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Au@*h***-Al2O3 analogic yolk-shell nanocatalyst for highly** 2 **selectively synthesis of biomass-derived _D-xylonic acid via regulation of structure effect**

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24 **Abstract**

25 Selective oxidation of biomass-based monosaccharides into value-added 26 sugar acids is highly desired, yet limited success of producing $D-XY$ lonic 27 acid has been achieved. Here, we report an efficient catalyst system, Au 28 nanoparticles anchoring on the inner wall of hollow Al_2O_3 nanospheres 29 ($Au@h-Al_2O_3$), which could catalyze the selective oxidation of $_D$ -xylose 30 into $D-xy$ lonic acid under base-free conditions. The mesoporous $A1_2O_3$ 31 shell as adsorption sites first adsorbed $_D$ -xylose, the interface of Au</sub> 32 nanoparticles and Al_2O_3 as active sites spontaneously dissociated O_2 , and 33 the exposed Au nanoparticles surface as catalytic sites drove the 34 transformation. With this catalyst system, the valuable D -xylonic acid was 35 produced with excellent yields in the aerobic oxidation of $_D$ -xylose.</sub> 36 Extensive works showed that the $Au@h-Al_2O_3$ is an efficient catalyst 37 with highly stability and recycling.

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46 **Key words:** p-xylose · p-xylonic acid · Au@h-Al₂O₃ · structural effect 47 regulation system · selective oxidation

Introduction

 Use of biomass-derived feedstocks provides an environmentally friendly and sustainable route for the production of chemicals. Organic acid is a versatile, renewable chemical with a wide industrial application profile 52 both as a solvent and as a building block.^{1,2} It is also widely used in food, 53 pharmaceutical, cosmetic, detergent, polymer, and textile.^{3,4} Transformation of biomass into value-added organic acids is highly desired, but is still very limited owing to the lack of efficient catalytic system and method. Recently, monosaccharide oxidation (especially selective oxidation), as a crucial element of organic acid production, offers an attractive option on the conversion of biomass resources into chemicals and sugar acids.

 $D-Xylose$, one of the main sugar units in hemicelluloses, is the 61 second most abundant sugar in nature.⁵ Many efforts have been made on 62 utilizing of $_D$ -xylose by fermenting⁶ or other processes to obtain fuels and 63 chemicals.⁷⁻¹⁰ Among these products, $_D$ -xylonic acid is one of the top 30 high-value chemicals identified by the US Department of Energy. It is a key compound for the preparation of 1,2,4-butanetriol or energetic 66 materials 1,2,4-butanetriol trinitrate.¹¹ Currently, as a concrete additive, D -xylonic acid can effectively improve concrete dispersion.¹² Furthermore, $D-xy$ lonic acid can be used as an efficient biocatalyst for organic 69 transformations.¹³ Other applications of p -xylonic acid have been 70 reported in food, pharmaceutical, and agriculture.^{14,15} Therefore, 71 innumerous scientists have devoted to themselves to the synthesis of D - xylonic acid research. Among all the literatures, microbial production of 73 $D-Ny$ p-xylonic acid plays a significant role in the synthesis of $D-Ny$ p-xylonic acid. As early as 1946, Lockwood and Nelson reported that the production of _D-xylonic acid could occur in the Pseudomonas.¹⁶ Later, a lot of studies

76 have found that $_D-xylonic$ acid could be produced in the oxidative</sub> 77 metabolism of $_D$ -xylose by some archaea and bacteria.^{9, 17-20} But until now, 78 commercial production of $_D-xylonic$ acid has not been developed,</sub> 79 reflecting the current limited market for D -xylonic acid. The reason lies in two facts: the one is that many others oxidizing enzymes are produced by bacteria strains which directly contribute to the conversion of other sugars present in lignocellulosic hydrolysates, the other is that the engineered yeast strains have low _D-xylonic acid accumulation rate and yield.⁹ 84 Besides, industrial scale production of $_D$ -xylonic acid needs the high cost</sub> of peptone and/or yeast extract media as nitrogen sources, which decide that the production methods are impractical. Furthermore, although the application of chemical synthesis in industrial production is widespread, 88 few studies have been reported for the synthesis of D -xylonic acid by chemical synthesis.

