

Efficient conversion of hemicellulose into high-value product and electric power by enzyme-engineered bacterial consortia

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- 2 enzyme-engineered bacterial consortia
- 3 Bo Liang¹, Jing Yang¹, Chen-Fei Meng¹, Ya-Ru Zhang², Lu Wang¹, Li Zhang¹, Jia Liu¹,
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23 Abstract:

As an abundant agricultural and forestry biomass resource, hemicelluloses are hard to be effectively degraded and utilized by microorganisms due to the constraints of membrane and metabolic regulations. Herein, we report a synthetic extracellular metabolic pathway with hemicellulose-degrading-enzymes controllably displayed on Escherichia coli surface as engineered bacterial consortia members for efficient utilization of xylan, the most abundant component in hemicellulose. Further, we develop a hemicellulose/O₂ microbial fuel cell (MFC) configuring of enzyme-engineered bacterial consortia based bioanode and bacterial-displayed laccase based biocathode. The optimized MFC exhibited an open-circuit voltage of 0.71 V and a maximum power density (P_{max}) of 174.33 ± 4.56 μ W cm⁻². Meanwhile, 46.6% (w/w) a-ketoglutarate was produced in this hemicellulose fed-MFC. Besides, the MFC retained over 95% of the P_{max} during 6 days' operation. Therefore, this work establishes an effective and sustainable one-pot process for catalyzing renewable biomass into high-value product and powerful electricity in an environmentally-friendly way.

45 Introduction

Lignocellulose, which is produced through photosynthesis by plant, is a huge solar energy 46 47 reservoir. As a green and sustainable alternative to fossil energy, it has attracted a broad interest to explore lignocellulose biomass for high added-value products and biofuel^{1, 2}. 48 49 Biological hydrolysis and fermentation process of lignocellulosic biorefinery are regarded as the most promise and eco-friendly approach compared to physical and chemical methods^{3, 4}. 50 As the most abundant component in lignocellulose, cellulose can be easily metabolized by 51 most microorganism owing to only glucose units⁵. Whereas, hemicellulose, accounting for 52 20-40% of lignocellulose, are hard to be effectively degraded and utilized by many 53 microorganisms due to the heterogeneous structure of hemicellulose and the constraints of 54 effective membrane transporters as well as the metabolic regulations *in vivo*^{6,7}. Thus, the low 55 utilization efficiency of hemicellulose greatly affect lignocellulosic biorefinery⁸. 56

Currently, hemicelluloses have been used to produce valuable chemicals such as ethanol, 57 furfural and xylitol⁹. Unfortunately, these processes still experience challenges of low 58 59 conversion activity and poor stability of cells. For example, the yield of hemicelluloses-bioethanol is lower than that of bioethanol fermented from cellulose owing to 60 61 the poor utilization efficiency of polysaccharide and pentose by microorganisms during fermentations¹⁰. Besides, bioelectric energy has been heralded as a one of the most potential 62 renewable energies in the future^{11, 12}. To realize the generation of energy from hemicelluloses 63 biomass, great efforts via recruitment of microbial community (microbial fuel cells, MFCs) 64 have been devoted¹³. However, diverse biological properties in one system result in the 65 instability in bacterial community, which severely limits its large-scale application¹⁴. To 66

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address these challenges, a strategy is highly desirable to realize high-efficient production of biofuels and biochemicals directly from hemicellulose instead of its monosaccharides.

69 As an efficient and green method, enzymatic catalysis has been employed to hydrolyze polysaccharides and oxidize monosaccharides. However, the high-cost and poor stability 70 greatly limit their large-scale applications¹⁵. Meanwhile, microbial fermentation is another 71 72 main biological approach to produce biofuels and biochemicals. The cell membrane barrier 73 and intracellular metabolic regulation always restrict the highly efficient substrate utilization, especially polysaccharides¹⁶. Enzymes can be displayed on microbial surface and the resulted 74 whole-cell biocatalyst was obtained by large-scale fermentation and directly applied in 75 biocatalytic reactions without tedious protein purification process, which enables to 76 77 implement complex biochemical reactions even metabolic pathways without the constraints of cell membrane and metabolic regulation^{17, 18, 19}. Similar with secreted proteins, displayed 78 enzymes could be expressed and folded in periplasm and exported directly across cell membrane 79 of E. coli to the culture medium using Sec secretory system²⁰. The displayed enzymes could be 80 81 assembled and anchored onto outer membrane by fusing with anchoring motif, which could 82 facilitate their abilities to be recycled and regenerated. In contrast, secreted proteins will be folded 83 in cytoplasm or periplasm, depending on the sec or tat signal sequence used, and then secreted and dispersed in the culture media without geographical restrictions of cell surface, failing to be 84 recycled or regenerated²¹. 85

