1	Hydrogen donation of bio-acids over transition metal			
2	facets: A Density Functional Theory study			
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11 Abstract

12 This research aims to evaluate and systematic compare two bio-based acidic hydrogen 13 donors as alternative hydrogen source for hydrodeoxygenation (HDO) of bio-oil from fast 14 pyrolysis of biomass. We investigated the hydrogen donation performance of acetic acid 15 (AcOH) and formic acid (FA) over the most stable facet of transition metal in comparison 16 with the dissociation of H₂, using Density Functional Theory (DFT) modelling. It was 17 found that AcOH adsorbed more strongly on Mo (110) than the other base transition 18 metals investigated. The binding of both acid molecules and Mo were dominated by 19 electrons migration from the molecule to the metal, while electron enrichment was also 20 observed for FA during the adsorption. The bonds strength for both acids were weakened 21 by Mo, therefore, facilitating their decomposition. The hydroxyl hydrogen and one methyl 22 hydrogen of AcOH could be readily cleaved with low energy barriers, but the methylene 23 hydrogen was found to be difficult to cleave when the carbon was unsaturated. Within

24 the FA molecule, the hydroxyl hydrogen would be donated more readily than the 25 hydrogen from aldehyde group. H_2 dissociated during adsorption on the Mo (110) facet 26 with negligible energy barrier. Within the three sources of hydrogen (acetic acid, formic 27 acid and H_2 gas), H_2 gas has the advantage of easy dissociation when compared to the 28 hydrogen generated from biomass-derived acids (bio-acids). However, the large 29 enthalpy change resulted from the exothermic decomposition of the bio-acid will probably 30 facilitate the activation and further migration of the single H atom for donation over the 31 metal facet compared to H₂ gas. AcOH exhibited a greater potential than FA as a 32 hydrogen donor over Mo (110) due to the release of more H atoms with low energy 33 barriers. The modelling results also suggested that AcOH is a promising alternative hydrogen source to replace H_2 gas over the facet of Mo (110). 34

Abbreviations: Acetic Acid, AcOH; Formic Acid, FA; Aromatic Hydrocarbons, AHs;
Electronic density difference, EDD

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Key words: Hydrogen donor compounds; Catalytic decomposition; Hydrodeoxygenation;
 DFT modelling; Transition metals

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41 **1. Introduction**

Nowadays biofuel is an attractive alternative fuel because it can lower our dependency on conventional fossil fuels, achieve "zero" net carbon emissions, and tackle the challenge of global warming [1]. Thermochemical conversion of waste biomass via catalytic fast pyrolysis offers a promising route to biofuels. However, the direct use of bio-oil is hindered by its undesirable physicochemical properties such as low calorific value due to its high oxygen content and water content, high viscosity and low stability owing to its high acid content. This explains why upgrading is being so intensively studied.

49 Hydrodeoxygenation (HDO) is considered an effective method for upgrading bio-oil to 50 more valuable and useful liquid products [2]. Conventionally, molecular H₂ gas is used 51 in industrial HDO processes, because of its dissociation on the surface of the catalysts, 52 thus providing high reactivity. However, there are some drawbacks associated with the 53 application of molecular H₂ gas such as the high process costs due to high-pressure 54 reactors, concerns over process safety regarding H_2 storage and utilisation, the 55 requirement to recover and recycle hydrogen. Additionally, using pure H_2 during HDO 56 may lead to the saturation of aromatic rings, producing undesired cycloalkanes instead 57 of aromatic hydrocarbons. Therefore, alternative sustainable hydrogen sources are 58 urgently sought-after to address the challenges of developing an efficient and cost effective HDO process. 59