 Au-based catalysts are found to be superior to Pt, Pd and Pt/Pd 91 bimetallic catalysts for the oxidation of glucose to gluconic acid, 21 as Au nanoparticles (NPs) have low sensitivity to oxygen poisoning and high activity in a wide range of pH values. However, the free Au NPs with high surface energy are easy to agglomerate, which affect its activity. To avoid this drawback, functional supports with a strong interaction with 96 Au,²²⁻²⁶ pores²⁷⁻³⁰ and core/yolk-shell architecture³¹⁻³³ are developed to improve the stability of Au NPs. The core/yolk-shell architecture featured the advantages of high permeability, low density, large surface area, 99 multifunctionality, and high loading capacity, $31, 34-38$ Furthermore, modification of the core/yolk-shell structure to give a nanocatalyst with metal NPs embedded on its inner wall can not only improve the stability of metal NPs, but also generate a synergistic effect on the interface 103 between the metal NPs and the shell. $^{39, 40}$

104 Herein, we report a new catalyst of Au NPs anchoring on the inner 105 wall of mesoporous hollow Al_2O_3 nanospheres, prepared via the 106 technique of core/yolk-shell architecture. The $Au@h-Al_2O_3$ catalyst can 107 selectively oxidize D -xylose to D -xylonic acid. The correlation between 108 the unique structure and catalytic activity of $Au@h-Al_2O_3$ was revealed 109 through comparative experiments. The adsorption sites were investigated 110 by a technique called quartz crystal microbalance with dissipation 111 monitoring (QCM-D) and DFT calculations. The active sites for the 112 spontaneous dissociation of $O₂$ were studied by DFT calculations. The 113 catalytic sites of $Au@h-Al_2O_3$ were verified by the comparison 114 experiment of sulfhydryl coated surface of Au NPs.

115 **Results and discussion**

116 The synthetic scheme of the $Au@h-Al_2O_3$ is shown in Fig. 1. A soap-free 117 emulsion polymerization method was adopted to prepare a core/shell 118 structure of $PS-co-P4VP₁⁴¹$ which was then used as scaffold for the 119 immobilization of Au NPs. The dispersed Au NPs were then anchored 120 onto the PS- co -P4VP core/shell microspheres by coordination of $Au³⁺$ 121 (from HAuCl4) with the P4VP shell first and then by reduction with 122 NaBH₄.⁴¹ Afterwards, uniform layers were gradually growing onto the 123 surface of Au@PS-*co*-P4VP by controlling the precipitation process in a 124 buffer solution of formic acid-ammonium formate.⁴² Finally, the resultant 125 NPs were calcined to remove PS-*co*-P4VP and allow the Au NPs to be 126 distributed on the inner wall of the hollow Al_2O_3 sphere. In this process, 127 PS-*co*-P4VP microspheres played a dual role, a scaffold for 128 immobilization of dispersed Au $NPs⁴¹$ and a template for the synthesis of 129 Al_2O_3 coating.

131 **Fig. 1** Schematic illustration of the synthetic route of Au ωh -Al₂O₃.

132 The morphology and chemical composition of the intermediate and 133 Au ω/h -Al₂O₃ were first characterized by scanning electron microscope 134 (SEM). As shown in Fig. 2A, the pure PS-*co*-P4VP microspheres with an 135 average size of about 490 nm were observed. As compared with the PS-136 *co*-P4VP microspheres, Au@PS-*co*-P4VP microspheres showed no 137 significant difference on morphology and size (Fig. 2B). The elements of 138 Au@PS-*co*-P4VP were mainly Au, C, O and N (the inset of Fig. 2B), 139 indicating that Au NPs were successfully deposited on the surface of PS-140 *co-P4VP.* After the precipitation of $A₁O₃$ on the surface of $Au@PS-co-$ 141 P4VP microspheres (using $Al_2(SO_4)$ ₃ as the precursor of Al_2O_3), a thin 142 shell was coated to produce Au@PS-*co*-P4VP@Al₂O₃ microspheres, 143 which was confirmed by a slight increase in the size from 490 to 510 nm 144 (Fig. 2C, 2H). The elemental mapping of O, Al, Au, C and N (Fig. 2I-2M) 145 further evidenced the successful preparation of $Au@PS-co-P4VP@Al_2O_3$ 146 microspheres. After Au@PS-co-P4VP@Al₂O₃ NPs were calcined at 550 147 °C, the polymeric template was removed, and Au NPs were left on the 148 inner wall of hollow A_1O_3 sphere, producing $Au@h-A1_2O_3$ (Fig. 2D and

149 Supporting Information Fig. S1). Only Al, Au and O were observed in the 150 EDX area scanning of $Au@h-Al_2O_3$, which indicates that the cavity 151 structure was achieved after the calcination (Fig. 2E-G).