Herein, we report on an artificial hemicellulose degrading pathway catalyzed by the engineered bacterial consortia whose enzymes from various origins are recruited to achieve the optimal overall reaction rate in a controlled manner, for the high-efficiently converting

renewable biomass into electric energy and high-value chemical simultaneously in an 89 environmentally-friendly way. The engineered Escherichia coli (E. coli) (up-stream pathway 90 members) degrade xylan, the most abundant biopolymer in hemicellulose²², into 91 monosaccharides, while the following recombinant E. coli (down-stream pathway members) 92 93 oxidize monosaccharides into α -ketoglutarate accompanied by generating the reduced form of nicotinamide adenine dinucleotide (NADH) (Fig. 1). Thus denovo MFC is designed to 94 achieve one-pot efficient production of high added-value chemical of α -ketoglutarate and 95 powerful electricity from hemicellulose biomass in a "one-stone-two-birds" manner, 96 97 demonstrating a model to efficiently utilize biomass in a sustainable way.

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99 **Results and discussion**

Bacterial surface displaying enzymes (engineered bacterial consortia) involving in the saccharification of xylan (up-stream pathway)

102 Herein, the utilization of xylan experienced two stages, firstly saccharified into 103 monosaccharides (mainly D-xylose), which are then oxidized to produce α -ketoglutarate and release electrons. For the initial phase, hemicellulases hydrolyze β-linkage in the xylan 104 105 backbone to release monosaccharides. A marine symbiont Teredinibacter turnerae was selected as the bacterial endo-β-1,4 xylanases (TtGH8) source. TtGH8 shows a wide variety 106 of glycoside hydrolases activities including β -1,4 xylanases, especially the highest activity on 107 mixed-linkage β -1,3 and β -1,4 xylanases²³, which could expand the panel of substrate used in 108 our system. B-D-xylosidase (SXA) from Selenominas ruminantium exhibits good 109 thermo-stability as well as superior activity towards 1,4-β-D-xylooligosaccharides^{24, 25}. 110 Besides, SXA also has α -arabinofuranosidase activity towards arabino xylanases²⁶. The 111

synergy of these two hemicellulases could break down xylan into monosaccharides.
Unfortunately, *E. coli* BL21 (DE3) can metabolize xylose under the action of xylose
isomerase (XylA) and xylulose kinase (XylB)²⁷. Therefore, in order to drive all of the
generated xylose by hemicellulase into the designed extracellular metabolic pathway and
avoid monosaccharides waste, *xylA* and *xylB* genes in *E. coli* BL21 (DE3) were knocked out.

During the first stage, when applying two types of hydrolase-displaying on the surface of 117 bacterial cell, polysaccharide generated from lignocellulose pretreatment could be 118 immediately degraded into monosaccharide by these displayed enzymes in a high efficient 119 way without passing through cell membrane²⁸. TtGH8 and SXA were displayed on cell 120 surface separately or simultaneously as fusion protein using N-terminal region of ice nuclear 121 protein from *P. borealis* (InaPb) as anchoring motif²⁹. The connection manner of cell surface 122 123 displayed TtGH8-SXA was optimized by introducing linkers Gly-Ser, Gly-Ser-Gly-Gly-Ser-Gly and (Ala-Pro)7 between TtGH8 and SXA, respectively. Results 124 show that the engineered strains harboring GS and GSGGSG linkers had similar activities. 125 126 However, (AP)₇ linker influenced the functions of cell surface displayed TtGH8-SXA, and the whole-cell activity reduced 50% compared to that of strain displayed TtGH8-SXA with GS 127 128 linker. Therefore, Gly-Ser was employed as the linker between TtGH8 and SXA. Surface displaying-enzymes demonstrated obvious expression levels, which were confirmed by 129 Western Blotting (Supplementary Fig. 1) and confocal imaging (Supplementary Fig. 2). Next, 130 biochemical activity assays were conducted to validate biological functions of 131 surface-displayed enzymes by using commercial xylan as substrate using 3,5-dinitrosalicylic 132 acid (DNS) method. The optimized protein expression conditions for displayed enzymes were 133