60 Much effort has been devoted to finding the alternative hydrogen sources, and it has 61 been reported that hydrogen can be produced in-situ via various reactions [2,3]. 62 Commonly reported alternative hydrogen donors include primary and secondary 63 alcohols such as methanol, ethanol and isopropyl alcohol (IPA), carboxylic acids 64 including acetic acid (AcOH) and formic acid (FA), and aromatic hydrocarbons (AHs) 65 including tetralin and decalin [4,5]. These hydrogen-rich compounds can supply 66 hydrogen for the HDO reaction through self-decomposition or dehydrogenation [1]. For 67 example, Wang et al. [6,7] reported the deoxygenation of phenol over Ni/Al_2O_3 and Ru 68 based catalysts using methanol and formic acid as hydrogen donor compounds. Guo et 69 al. [8] explored the HDO of p-cresol over a Ru based catalyst with IPA as the hydrogen 70 donor, achieving 98.5% conversion and 84.0% yield of toluene. IPA was also found to 71 be effective in humins conversion [9]. Vasiliadou and Lemonidou reported the HDO of 72 glycerol at 220 – 250 °C. The highest 1,2-propanediol yield of 53% was obtained using 73 ethanol as the source of hydrogen [10]. Pajak et al. [11] investigated the hydrogen 74 transfer from tetralin and decalin to coal derived tars, revealing hydrogen transfer from decalin was two times greater than that of tetralin. Similar conclusions were drawn by other researchers when decalin and tetralin were used for bio-oil upgrading through hydrogen transfer deoxygenation [12,13]. In-situ H₂ gas production can also take place through redox reactions. For example, water-gas shift reaction (CO + H₂O \Rightarrow CO₂ + H₂) may occur within the HDO process to supply hydrogen [14].

The mechanism of various hydrogen transfer reactions for HDO have been extensively investigated [15–18]. The adsorption and dissociation of molecule hydrogen over various transition metals were reported by Pozzo et al., indicating Ni was the most effective metal in such a process [19]. Experimental and modelling investigation performed by Grilc and Likozar and other researchers revealed that bi-functional Ni-Mo based catalysts are promising for HDO with molecular H₂ [20,21]. Numerical models have been reported as well [22–24] to describe the mechanism and kinetics of catalytic HDO reactions.

87 DFT based studies regarding decomposition of bio-alcohols have been also intensively 88 reported; the decomposition mechanism of methanol over Co(0001) and Co(111), 89 Pd(111), Pt(111), and Ni(111) facets has been investigated, predicting all the C-H bond 90 and C-O bond would cleave prior to the CO bond in these scenarios [25,26]. Alcalá et al. 91 reported the decomposition mechanism of ethanol, predicting that demethylation 92 reaction would happen much faster than dehydroxylation over Pt(111), and methane and 93 CO were more likely to be produced [27,28]. In addition, detailed mechanisms for ethanol 94 decomposition over various transition metals, propanol decomposition over Ni-Fe and 95 Ni-Cu bimetallic facet, and butanol decomposition over Pd(111) facet have been widely 96 reported [28–31].

97 There are limited reports on the DFT modelled decomposition of AHs (as H donors). Kim 98 et al. elucidated the dehydrogenation mechanism of decalin to tetralin and further to 99 naphthalene on the (111) facet of Pd and Pt respectively, revealing that according to the modelling, Pd would outperform Pt by lowering the energy barrier of tetralin conversion
to naphthalene by 0.31 eV [32].

102 Regarding the bio-acids, the decomposition of FA over Cu (111) facet has been analysed 103 by DFT, predicting the most favourable reaction pathway was HCOOH \rightarrow HCO \rightarrow CO 104 [33]. The decomposition of acetic acid over a variety of catalysts were also reported 105 [34,35]. Xinbao et al. investigated the decomposition mechanism of AcOH over Co (111) 106 stepped facet, and revealed that the reaction pathway follows $CH_3COOH \rightarrow CH_3CO \rightarrow$ 107 $CH_2CO \rightarrow CH_2 \rightarrow CH$ [36]. Although, there are other different reaction mechanisms for 108 AcOH decomposition proposed in the literature [37–39]. The detailed description of the 109 decomposition of bio-acids over different catalysts, especially over base transition metals, 110 are still unclear and scarce. Moreover, previous studies have reported the performance 111 of various hydrogen donors, however most of them cannot be compared because 112 different catalysts were used. There are very few reports on the evaluation of the 113 reactivity of different hydrogen donor compounds over one selected catalyst and their 114 capacity for HDO reactions.

115 Hence, this research is aimed at the evaluation and systematic comparison between two 116 bio-based acidic hydrogen donors, since they abundantly exist in the primary bio-oil. The 117 decomposition of these carboxylic acids leads to in-situ hydrogen formation, which 118 subsequently can take part in HDO reactions. With respect to catalyst selection, the 119 performance of noble metal based catalysts e.g. Pt and Ru, have been widely studied, 120 while not enough attention have been paid to the base transition metals. Therefore, this 121 study has sought to predict the decomposition of AcOH with a focus on the cleavage of 122 hydrogen related bonds over various transition metals. Ten transition metals (Ni, Mo, Fe, 123 Co, Pt, Rh, Ru, Zn, Cu, and Pd) are compared regarding their binding energy in 124 adsorbing AcOH and the one with strong binding was selected. Then, AcOH and FA are 125 compared with H₂ in terms of their adsorption process and hydrogen donation 126 performance over the most stable facet of the selected catalyst. Electronic density 127 difference (EDD) of the hydrogen donor compounds was analysed to investigate the 128 interface interactions. Bond lengths of the donor compounds were also tracked 129 throughout their decomposition to evaluate the impact of metal catalyst.