152 The transmission electron microscopy (TEM) image of pure PS-*co*-153 P4VP microspheres in Fig. S2A is consistent with the SEM results. In the 154 TEM images of Fig. S2B, the Au@PS-co-P4VP@Al₂O₃ microspheres 155 with a diameter of ~510 nm. The inter bright spots represent the PS-*co*-156 P4VP core; the dark spots indicate the Au NPs, while the peripheral dark 157 circle corresponds to $A₁Q₃$ shell. After PS- co -P4VP core was removed, 158 the shell (of 10 nm in thickness) and the cavity were clearly observed by 159 TEM and STEM (Fig. 3A and 3B); and Au NPs with a diameter of ~1.87 160 nm were observed on the inner wall of $A₁O₃$ microspheres (the inset in 161 Fig. 3B). Besides the diameter of Au NPs slightly increased from 1.87 to 162 2.20 nm (Supporting Information Fig. S3C), and no significant difference 163 in size was observed for $Au@h-Al_2O_3$ after being used for 10 times 164 (Supporing Information Fig. S3A, S3B and S4), indicating the high 165 stability of $Au@h-Al_2O_3$. The Au element analyzed with XPS is shown in 166 Fig. 3D. The signal of Au 4f can be resolved into two peaks. The binding 167 energies of Au 4f_{5/2} and Au 4f_{7/2} locating at 87.7 and 83.9 eV indicate Au⁰, 168 while the 89.6 and 86.3 eV demonstrate Au_2O_3 , respectively.⁴³ There was 169 no significant difference in Au valence for $Au@h-Al_2O_3$ before and after 170 catalysis for D -xylose oxidation. For the freshly prepared $Au@h$ -Al₂O₃, 171 the peaks locating at 83.9 and 87.5 eV reveal only the valence of Au 172 species. After being used for the synthesis of $_D$ -xylonic acid, the peaks</sub> 173 shift slightly to the lower binding energies at 83.5 and 87.3 eV 174 (Supporting Information Fig. S5), respectively, indicating that the valence 175 of Au species did not change much during the catalytic oxidation process. 176 In addition, the difference in O 1s and Al 2p spectra for $Au@h-Al_2O_3$ was

177 not significant before and after being used for D -xylose oxidation 178 (Supporting Information Fig. S6). The Au contents determined by ICP-179 MS were 2.80 wt% and 2.49 wt% for $Au@h-Al_2O_3$ before use and after 180 10 catalytic cycles, respectively, indicating that $Au(\hat{a}_i h - A_1 O_3)$ has 181 excellent stability (Supporting Information Table S1). Fig. S7A shows the 182 HAADF-STEM spectrum of $Au@h-Al_2O_3$. The white spots indicate the 183 Au NPs and the peripheral white circle corresponds to the $A₁O₃$ shell. 184 The elemental mapping of O, Al and Au in $Au@h-Al_2O_3$ are shown in 185 Fig. S7B-D, demonstrating the actual distributions of O, Al, and Au 186 elements in the catalyst. Fig. 3C shows the HRTEM image of Au@*h*- 187 Al₂O₃. There were mainly two types of lattice fringes with interplanar 188 spacing of 2.32 and 2.0 Å, corresponding to (111) and (200) crystal 189 planes of Au NPs, respectively. The inset in Fig. 3C shows the diffraction 190 pattern of Au NPs. To illustrate the position of Au NPs in $Au@h-AbO_3$, a 191 STEM spectrum of Au NP is given in Fig. S8A; then a line scan of the 192 Au NP is shown in Fig. S8B, which gives a preliminary explanation of 193 the position of Au NPs in $Au@h-Al_2O_3$. Fig. 3F shows the XRD pattern 194 of $Au@h-Al_2O_3$ catalyst. $Au@h-Al_2O_3$ displays a good crystallinity and 195 the diffraction peaks at $2\theta = 38.27^{\circ}$, 44.6°, 64.68° and 77.55° were 196 assigned to (111), (200), (220) and (311) reflections of the gold lattice, 197 well agreeing with the results from Fig. 3C. The 10^{th} reused Au ωh -Al₂O₃, 198 Au/Al₂O₃ and other Au catalysts (Supporting Information Fig. S4F and 199 S9a-d) also showed the similar diffraction peaks.