0.5 mM isopropyl-β-D-thiogalactopyranoside (IPTG) and 30 °C (Supplementary Fig. 3). 134 However, when IPTG concentration was over 1.0 mM, the whole-cell activities of engineered 135 strains were declined due to the imbalance of transcription and secretion³⁰. According to 136 results of Quantitative Immunoassay (Supplementary Fig. 4), the number of cell surface 137 displayed proteins were estimated. Strains displayed different amounts of these enzymes, 138 approximately 25560, 25300 and 18020 enzyme molecules for TtGH8, SXA and TtGH8-SXA, 139 respectively. The fewer number of TtGH8-SXA displayed on the surface of per cell suggests 140 141 the decreased display efficiency for protein with a larger molecule (108 kDa). On the basis of 142 the number of enzymes displayed on cell surface, the effects of single display system and dual-display system on saccharification of hemicellulose were investigated. As can be seen 143 144 from Supplementary Fig. 5, the strain displaying fusion protein TtGH8-SXA exhibited a 145 30.37% lower activity towards xylan than the mixture of two strains displaying TtGH8 and SXA individually with molar ratio of 1:1 at the same levels of displayed enzyme molecules, 146 probably due to the lower protein numbers of TtGH8-SXA on cell surface compared to the 147 148 sum of TtGH8 and SXA single-displayed numbers. Then, the ratio of cell density of two engineered strains was optimized to efficiently hydrolyze xylan into monosaccharides. As 149 150 indicated in Fig. 2a, different ratios of strain E. coli-TtGH8 to E. coli-SXA resulted in different hydrolysis efficiency with the best ratio of 3:7. TtGH8 and SXA possess different 151 functions during the degradation process of xylan. TtGH8 is responsible for degrading xylan 152 into xylooligosaccharide as well as a small amount of D-xylose and L-arabinose²³. The 153 xylooligosaccharide could be further hydrolyzed into D-xylose and L-arabinose under the 154 action of SXA. So, the amounts of required SXA were higher than those of TtGH8 for full 155

hydrolyzation of xylan into monosaccharides. The hydrolysis efficiency of up-stream 156 bacterial consortia including E. coli-TtGH8 and E. coli-SXA towards xylan at different 157 concentrations were examined. To quantify the proportion and amounts of pentose in the 158 hydrolysate after strains' treatment, the resultant D-xylose and L-arabinose were determined 159 160 by HPLC method, accounting for 97.43% and 2.57%, respectively. Although the efficiency of saccharification was increased with the decreasing xylan concentration from 1 g/L to 0.1 g/L, 161 the yields of pentose monomers also reduced accordingly (Supplementary Fig. 6). When 162 corncob xylan was 1 g/L, the hydrolysis efficiency was 65.43% (w/w) after 6 h reaction 163 (Supplementary Fig. 6), which was about 1.42-fold higher than in vitro process³¹ and 164 1.65-fold higher than *in vivo* process reported previously³². When the concentration of xylan 165 continued to increase, the production of pentose failed to rise proportionally. Anyway, this 166 167 simple bacterial consortium could generate soluble monosaccharides from xylan ready for the subsequent oxidative degradation. 168

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170 Engineered bacterial consortia involving in oxidation of pentose monosaccharides 171 (down-stream pathway)