130 **2. Computational details**

The first-principle density functional theory plus dispersion (DFT-D) calculations were 131 132 implemented in the Cambridge Sequential Total Energy Package (CASTEP) module 133 available in Materials Studio 2017 R2 from BIOVIA [40,41]. The generalized gradient 134 corrected approximation (GGA) [42] treated by the Perdew-Burke-Ernzerhof (PBE) 135 exchange-correlation potential with long-range dispersion correction via Grimme's 136 scheme was used to calculate the exchange-correlation energy [43]. The On-the-fly 137 generated (OTFG) ultrasoft pseudopotential was employed as the scheme in the 138 representation of reciprocal space for all the elements [44,45]. The plane-wave cut-off 139 energy was set to 500 eV for all the calculations based on its independence test (Fig. 140 S1(a)). The Brillouin zone was sampled using a 2×2×1 Monkhorst-Pack k-point (spacing 141 of 0.04 Å⁻¹) with a smearing of 0.1 eV, based on its independence test (Fig. S1(b)). The self-consistent field (SCF) tolerance was set to 10⁻⁶ eV/atom. All the modelling was 142 performed with a convergence threshold of 10⁻⁵ eV/atom on energy, 0.03 eV/Å on 143 maximum force, and 10⁻³ Å on the maximum displacement. No symmetry constraints 144 145 were used for any modelling. The impact of zero point energy (ZPE) was evaluated for 146 the adsorption of AcOH and FA. It was found that the net ZPE corrections (difference 147 between ZPE of the systems before and after the adsorption) were less than 0.01eV, so 148 that the ZPE correction was not included in the calculations of this study. The 149 computational method is believed to give high precision results, according to the 150 validation of lattice constant for each metal element lattice in this study; the variations between computational and experimental values are less than 0.10 Å for most systems(full comparison is shown in Table S1).

153 The most stable facets of the ten metals were simulated; facet (1 1 1) was simulated for 154 face-centred cubic (FCC) metals including Ni, Cu, Pd, Pt, and Rh, and facet (1 1 0) was 155 simulated for body-centred cubic (BCC) metals of Mo, Fe. For hexagonal close-packed 156 (HCP) metals including Co, Zn, and Ru, facet (0 0 1) was simulated [28,46]. All the metal 157 facet models were created from the optimized metal lattices, and a four-layer slab of P 158 (5×5) super-cell was used with the adsorbate coverage of 1/25ML (Fig. S2). 15Å vacuum 159 region was created above the metal facet. The transition state (TS) was completely 160 determined by the LST/QST method, and the TSs for the dominated reaction steps were 161 confirmed by the unique imaginary frequency. Mulliken charges were assigned to each 162 bond to address the bond order [47]. The adsorption energy E_{ad} was determined by Eq.1, 163 where E_{catalyst}, E_{adsorbate} and E_{adsorbate/catalyst} are the total energies of clean metal facet, free 164 adsorbate molecule and metal facet with adsorbed molecule respectively. The energy 165 barriers of reactions E_{barrier} were determined by the difference between the transition 166 state and reactant energies, as shown in Eq.2, where Etransition state and Ereactant are the total 167 energies of the transition state and reactant of a reaction.

168
$$E_{ad} = E_{catalyst} + E_{adsorbate} - E_{adsorbate/catalyst}$$
 Eq.1

$$E_{barrier} = E_{transition \ state} - E_{reactant}$$
 Eq.2

170 EDD was determined by Eq.3.

171
$$\Delta \rho = \rho_{adsorbate@Mo(110)} - (\rho_{adsorbate} + \rho_{Mo(110)}) \qquad Eq.3$$

where $\rho_{adsorbate@Mo(110)}$ is the electron density of the total adsorbate + Mo(110) system, and $\rho_{adsorbate}$ and $\rho_{Mo(110)}$ are the unperturbed electron densities of the adsorbate and the Mo(110) facet, respectively.