201 **Fig.** 2 SEM of PS-co-P4VP (A), $Au@PS-co-P4VP$ (B) (the inset is the EDX of 202 $Au(\partial P)S-co-PAVP$, $Au(\partial P)S-co-PAVP(\partial A)_{2}O_{3}$ (C, H), $Au(\partial P)A_{2}O_{3}$ (D) and element 203 mapping images of $Au@h-Al_2O_3$: Al element (E), Au element (F), and O element (G), 204 the element mapping images of Au@PS-co-P4VP@Al₂O₃: Al element (I), Au 205 element (J), C element (K), N element (L) and O element (M).

206 70 To further determine the distribution patterns of Al, O and Au in 207 $Au(\omega, h-A_1, O_3, Au(\omega, h-A_1, O_3)$ with 10 times $HAuCl_4$ addition was prepared 208 and the obtained products were named $Au(a/h - Al₂O₃ - 10$. Besides an 209 obvious increase in diameters of Au NPs (the inset in Supporting 210 Information Fig. S10B), no significant difference was observed between 211 $Au(\partial h - Al_2O_3 - 10$ and $Au(\partial h - Al_2O_3)$ (Supporting Information Fig. S10A-E). 212 The elemental mapping of $Au@h-Al_2O_3-10$ in Fig. S10F-H gave a clear 213 distribution of O, Al and Au. The TEM and STEM of the cross section 214 diagram of $Au@h-Al_2O_3-10$ visually showed the distribution of Au NPs 215 (Fig. 3E, 3H and Supporting Information Fig. S11). The line scan 216 spectrum and elemental mapping of the cross section diagram of Au@h- 217 Al_2O_3 -10 further presented the actual distributions of O, Al, and Au 218 elements (Fig. 3G and 3I-3K). All the results demonstrated that Au NPs 219 are well distributed on the inner wall of the Al_2O_3 hollow sphere.

221 **Fig. 3** TEM and STEM of $Au@h-Al_2O_3$ (A, B). The inset B is the size distributions of 222 Au NPs from 300 particles. (C) High-resolution TEM image of $Au@h-Al_2O_3$ (the 223 inset is the diffraction patterns of Au NPs). (D) Au 4f spectra for $Au(\partial h-A_2O_3$. (E) 224 TEM of the cross section diagram of $Au@h-Al_2O_3-10$. (F) The XRD pattern of 225 Au $\omega_a h$ -Al₂O₃. (G) The line scan of cross section diagram of Au $\omega_a h$ -Al₂O₃-10. (H) 226 STEM-HAADF image of the cross section diagram of $Au@h-AbO_3-10$ and element 227 mapping images: (I) O element, (J) Al element, and (K) Au element.

228 The N₂ sorption isotherms of Au/Al₂O₃, Au@PS-*co*-P4VP@Al₂O₃ 229 and $Au@h-Al_2O_3$ exhibit type IV isotherm patterns (Supporting 230 Information Fig. S12), suggesting the presence of mesopores in these 231 materials. Compared to Au/Al_2O_3 and Au/QPS -co-P4VP $@Al_2O_3$, $Au/Q/h$ - 232 Al₂O₃ showed more uniform and well-developed mesopores with a 233 diameter of 2.19 nm, favoring the diffusion of reactants and products in 234 and out of the catalyst. The BET results showed that the surface area of 235 Au@h-Al₂O₃ increases from 10.09 to 100.45 m²/g after removing PS-*co*-236 P4VP, indicating a higher contact area between D -xylose and Au ω *h*- 237 Al₂O₃. Fig. S13 shows no significant difference in the thermal weight loss 238 between $Au@h-Al_2O_3$ and Au/Al_2O_3 , demonstrating that the catalysts 239 have a good thermostability. Although weight loss occurrs to Au@PS-*co*-

240 P4VP@Al₂O₃ when the temperature was higher than 350 °C, it has no 241 impact on the aerobic oxidation of $_{D}$ -xylose because the reaction was 242 carried out in the temperature range of $100-160$ °C.