To date, D-xylose metabolism pathway has been found in a few microorganisms, which harbors three catabolic routes, including the Weimberg or Dahms pathway, the xylulose-1-phosphate or ribulose-1-phosphate pathway and the xylose isomerase or xylose reductase-xylitol dehydrogenase pathway³³. *E. coli* can utilize xylose as a carbon source for growth through the native route mediated by XylA and XylB as well as pentose phosphate pathway and the glycolysis. However, the low efficiency limited the application of this pathway in metabolic engineering³⁴. Weimberg route in *Caulobacter crescentus* involves 179 conversion of D-xylose to α -ketoglutarate by five steps successively catalyzed by XDH, XylC, 180 XylD, XylX and KGSADH³⁵ (Fig. 1). This pathway has been proved to be an attractive route 181 for biosynthesis of various chemicals from xylose^{34, 36, 37}. In our current study, the oxidation of 182 pentose and the transferring of electrons could be realized by employing this efficient 183 pathway, during which 1 molecule of α -ketoglutarate and 4 electrons per pentose unit can be 184 generated.

To identify the biological activity of each enzyme involved in Weimberg pathway, the in 185 vitro activities of the purified enzymes were measured. Proteins expression conditions were 186 optimized (Supplementary Fig. 7) and enzymes were purified to homogeneity through Ni²⁺ 187 column affinity chromatography before SDS-PAGE analysis (Supplementary Fig. 8). In this 188 first step of Weimberg pathway, D-xylose is oxidized into D-xylono-lactone and generates 189 190 NADH catalyzed by XDH using NAD⁺ as coenzyme. So, the activity of XDH was monitored by measuring the absorbance at 340 nm, typical absorption peak of NADH. As listed in 191 Supplementary Table 3, the enzymatic activity of XDH was 1195.00 ± 8.60 U/mg. To 192 193 examine the activity of XylC, D-xylono-lactone was used as substrate to produce D-xylonic acid. The enzymatic activity of XylC was 146.93 ± 2.16 U/mg, which was 7-fold lower than 194 195 that of XDH (Supplementary Table 3). Subsequently, D-xylonic acid would be transformed into 2-keto-3-deoxyxylic acid by XylD with an activity of 77.00 ± 3.00 U/mg. The activity of 196 KGSADH is 54.00 ± 6.00 U/mg, which was also monitored by measuring the generation of 197 NADH using analogous glutaraldehyde as substrate. Finally, the enzymatic activity of XylX 198 was determined only 35.65 ± 0.25 U/mg (Supplementary Table 3) by a coupled assay with 199 XylD and KGSADH. Thus XylX possessed the lowest activity among these five enzymes 200

involved in Weimberg pathway. Therefore, all of the enzymes had biological functions, but
the conversion of 2-keto-3-deoxyxylic acid to 2,5-dioxopentanoate catalyzed by XylX may be
a limiting step in the overall route, as proposed in other literatures³⁷.

On the basis of the *in vitro* analysis of these five members in Weimberg pathway, 204 205 cell-surface display systems were constructed. To confirm that the localization of enzymes on the surface of E. coli using InaPb anchoring motifs, Western-Blot analysis of outer membrane 206 fraction and immunofluorescent labeling of cells were conducted. All the distinct bands of 207 expected sizes from outer membrane fractions were found (Supplementary Fig. 9), confirming 208 209 that the introduced genes in different recombinant plasmids were expressed obviously. Compared with the control strain, green fluorescence was visualized for strains samples by 210 211 confocal microscope (Supplementary Fig. 10). To ascertain the displayed enzymes possessed 212 functions on the surface of bacteria, the whole-cell was regarded as the catalyst in the enzymatic activity assay. After optimizing the protein expression conditions for displayed 213 enzymes (Supplementary Fig. 11), the whole-cell activities were determined. As depicted in 214 215 Tables S3 and S4, XDH-displaying E. coli exhibits the highest activity of 1.25 ± 0.04 U/OD₆₀₀ with k_{cat} value of 11.34 \pm 1.21 s⁻¹. The whole cell activity of *E. coli*-XylC was significantly 216 217 lower than that of E. coli-XDH. So, the co-display of XDH and XylC would lead to the imbalance of cascade reaction. Therefore, the maximum cascade reaction rate could be 218 achieved by separately displaying XDH and XylC on cell surface and regulating the ratio of 219 strain E. coli-XDH and E. coli-XylC. On the contrary, XylX-displaying E. coli shows the 220 lowest activity (0.044 \pm 0.003 U/OD₆₀₀) with k_{cat} value of 0.07 \pm 0.006 s⁻¹, which corresponds 221 to the in vitro data. These results reveal that XylX was the rate-limiting enzyme in the whole 222