- 175 Geometry optimization was implemented to every model before energy was calculated.
- 176 Energy of all the geometries was calculated at 0K in the DFT investigation.

177 **3. Results and discussion**

178 3.1 Metal catalyst selection based on AcOH adsorption

179 Most stable facets of ten transition metals were established to identify the metal catalyst 180 that can adsorb AcOH more strongly than the others. The adsorption modelling was 181 carried out by placing AcOH molecule vertically, and binding onto the top site of each 182 metal facet with the carbonyl oxygen atom. More details of the stable adsorption models 183 are shown in the Fig. S2. The adsorption energies are summarized in Table 1, along with 184 relevant geometry parameters (bond length of O-metal, which refers to the distance 185 between the centres of the bounded oxygen atom and the nearest metallic atom, and 186 the corresponding angle of C-O-metal) of the most stable geometry.

187 Table 1. Adsorption energies and geometrical parameters of AcOH on common facets

Metal facets	Bond length (Å)	∠C-O-metal (degree)	Adsorption energy (eV)
Ru (0001)	2.16	143.22	0.91
Mo (110)	2.10	170.16	0.88
Ni (111)	1.93	169.33	0.71
Pd (111)	2.14	141.28	0.70
Fe (110)	1.97	170.92	0.67
Co (0001)	1.95	176.82	0.66

188 of transition metals

Rh (111)	2.21	175.10	0.60
Pt (111)	2.32	137.75	0.58
Cu (111)	2.15	148.03	0.47
Zn (0001)	2.72	127.95	0.30

189 Note: Considering the effects of adsorption sites and compounds configurations on adsorption energy, more 190 comprehensive adsorption energy of AcOH onto the top three metal facets of Ru, Mo and Ni in different 191 adsorption sites and configurations are calculated and shown in Table 2 and Table. S2.

The adsorption energy of AcOH onto all of the modelled transition metals facets were less than 1.00 eV. The highest adsorption energy of 0.91 eV and 0.88 eV associated with AcOH adsorption onto Ru (0001) and Mo (110) respectively, followed by 0.71 eV for Ni (111) and 0.70 eV for Pd (111). Adsorption of AcOH on Zn (0001) exhibited the lowest adsorption energy of 0.3 eV. The bonds generated between metal facets and AcOH molecule ranged from 1.93 to 2.72 Å, and the angles generated between C-O bond and O-metal bond ranged between 127.95° and 176.10° for all cases.

199 In the perspective of thermodynamic, the high adsorption energy for Ru, Mo, Ni, Pd 200 implies that chemisorption happens with a strong binding between AcOH molecule and 201 the metals facets. These four metals are commonly known as the catalysts being 202 adopted for the HDO using H₂ gas and for the decomposition of organic compounds with 203 good performance [5,48-50]. Specifically for acids decomposition, Ru, Ni and Pd have 204 been widely reported based on experimental investigation as well as DFT modelling 205 [6,7,34,39,51]. Ru was found guite active in HDO with formic acid as hydrogen donor [7]. 206 Ni was reported to exert high activity in HDO with H₂ gas but poor performance with 207 formic acid as hydrogen donor [6]. Pd is also active but has the potential to lead to the 208 direct formation of CH₄ instead of hydrogen donation [34]. Besides, Ru and Pd are noble

- 209 metals, which would lead to high cost. In comparison, Mo as catalyst has been less210 investigated thus it is the focus for the further investigations in this study.
- 211 3.2 AcOH as hydrogen donor on Mo (110) facet
- 212 The adsorption of AcOH onto Mo (110) facet is investigated in terms of the preferred
- absorption sites and configuration. Four probable adsorption sites are considered for the
- adsorption modelling, including top site (Top), short bridge site (SB), long bridge site (LB)
- and hollow site, as shown in Fig. 1.



- 216
- Fig. 1. The probable adsorption sites on the facet of Mo (110)

The adsorption of AcOH was simulated over four different adsorption sites, and four configurations for AcOH were compared. The structures of each stable adsorption are shown in Table 2.