243 Prior to comprehensively investigate the catalytic activity of Au@*h*-244 Al₂O₃ on the aerobic oxidation of $_D$ -xylose to $_D$ -xylonic acid, the affinities 245 of $_D$ -xylose to various surfaces $(Al_2O_3,$ silica, and gold) were compared 246 by a technique of quartz crystal microbalance with dissipation monitoring 247 (OCM-D). Fig. S14 shows that the frequency change of $A₁O₃$ surface is -248 7.3 Hz after water rinsing, which is equivalant to 129.2 mg/m², higher 249 than those of silica $(-1.4 \text{ Hz}$ and $24.8 \text{ mg/m}^2)$ and gold $(-4.9 \text{ Hz}$ and 86.7 Hz 250 mg/m^2) surfaces. The results suggest that Al_2O_3 has a much stronger 251 affinity to $_D$ -xylose than silica and gold.

252 The adsorption energy of $_D$ -xylose on Al₂O₃ was calculated by DFT 253 methods using DMOL3 module from Materials Studio 2017.⁴⁴ A model 254 of the dehydrated Al_2O_3 (110) surface (Fig. 4A), the same as that used by 255 Falamaki and co-workers, 45 was chosen to investigate the interactions 256 between $D-xy$ lose and the Al₂O₃ crystal. The adsorption energy of $D-$ 257 xylose on Al_2O_3 (110) was -70 kJ/mol. In addition, the dissociation 258 energy O_2 on the interface of Au NPs and Al_2O_3 shell was also calculated 259 by DFT as -245.68 kJ/mol (Fig. 4B and 4C), suggesting that the O_2 260 molecule can be dissociated spontaneously on the interface of Au NPs 261 and Al_2O_3 shell. The generated negative oxygen ion is expected to react 262 with water to produce oxidizing substances. These results indicate that it 263 is feasible to synthesize D -xylonic acid from D -xylose catalyzed by 264 Au ω *h*-Al₂O₃.

265 For the aerobic oxidation of $_D$ -xylose to $_D$ -xylonic acid catalyzed by 266 Au $@h$ -Al₂O₃, the reaction was carried out at different temperatures (100-

267 160 °C) in the presence of O_2 (3.0 MPa). As shown in Table S2, the yield 268 of p -xylonic acid first significantly increased from 38.6% to 83.3% when 269 reaction temperature was raised from 100 to 130 °C (Supporting 270 Information Table S2, entries 1-4), and then gradually decreased as the 271 temperature reached 160 °C (Supporting Information Table S2, entries 4- 272 7). The decreasing yield may be due to the production of byproducts (e.g., 273 formic acid, acetic acid and lactic acid) at high temperature. The 274 selectivities of $_D$ -xylonic acid increased first and then decreased with the 275 increasing temperature (Fig. 4E), while the conversion of $_D$ -xylose</sub> 276 increased with the increase of temperature (Fig. 4F), implying 277 endothermic nature of the overall reaction. Thus, $130 \degree C$ was chosen as 278 the optimal reaction temperature. After that, the reaction was tested under 279 different O_2 pressure. It was found that only 14.7% of $_D$ -xylonic acid was 280 obtained when air was used as oxidizer (Supporting Information Table S3, 281 entry 1). As the O_2 pressure increased to 3.0 MPa, the yield remarkably 282 increased from 14.7% to 83.3% (Supportiing Information Table S3, 283 entries 1-6). A slight decrease of $_D$ -xylonic acid yield (79.6%) 284 (Supporting Information Table S3, entry 7), however, was observed when 285 O₂ pressure increased to 4.0 MPa. The selectivity of $_D$ -xylonic acid under 286 different O_2 pressure also showed similar change to that of the yield of D -287 xylonic acid, and the conversion of $_D$ -xylose increased with the increase 288 of O_2 pressure (Supporting Information Fig. 4E and 4F).