pathway, so the highest overall reaction rate could be realized by adjusting the level of *E*. *coli*-XylX in engineered bacterial consortia.

225 Considering the imbalance of activities between E. coli displaying enzymes, the dual-display and tri-display systems were developed to accelerate the overall reaction rate of 226 227 the latter half of the pathway. The Western Blotting analysis and confocal imaging micrographs indicate the successful expression and display of fusion enzymes on cell surface 228 (Supplementary Fig. 9 and Supplementary Fig. 10). Unfortunately, the degradation of triple 229 230 fusion protein XylX-XylD-KGSADH was observed, implying the unstable presentation of 231 much larger protein (157 kDa) on cell surface. The overall reaction rates from D-xylonic acid to α-ketoglutarate were varied using different display systems as biocatalysts. Among them, 232 233 the bacterial consortia containing XylD-KGSADH-displaying E. coli (E. coli-XylDK) and 234 XylX-displaying E. coli (E. coli-XylX) exhibited the best catalytic efficiency when using D-xylonic acid as substrate and NADH as indicator (Fig. 2b). Then, the cell density ratios of 235 down-stream pathway were optimized to realize the maximum overall reaction rate from 236 237 D-xylose to α -ketoglutarate. Considering the highest activity of E. coli-XDH in the five whole-cell catalysts, the ratio of E. coli-XDH in the prepared bacterial consortia was 238 controlled at a lower level. The ratios of other three strains, including E. coli-XylC, E. 239 coli-XylDK and E. coli-XylX, were varied to realize the highest overall reaction rate from 240 D-xylose to a-ketoglutarate. As shown in Fig. 2c, the ratios of E. coli-XylDK and E. 241 *coli*-XylX had obvious synergistic effect on the generation of the final product α -ketoglutarate. 242 243 Thus, the Weimberg pathway can function in the engineered bacterial consortia.

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246 The integration of the engineered bacterial consortia for producing α-ketoglutarate

To integrate the entire pathway for degrading xylan, according to above optimized 247 248 enzyme-displaying strain ratios in up-stream pathway and down-stream pathway, these involved enzyme-displaying strains were mixed into bacterial consortia with different ratios 249 250 of cell densities in the saccharification of xylan into pentose and the oxidation pentose into 251 α -ketoglutarate. The most appropriate ratio was 3:7 for bacterial consortia in up-stream route to down-stream route with the total OD₆₀₀ of 10.0 taking the titer of α -ketoglutarate as 252 253 evaluation criterion (Fig. 2d). These results demonstrate that the imbalanced enzymes 254 activities could be adjusted by optimizing strain ratios.

α-Ketoglutarate is not only an important intermediate metabolite in cells, but also is 255 widely used as antioxidant and nutrient supplements in food and pharmaceutical fields³⁸. Due 256 to the low efficiency and complex process of chemical synthetic route³⁹, α -ketoglutarate is 10 257 times more expensive than other common organic acids such as citric acid and lactic acid. 258 Moreover, the usage of toxic chemical reagent cyanide in chemical reaction process has 259 260 aroused the concerns of environmental pollution. The microbial synthesis method is characterized as cost-effective, high efficient and eco-friendly⁴⁰. Until now, Yarrowia 261 262 lipolytica and Corvnebacterium glutamicum have been engineered as microbial cell factories to produce α -ketoglutarate using various substrates, such as raw glycerol and xylose (Table 1). 263 Compared with these substrates, hemicellulose is considered to be the most promising 264 feedstock due to its global availability and cost-effective benefits. Herein, to the best of our 265 266 knowledge, the employment of renewable hemicellulose to produce α -ketoglutarate was realized for the first time with the yield of 47% (g/g) within 6 h in vitro one-pot reaction by 267