Table 2. Adsorption energy of AcOH, FA and H₂ onto Mo (110) at different sites and configurations

Adsorption Energy			1162	5.511
of AcOH (eV)				
	① Ο α- Μο	 Ο_β-Mo 	③ Parallel	④ C _α -Mo
Тор	0.88	0.70	0.28*	0.22*
Short Bridge (SB)	1.24	-	-	-
Long Bridge (LB)	0.28*	-	-	-
Hollow	0.95	-	-	-
Adsorption Energy of FA (eV)				
	 O_α-Mo 	② C-Mo**	③ O _β -Mo	(4) Parallel
Тор	1.15	0.75	0.51	0.18*
Short Bridge (SB)	0.15*	-	-	-
Long Bridge (LB)	0.16*	-	-	-
Hollow	0.12*	-	-	-
Adsorption Energy of H ₂ (eV)				
	1) Parallel	2 Vertical		
Тор	0.57	0.56	-	-
Hollow	1.64	-	-	-

223 * Only stable physical adsorption observed

224 ** Transformed binding from C-Mo to O-Mo (as shown) after geometry optimization

Configuration ① led to the highest AcOH adsorption energy of 0.88 eV compared to other configurations, followed by configuration ④, for 0.70 eV. While configuration ② and ③ resulted in lower adsorption energies of 0.28 eV and 0.22 eV respectively.
Regarding different adsorption sites based on configuration ①, AcOH was able to develop strong bindings on to the top, SB and hollow site of Mo facet, resulting in the

230 adsorption energy of 0.88 eV, 1.24 eV and 0.95 eV respectively. The adsorption of AcOH 231 onto LB site gave rise to the adsorption energy of 0.28 eV.

232 The results reveal that AcOH molecule most likely will adsorb on the SB site of Mo (110) 233 facet through the binding between O^{α} and Mo facet, leading to the highest adsorption 234 energy. Carbon and hydrogen in the molecule are found hardly to form strong bond with 235 the Mo facet independently. The adsorption onto the positions of Top (1) and SB (2), and 236 hollow are likely to be chemical adsorption, because of the large adsorption energy. To 237 confirm this, electrons migration between the interfaces was investigated by the 238 calculation of EDD, as shown in Fig. 2.



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Fig. 2. Colour mapping of electron density difference (threshold value: +/- 0.05 electrons/Å³) for AcOH adsorption configurations of (a) Top ①, (b) Top ②, (c) SB, and (d) hollow. (Loss of electrons is indicated in blue, while electron enrichment is indicated

in red)

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247 Obvious electrons migration around the interfaces of the above four adsorptions 248 indicates that chemical adsorption happened. The adsorption was likely to happen via 249 O^{α} , where was also the centre for electron migration. When only O^{α} atom (Fig. 2(a) and 250 (b)) bonded to the facet, fewer electrons were lost but apparent electrons enrichment 251 was observed around the O^{α} atom (Fig. 2(a)) and around the adsorbed hydrogen atom 252 (Fig. 2(b)). However, severer electron loses happened when AcOH bound to two Mo 253 atoms through O^{α} atom over the SB and hollow positions (in Fig. 2(c) and (d)). These 254 results are in agreement with the highest adsorption energy on these two sites. They 255 also reveal that the stable adsorption of AcOH on Mo (110) is achieved by strong electron migration from the molecule to the substrate through the bridge of C^{β} - O^{α} -Mo. 256

During modelling the decomposition of AcOH over Mo (110), the cleavage energy and reaction enthalpy were also calculated, in comparison with the free molecular decomposition of AcOH molecule in the absence of metal catalyst, which is shown in Fig. 3. The most favourable decomposition pathway of AcOH is illustrated in Fig. 4. The bond lengths and bond orders for the AcOH molecule and its most probable intermediates during the decomposition are shown in Table. S3 (a) and (b).

For the free molecular decomposition of AcOH, the $C^{\beta}-O^{\alpha}$ bond showed the largest cleavage energy of 8.46 eV, which was followed by the C^{α} -H bond with 5.85 eV and the C-C bond with 5.17eV. The bonds of O^{β} -H bond and $C^{\beta}-O^{\beta}$ exhibited similar cleavage energy of 4.51 eV and 4.67 eV respectively. All the cleavage reactions in free molecular decomposition were endothermic as there was no new bond formed.



Fig. 3. The cleavage energy for the free molecular decomposition of AcOH and FA





Fig. 4. The decomposition pathways of AcOH over Mo (110) facet

271 The decomposition of AcOH over Mo facet was the most likely to start with the cleavage 272 of the hydroxyl group with a small energy barrier of 0.11 eV, producing acetate and one 273 H atom. The acetate could then lose the O^{β} atom and one H sequentially to form CH₂CO. 274 All the cleavage reactions in the decomposition of AcOH over Mo facet were exothermic, 275 except for the final H-CC cleavage step, which was endothermic with $\Delta H = 0.43$ eV.