290 **Fig. 4** The model of $_D$ -xylose adsorption energy on Al_2O_3 (110) (A) and O_2 291 dissociation energy on the interface of Au and Al_2O_3 in $Au@h-Al_2O_3$ (B and C) 292 calculated by DFT (grey: C, white: H, pink: Al, red: O, yellow: Au). (D) Proposed 293 reaction mechanism for the base-free oxidation of $_D$ -xylose to $_D$ -xylonic acid 294 catalyzed by $Au@h-Al_2O_3$ (left) and oxygen dissociation reaction mechanism (right). 295 The selectivity of $_D$ -xylonic acid (E) and conversion of $_D$ -xylose catalyzed (F) by 296 Au $@h$ -Al₂O₃ under different temperatures and oxygen pressures. The selectivity of _D-

297 xylonic acid (G) and conversion of $_D$ -xylose (H) catalyzed by $Au@h$ -Al₂O₃ under 298 different reation times and catalyst dosages.

299 The dosage (relative to 0.25 g $_D$ -xylose) of Au ω/h -Al₂O₃ also played 300 an important role in the aerobic oxidation of $_D$ -xylose to $_D$ -xylonic acid. 301 The yield of $_D$ -xylonic acid increased with increasing dosage of $Au@h$ - 302 Al₂O₃ (Supporting Information Table S4, entries 1-4). An increase in 303 dosage from 20 to 30 mg, however, did not lead to a significant increase 304 in the yield of $_D$ -xylonic acid (Supporting Information Table S4, entries 305 4-5). It is possible that $_D$ -xylose molecules adsorbed on the Al₂O₃ shell 306 became unstable species (more active), which reduced the activation 307 energy of the reaction. Similar trend was observed for the selectivity of D -308 xylonic acid, as shown in Fig. 4G. The conversion of $_D$ -xylose increased</sub> 309 with increasing dosage of $Au@h-Al_2O_3$ (Fig. 4H). Therefore, 20 mg of 310 Au ω/h -Al₂O₃ is optimal for this reaction. Furthermore, the effect of 311 reaction time on the oxidation of $_D$ -xylose was also examined. With the 312 growth of reaction time, both the yield (Supporting Information Table S5, 313 entries 1-4) and selectivity (Fig. 4G) increased, and then decreased after 314 60 min. This phenomenon might be due to the generation of byproduct 315 (e.g., formic acid and acetic acid) from the produced $_D$ -xylonic acid with 316 increasing reaction time (Supporting Information Table S5, entries 4-6). 317 The conversion of $_D$ -xylose also increased with the increase of reaction</sub> 318 time (Fig. 4H). Furthermore, the $_D$ -xylonic acid yield and the conversion 319 of $_D$ -xylose decreased slightly from 83.3% to 80.5% and from 93.8% to 320 92.8% in the tenth recycling run, respectively. Thus, $Au@h-Al_2O_3$ is 321 highly stable and recyclable (Supporting Information Fig. S15).

322

Entry	Al_2O_3 -based Au catalyst	p -Xylonic acid $(\%)$
1	Au/Al_2O_3	21.6
$\overline{2}$	Au $@$ PS-co-P4VP $@$ Al ₂ O ₃	17.2
3	Au $(\partial h - Al_2O_3)$	83.3
$\overline{4}$	Au-SH- $(CH_2)_3$ -Si $(OCH_3)_3/Al_2O_3$	9.6
5	Au-SH-(CH ₂) ₃ -Si(OCH ₃) ₃ @PS-co-P4VP@Al ₂ O ₃	5.2
6	Au-SH-(CH ₂) ₃ -Si(OCH ₃) ₃ $@h$ -Al ₂ O ₃	13.0

324 **Table 1.** Base-free oxidation of $_D$ -xylose with structurally different Al_2O_3 -based Au catalyst.^a 325

326 ^[a] Typical reaction conditions: $_{D}$ -xylose (0.25 g), O₂ (3.0 MPa), water (25 mL), 327 Al₂O₃-based Au catalyst (20 mg), 130 °C, 60 min.