the engineered bacterial consortia saccharification and oxidation pathway. Our yield reaches 268 the highest level within 6 h, which is significantly shorter than over 90 h for microbial cell 269 270 factories using cost-ineffective glycerol and xylose as substrates (Table 1). It is worth noting that, as the pivot metabolites among tricarboxylic acid cycle (TCA cycle), glyoxylate cycle 271 272 and amino acid metabolism, the generation of α -ketoglutarate would be impacted by complex metabolic regulation in vivo. In this study, extracellular pathway circumvented this bottleneck 273 and improved the synthetic efficiency. Thus, the bioproduction of α -ketoglutarate from 274 hemicellulose in this work realized the efficient conversion of agricultural and forestry 275 276 residues to high-added value compound.

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The integration of the engineered bacterial consortia for generating electricity in MFC 278 279 Fig. 3 describes the mechanism of the two-compartment xylan/O2 MFC separated by a Nafion® membrane. The mediator-less anodic compartment containing multi-walled carbon 280 nanotubes (MWCNTs) covered carbon cloth (CC) electrode, which was modified with 281 282 engineered bacterial consortia (CC/MWCNTs/bacterial-consortia) and fuel of xylan, while the 283 cathodic compartment contained MWCNTs-coated CC electrode, which was modified with E. 2,2'-azino-bis *coli*-displayed laccase (Lac) (CC/MWCNTs/*E.coli*-Lac) 284 and (3-ethylbenzothiazoline-6-sulfonic acid) (ABTS) as the redox mediator. Lac was confirmed to 285 display on the cell surface by SDS-PAGE, Western-Blot and confocal imaging 286 (Supplementary Fig. 12) with good catalytic activity (Supplementary Fig. 13), which 287 catalyzes the O₂ reduction at the cathode (Fig. 4a). *E.coli*-Lac exhibits the maximum activity 288 at pH 5.0 (Supplementary Fig. 13c). The synthetic metabolic pathway on the anode 289

- 290 compartment was designed to degrade xylan into α-ketoglutarate, generating 4 electrons per
- 291 pentose unit. The reactions can be expressed as:
- 292 Anode: Xylan $\xrightarrow{\text{engineered bacterial consortia}} \alpha$ -ketoglutarate + 4e⁻ + 4H⁺ (1)
- 293 Cathode: $O_2 + 4e^- + 4H^+ \rightarrow 2H_2O(2)$
- 294 Overall: Xylan + $O_2 \rightarrow \alpha$ -ketoglutarate + 2H₂O (3)
- 295 The performance of the biocathode and bioanode was examined by Cyclic voltammetry (CV). At the CC/MWCNTs/E.coli-Lac, ABTS exhibited a redox pair in the absence of oxygen 296 (red line), while the cathodic specific current significantly enhanced in the presence of O₂ 297 (Fig. 4a, light blue line), indicating that with ABTS as the electron transfer mediator, the 298 bacteria-displayed laccase catalyzes the reduction of oxygen. For bioanode, as shown in Fig. 299 4b, an onset potential at 0.13 V (vs. Ag/AgCl) for enzyme-displayed E. coli was observed (red 300 301 line). Interestingly, the onset potential for CC/MWCNTs/bacterial-consortia was negatively 302 shifted to 0 V (blue line), suggesting that the generated NADH from NAD⁺ was catalytically oxidized by MWCNTs⁴³. 9,10-anthraquinone-2,7-disulfonic acid (AQDS) was used as 303 electron mediator to investigate the impact of electron transfer mechanisms on the 304 performance of bioanode. As shown in Fig. 4b, the two reduction peaks (green curves) at -0.6 305 V and -0.9 V were the reduction peaks of AQDS with MWCNTs modified CC as electrode. 306 307 The oxidation peak of AQDS was around -0.5 V on the purple curve, which was shifted to around -0.4 V due to the influence of CC modified with bacterial consortia. The oxidizing 308 309 redox peak around -0.1 V on the purple curve was the oxidation peak of NADH produced by 310 the catalysis of bacterial consortia in the presence of AQDS. AQDS can lower the onset 311 potential of NADH which was produced by cell-surface displayed enzymes' catalysis. The 312 onset potential was negatively shifted to -0.20 V (purple line). Therefore, this mediator was

implemented in the electrolyte of anodic chamber in the following studies.