276 The reaction modelling results reveal that the bulk decomposition of AcOH over Mo (110) 277 is exothermic, which is much more thermodynamically favoured compared to its free 278 molecular decomposition. Besides, the significant decrease in cleavage energy 279 observed during the catalytic decomposition indicates the excellent effect of Mo facet in 280 facilitating the cleavage of AcOH. The most favourable decomposition pathway of AcOH 281 over Mo (110) is CH₃COOH \rightarrow CH₃COO+H \rightarrow CH₃CO+O+H \rightarrow CH₂CO+H+O+H \rightarrow 282 $CH_2C+O+H+O+H \rightarrow CHC+H+O+H+O+H \rightarrow CC+2H+O+H+O+H$, which is similar to the 283 reported decomposition pathways [36]. The highest energy barrier of 1.19 eV among all 284 the elementary reactions implies that the decomposition of CHC to CC is the rate-285 determining step. It is also found that the decarboxylation of CH₃COO was suppressed 286 at the beginning, and hydrogen is released during the whole decomposition process. The 287 C-C bond is stable throughout the decomposition, and the remaining C-C anchoring on 288 the Mo facet would result in carbonaceous deposits after further accumulation or they 289 crack to produce CO₂ & CH₄ with hydrogen and oxygen generated in previous steps [34].

290 3.3 FA as hydrogen donor on Mo (110) facet

The adsorption energies of FA onto the four different adsorption sites and in four different configurations are specified in Table 2. The configurations with the binding of O^{α} -Mo ((1) and (2)) showed the highest adsorption energy of 1.15 eV and 0.75 eV respectively. Configuration with O^{β} -Mo led to the adsorption energy of 0.51 eV. Parallel adsorption (configuration (4)) exhibited the lowest adsorption energy of 0.18 eV. Regarding different sites, the adsorption onto the top site led to the highest adsorption energy, followed by SB, LB and hollow sites with adsorption energies of 0.15 eV, 0.16 eV and 0.12 eV respectively. These results predict that FA is more likely to adsorb onto the top site of Mo (110) facet through a single O^{α} -Mo (110) bond. EDD was analysed for the adsorption configurations (1), (2), (3) with high adsorption energies to investigate the interface interaction, as shown in Fig. 5.



302

303

(a)

(b)



304

305

(C)

Fig. 5. Colour mapping of electron density difference (threshold value: +/- 0.05
electrons/Å³) for FA adsorption configurations of (a) Top ①, (b) Top ②, and (c) Top ③
(Loss of electrons is indicated in blue, while electron enrichment is indicated in red)

309 The electron migration exhibited in Fig. 5 indicates that all three adsorption 310 configurations are in the form of chemisorbed species, where O atom locates at the 311 centre of electron migration. Significant electrons migration has been observed for the 312 configuration with O^{α} -Mo bond. Fig. 5(a) shows that strong electron lose has taken place 313 around the O^{α} , but apparently electrons were enriched around O^{α} and O^{β} as well as the 314 hydroxyl hydrogen atom. Similar but minor electron migrations were also observed in the 315 other two cases, as demonstrated in Fig. 5 (b) and (c). The strong electron migration 316 implies large adsorption energy, in line with what has been observed in the AcOH 317 adsorption. However, AcOH leads to mainly electron lose, while the binding between FA 318 molecule and Mo (110) facet is found not only results from electron lose, but also from a 319 strong electron enrichment. This could be ascribed to the electron-donating nature of the 320 methyl in AcOH molecule.

Based on the reaction modelling for the decomposition of FA, the most probable reaction pathways with the lowest energy barriers are shown in Fig. 6, and the bond lengths and bond orders for the FA molecule and its most probable intermediates during the decomposition are presented in Table S4. (a) and (b). The free molecular decomposition of FA in the absence of metal catalyst is also shown in Fig. 3 for comparison.

In the decomposition via the free molecular mechanism, FA showed similar cleavage energy as AcOH for the bond $C^{\beta}-O^{\alpha}$ (8.59 eV), and slightly lower C-H cleavage energy (4.38 eV) than that of C_{α}-H in AcOH. While the bond C-O^{β} (4.91 eV) and bond O^{β}-H (5.66 eV) were more stable compared to AcOH. All the cleavage reactions in the free molecular decomposition were endothermic as no new bonds were formed.