328 To better understand our catalyst, the catalytic efficiency of Au@*h*- 329 Al₂O₃ was compared with those catalysts that prepared in the literatures. 330 A series of experiments were carried out at the optimal condition and the 331 results are presented in Table S6. Although there was no significant 332 difference in the conversion of $_D$ -xylose between Au ω/h -Al₂O₃ and other 333 catalysts, the yield of $_D$ -xylonic acid (thus selectivity) was different. The 334 yield of $_D$ -xylonic acid catalyzed by $Au@h$ -Al₂O₃ was significantly 335 higher than those of other catalysts (Au/SBA-15⁴⁶: 16.8%, Au@PS-*co*-336 P4VP: 11.4% , $SiO_2-SH-Au$: 1.6% , C-SH-Au: 10.8%). Furthermore, the 337 effect of morphology of catalyst on the oxidation of $_D$ -xylose was also 338 investigated. As shown in Table 1, the yield of $_D$ -xylonic acid catalyzed</sub> 339 by Au/Al₂O₃ was higher than that catalyzed by Au@PS-*co*-P4VP@Al₂O₃, 340 while $Au@h-Al_2O_3$ showed a even higher yield than $Au/Al_2O_3^{47}$, 341 implying that the $Au@h-Al_2O_3$ catalyst with special structure can 342 effectively promote the formation of $_D-xylonic$ acid. All these results</sub>

343 suggest that the $Au@h-Al_2O_3$ has very high catalytic activity for the 344 aerobic oxidation of $D-xy\log$ into $D-xy\log$ acid compared to other 345 catalysts. In addition, the possible reaction pathways and descriptions was 346 given in supporting informatin (Table S7, Fig. S16 and Fig. S17)

347 To determine the catalytic sites of $Au@h-Al_2O_3$, we used 3-348 mercaptopropyltrimethoxysilane to treat the $Au@h-Al_2O_3$, and the 349 exposed Au NPs surface was covered with sulfhydryl groups. The 350 resulting $Au-SH-(CH_2)3-Si(OCH_3)3@h-Al_2O_3$ (Supporting Information 351 Fig. S18) was used to catalyze $_D$ -xylose. Only 13.0% yield of $_D$ -xylonic 352 acid was obtained, which well agrees with the results of Table S6 (entries 353 3 and 4). The phenomenon suggested that the catalytic sites of Au@*h*- 354 Al₂O₃ were the exposed Au NPs surface. Hence, we proposed a plausible 355 reaction mechanism of synthesis of D -xylonic acid from D -xylose 356 catalyzed by $Au@h-Al_2O_3$ (Fig. 4D). According to the early reports, ⁴⁸ 357 hydrogen peroxide, as a byproduct, was produced from the aerobic 358 oxidation of glucose to gluconic acid. The hydrogen peroxide acted as an 359 oxidant to oxidize glucose. In this work, our DFT calculations showed 360 that the O_2 molecule was dissociated spontaneously on the interface of 361 Au NPs and Al_2O_3 (Fig. 4B and 4C), and $_D$ -xylose was preferably 362 adsorbed on Al_2O_3 (Fig. 4A and Supporting Information Fig. S14). The 363 resulting negative oxygen ion can react with water to produce O_2 , 364 hydrogen peroxide and hydroxyl ions (Fig. 4D, right panel). Then the 365 isomerization reaction between D -xylose and chain intermediate may 366 occur in the presence of hydroxyl ions. Finally, the chain intermediate 367 was oxidized by hydrogen peroxide to produce D -xylonic acid.

368 **Conclusions**

369 In summary, we demonstrated a highly effective chemical method for the 370 synthesis of $_D$ -xylonic acid from $_D$ -xylose, catalyzed by $Au@h$ -Al₂O₃. 371 The catalytic oxidation of $_D$ -xylose was facile with high yields. The 372 correlation between the unique structure and catalytic activity of Au@*h*- 373 Al₂O₃ was revealed through comparative experiments. The Al₂O₃ shell 374 played a dual role in the stabilization of Au NPs and the adsorption of $_{D}$ -375 xylose. DFT calculations revealed that the unique structure between Au 376 NPs and Al_2O_3 provided a synergism for spontaneous dissociation of O_2 , 377 and hence promoted the formation of $_D$ -xylonic acid. At the condition of</sub> 378 130 °C, 3.0 MPa O_2 and 60 min reaction time, the _D-xylose conversion 379 was 93.8%, and the $_D$ -xylonic acid yield was 83.8%. Compared with other 380 Au supported catalysts (e.g. Au/SBA-15, SiO2-SH-Au, Au@PS-*co*-P4VP, 381 etc.), $Au@h-Al_2O_3$ showed selectivity toward the production of D -xylonic 382 acid. Au $\omega/h - A_2O_3$ was stable after 10 rounds of catalytic recycling. This 383 work provides a new facile route to selectively oxidize monosaccharides 384 into value-added sugar acids such as D -xylonic acid, using a type of 385 hermit crab typological nanocatalyst $Au@h-A_2O_3$.

