26.8 kcal/mol. In comparison with the gaseous unimolecular decomposition of formic acid, the energetic barrier of the dehydrogenation and dehydration reaction can be decreased 34.7 and 35.3 kcal/mol by the α -Al₂O₃(0001) surface.

Furthermore, two formic acid molecules are considered to be s81 stable to adsorb physically on the same surface Al atom as s82 indicated by the calculated binding energy of 42.7 kcal/mol, but s83 just one dissociated HCOOH molecule can bond with each s84 surface Al atom.

The influence of the hydroxylation on the dissociation of formic acid at the α -Al₂O₃(0001) surface is analyzed. It finds that the 1,2-dissociation involving the carboxyl O atom and OH hydrogen has the lowest energy barrier of 14.9 kcal/mol to form HOCO- and H₂O-covered surfaces.

The dissociated configuration of lauric acid at the α 591 Al₂O₃(0001) surface is tested. It can dissociated into

 $C_{11}H_{23}OCO$ - and HO-covered surfaces with the binding s92 energy of 60.7 kcal/mol and free energy barrier.

ASSOCIATED CONTENT

S Supporting Information

Table S and Figures S1—S4. This material is available free of 596 charge via the Internet at http://pubs.acs.org.

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Notes 602

The authors declare no competing financial interest.

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