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Reaction Coordinate



The decomposition of FA over Mo (110) started by the cleavage of O-H with a small energy barrier of 0.79 eV, and was followed by the cleavage of the bonds $C-O^{\beta}$ and $C=O^{\alpha}$ in sequence with energy barriers of 1.29 eV and 0.71 eV respectively. Finally, the C-H bond cleaved with an energy barrier of 1.01 eV. All the elementary decomposition reactions were exothermic, as shown in Fig. 6.

The modelling results confirm that Mo significantly decreases the cleavage energy of each bond. For instance, the cleavage energy of hydroxyl O-H bond was lowered from 5.66 eV in the free molecular decomposition to 0.79 eV over Mo facet. Besides, the exothermic property of each step in the catalytic decomposition makes it more thermodynamically favoured compared to the free molecular decomposition. The most probable decomposition pathway for FA over Mo facet is HCOOH \rightarrow HCOO+H \rightarrow

344 $HCO+O+H \rightarrow HC+O+O+H \rightarrow H+C+O+O+H$, indicating that O-H and C-H remain the 345 weakest and strongest bond respectively. The highest energy barrier among the elementary reactions was 1.29 eV for the C-O^{β} cleavage, revealing that it is the rate-346 347 determining step. When O^{β} is cleaved, the bond between C and the other oxygen (O^{α}) is 348 weakened significantly with the cleavage energy barrier of only 0.71 eV. The cleavage 349 of C=O^{α} was the exothermic reaction with a significant enthalpy change of 2.15 eV. This 350 result indicates that the cleavage of the second C-O bond is the most thermodynamically 351 favoured step for both FA and AcOH. One H atom was released at the beginning of the 352 FA decomposition process and another one at the final step. The single C atom left on 353 the Mo facet might result in carbonaceous deposit or CO₂ generation with additional 354 synthesis reactions [33].

The electron migration within the FA molecule under the effect of Mo is not as significant as that of AcOH. However, the loss of electrons in C=O^{α} still makes it cleave easily prior to C-H bond. The most electron enrichment takes place to the C-O^{β} bond, leading to the highest energy barrier of 1.29 eV for its cleavage reaction.

359 In the catalytic decomposition of AcOH and FA, Mo causes electron migrations between 360 the interfaces. Consequently most of the bonds in the two acid molecules are significantly 361 weakened. For instance, it was found that both decompositions start with hydroxyl O-H 362 cleavage with much cleavage energy compared to the free molecular decomposition. 363 Besides, the C=O $^{\alpha}$ bonds remain to be the strongest bond in both unabsorbed molecules, 364 but is readily cleaved over Mo facet with much smaller cleavage energy (energy barrier 365 for catalytic reactions), in comparison with the free molecular decomposition. In presence 366 of Mo, AcOH can release two H atoms from hydroxyl and methyl group respectively with 367 low energy barriers, but it requires larger energy to donate another two H atoms from the 368 methylene; the C-H bond becomes stronger when the C was unsaturated inside the 369 molecule. In FA decomposition, the hydroxyl hydrogen is also released at the early stage of its decomposition, however it exhibits higher energy barrier compared to the hydroxyl hydrogen cleavage in AcOH. The other hydrogen in FA is found to be more difficult to release prior to the cleavage of carboxyl oxygen atom, so that it remains to be the final step of the decomposition. The modelling results predict that AcOH possesses a better potential as a hydrogen donor compound over Mo (110) facet.

375 3.4 Dissociation of H₂ over Mo (110)

376 The adsorption of H_2 was modelled and the adsorption energy is presented in Table 2. 377 The results reveal that there were only two possible configurations (vertical and parallel) 378 and two probable sites (top and hollow) for H₂ adsorption. The adsorption showed similar 379 adsorption energy in parallel configuration (0.57 eV) and vertical configuration (0.56 eV). 380 A higher adsorption energy of 1.64 eV resulted from the adsorption onto the hollow 381 position, which is the only plausible adsorption position observed in the modelling other 382 than the top position. It is noteworthy that hydrogen molecule was dissociated into two 383 atoms, anchoring on two hollow positions as shown in Fig. 7, known as dissociative 384 adsorption [52]. The overall dissociative adsorption process of H₂ remained to be 385 exothermic.