386 **Experimental**

387 **Synthesis of Au@***h***-Al2O3 nanocatalyst**

 The styrene was pretreated before use. First, a certain amount of commercially available styrene was added into 10% sodium hydroxide 390 solution, and the system was placed stably for 10 min at 4 $^{\circ}$ C. The process was replicated for 3 times. Thereafter, the styrene was washed with water and dried with anhydrous magnesium sulfate overnight. Finally, the pure styrene was obtained by vacuum distillation and kept at 4° C.

 The polystyrene-*co*-poly(4-vinylpyridine) (PS-*co*-P4VP) core-shell microspheres were synthesized by one-stage, soap-free emulsion 397 polymerization.⁴⁹ Typically, 80 mmol of 4-vinylpyridine was mixed with 360 mL of water at room temperature, then 160 mmol of styrene was added to the above system under magnetic stirring. The mixture was stirred vigorously for another 30 min at room temperature. Thereafter, 401 1.30 g of $K_2S_2O_8$ was added into the reaction system, and the pH of the mixture was adjusted to 7 with 0.1 mol/L HCl aqueous solution. The reaction system was then filled with nitrogen and carried out at 80 °C for 24 h under vigorous stirring conditions. Finally, the excessive volume of 0.1 mol/L NaOH aqueous solution was injected into the above colloidal dispersion, and the obtained precipitates were washed with deionized 407 water and then dried in a vacuum at 50 °C overnight.

408 0.480 g PS-*co*-P4VP was dispersed in HAuCl₄ aqueous solution and stirred at room temperature for 17 h. After that, the pH of the mixture was adjusted to 7 with 0.1 mol/L NaOH aqueous solution, and then a 5-fold 411 excess volume of 0.1 mol/L NaBH₄ aqueous solution was added drop wise with vigorous stirring. The obtained Au@PS-*co*-P4VP core-shell microspheres were collected by centrifugation, washed thrice with 414 deionized water, and dried in a vacuum at 80° C.

 A certain volume of formic acid-ammonium formate buffer solution (pH = 4.4) was first prepared. Then, a certain amount of Au@PS-*co*- P4VP core-shell microspheres and aluminum sulfate were added and 418 dispersed by ultrasound for 15 min. The system was carried out at 70 $^{\circ}$ C 419 for 2 h with vigorous stirring. The obtained $Au@PS-co-P4VP@Al_2O_3$ was collected by centrifugation, washed with deionized water and oven-421 dried at 80 °C overnight. The Au $@h$ -Al₂O₃ was obtained by calcinating

422 Au@PS- $co-P4VP@Al_2O_3$ at 550 °C for 4 h. Changing the concentration of aluminum salts can easily tune the thickness of aluminum oxide shell.

Activity Tests. The catalytic synthesis of _D-xylonic acid were carried out in a 60 mL Teflon-lined stainless-steel autoclave. In a typical 426 procedure, 0.25 g $_{D}$ -xylose was dissolved in 25 mL water, and then a 427 certain amount of $Au@h-Al_2O_3$ was added into the solution and dispersed by ultrasound for 10 min. Then oxygen was purged into the reactor for three times before the reactor being sealed and pressurized with oxygen. The reaction was then heated to a required temperature for different times under constant stirring (1000 rpm). After the completion of the reaction, the autoclave was immersed in a water bath to cool down, and the oxygen was expelled from the stainless-steel autoclave at the same time. Finally, the sample was immediately syringed out, filtered and analyzed by high- performance liquid chromatography (HPLC, Agilent 1260 series) with a UV detector.

Conflicts of interest

- The authors declare no conflict of interest.
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