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387

Fig. 7. Configuration of H₂ adsorption onto hollow position

388 3.5 Comparison of enthalpy changes

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Modelling results have predicted that the overall decomposition reactions of AcOH and FA over Mo(110) facet are both exothermic, implying the energy released by new binding formation between their intermediates and the Mo facet is more than the enthalpy change for the old binding cleavage [33,36,39]. Thus, the decomposition of both hydrogen donors over Mo (110) facet is hydrodynamically favoured, similar to the dissociative adsorption of H₂. The enthalpy changes for the adsorption and decomposition of AcOH, FA, and H₂ are calculated respectively, and summarized in Fig. 8.



Fig. 8. Overall enthalpy change of adsorption and decomposition of hydrogen donor compounds

Hydrogenation and hydrogenolysis reactions normally happen with single hydrogen migration or spillover during the HDO process [53,54], suggesting that the activation of single hydrogen atom would be an essential step for the hydrogen donation. Therefore, the cleavage of metal-single hydrogen bond plays a key role in such a process. The cleavage (activation) enthalpy of the hydrogen-metal bond is calculated and compared with the adsorption and decomposition enthalpy for each hydrogen donor compound, asshown in Fig. 8.

406 Additional calculation for the activation of H atom was performed, and the results show 407 that 3.94 eV was required to activate a single hydrogen atom on Mo (110) facet. Previous 408 results predicts that the enthalpy change for the dissociative adsorption of hydrogen 409 molecule is -1.64 eV, which is bigger than those for AcOH and FA (-1.24 eV and -1.15 410 eV respectively), nevertheless it is inadequate for one single H atom activation. Due to 411 the exothermic decomposition of AcOH and FA, the overall enthalpy change for the bulk 412 adsorption and decomposition process of AcOH and FA were -6.50 eV and -6.19 eV 413 respectively, which are adequate for the activation of H-metal bond. Therefore, the 414 cleavage of H-metal bond could be more facilitated during the decomposition of AcOH 415 and FA. In other words, more hydrogen atoms as well as other decomposed atoms would 416 be likely to migrate and transfer during the decomposition of AcOH and FA compared to 417 the H₂ gas. In this sense, the bio-acids showed good potential to replace H₂ gas as 418 alternative hydrogen donors.

419 **4. Conclusions**

420 The performance of AcOH and FA as hydrogen donors over transition metal facets were 421 investigated and compared with H_2 . The adsorption model showed that facets of Ru 422 (0001), Mo (110), Ni (111) and Pd (111) led to the strongest binding to the AcOH 423 molecule among the ten metal facets. Mo (110) was selected as the metal catalyst for 424 further investigating the hydrogen donation of bio-acids as it is a base metal and the lack 425 of atomic-level investigation. EDD analyses revealed that the bindings between bio-acids 426 and Mo were dominated by the electron migration from the molecule to the Mo (110) 427 facet, while electron enrichments in the acid molecule has also been observed for FA 428 adsorption. Mo exhibited positive effect on weakening the bond strength for both acids, 429 and facilitated their cleavage. The reaction model predicted that the most favourable

430 decomposition pathway for acetic acid over Mo (110) was via $CH_3COOH \rightarrow CH_3COO+H$ 431 \rightarrow CH₃CO+O+H \rightarrow CH₂CO+H+O+H \rightarrow CH₂C+O+H+O+H \rightarrow CHC+H+O+H+O+H \rightarrow 432 CC+2H+O+H+O+H, where hydroxyl hydrogen and the methyl hydrogen were more 433 readily to be released than the H atoms from the unsaturated methylene. The most 434 favourable decomposition pathway for FA over Mo (110) facet was through HCOOH \rightarrow 435 $HCOO+H \rightarrow HCO+O+H \rightarrow HC+O+O+H \rightarrow H+C+O+O+H$, where the hydroxyl hydrogen 436 can be cleaved and donated at an early stage. Hydrogen showed its undoubtable merits 437 in hydrogen donation because of its dissociative adsorption property over Mo facet. 438 However, compared to the decomposition of H_2 gas, the larger enthalpy change resulted 439 from the exothermic decomposition of the bio-acid compounds would more benefit the 440 activation of decomposed atoms from the catalyst surface, including the hydrogen 441 donation in terms of the single H atom activation and migration. Within the bio-acids, 442 AcOH exhibited a better potential than FA as a hydrogen donor because it released more 443 hydrogen atoms with lower energy barriers. The modelling predicted that AcOH has good 444 potential as an alternative hydrogen donor compound over Mo (110) facet.

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448 Author Contributions

- All the authors have given approval to the final version of the manuscript.
- 450 **Notes**
- 451 The authors declare no competing financial interest.

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