314 Then, we studied the performance of MFCs composed of the above prepared bioanode and biocathode by varying the way of loading engineered strains. The current density (j_{max}) of 315 the generated MFC was $371.67 \pm 10.36 \,\mu\text{A}\cdot\text{cm}^{-2}$ when bacterial consortia were immobilized 316 317 on electrode (Fig. 4c, light green line). However, the j_{max} of MFC dropped to 297.54 \pm 10.24 μ A·cm⁻² when bacterial consortia were dispersed in bioanode chamber (Fig. 4c, blue line). 318 Then the effects of different concentrations of AQDS on the performance of xylan/O₂ MFC 319 320 were investigated. The results show that the highest value of power was achieved when 10 321 mM of AQDS was used (Fig. 4c, peach pink line). The similar trends were observed in Fig. 4d, and the constructed MFC reached the maximum power output (P_{max}) of 68.25 ± 2.38 322 μ W·cm⁻² when bacterial consortia immobilized on bioanode in the presence of 10 mM of 323 324 mediator AQDS.

Shewanella oneidensis is one of the most well-known electricigen in nature⁴⁴. The 325 attempt to engineer S. oneidensis MR-1 using D-xylose as fuel to generate bioelectricity in 326 327 MFC by introducing D-xylose metabolic route shows restricted power owing to poor efficiency of this exogenous pathway⁴⁵. Herein, we reconstituted a D-xylose oxidative 328 pathway and electron transfer chain on cell surface by displaying Weimberg pathway 329 members. There are two steps can produce NADH for generating power, which were 330 catalyzed by XDH and KGSADH. Finally, the oxidation of one D-xylose can yield two 331 NADH and four electrons during the whole process, and the final metabolite α-ketoglutarate 332 333 can be used as the key precursor of several medicine and nutrient substance in food as well as feed⁴⁶. 334

335 The optimization of MFC performance

To boost the power output of MFC, effects of several parameters on electricity generation 336 337 were investigated. First, the loading amounts of bacterial consortia on the bioanode was optimized. The i_{max} of MFCs was 671.33 ± 26.21 μ A·cm⁻², corresponding P_{max} of 112.87 ± 338 7.83 μ W·cm⁻² when loading 10 OD₆₀₀ bacterial consortia (Fig. 5a, peach pink line). The 339 results indicate that the bacterial cell loading exerted effects on power output since the 340 insulated nature of concentrated cells would cause ohmic losses⁴⁷. Then, MFCs performance 341 342 fueled by commercial xylan from corncob was tested under the optimal conditions established 343 according to above experiments. As shown in Fig. 5b, when the concentrations of corncob xylan increased from 0.5 g/L to 1 g/L, the P_{max} boosted from 111.82 ± 5.35 μ W·cm⁻² (green 344 line) to $174.33 \pm 4.56 \,\mu\text{W}\cdot\text{cm}^{-2}$ (peach pink line). However, further increase in concentrations 345 346 of xylan resulted in 12.22% decrease in power density (blue line), probably due to the poor solubility and high viscosities of commercial xylan prepared in buffer solution. The P_{max} of 347 our MFC fueled by 1 g/L xylan is $174.33 \pm 4.56 \ \mu\text{W} \cdot \text{cm}^{-2}$, which is significantly higher than 348 those for xylan-fueled MFCs inoculated with activated sludge (0.609 μ W·cm⁻²) or rumen 349 microorganisms $(40.5 \ \mu W \cdot cm^{-2})^{13, 48}$. 350

In order to investigate the release and transfer of electrons in bioelectrochemical system, Faraday efficiency (η_F) was determined (Supplementary Fig. 14). The current peaked at 2.07 mA at the onset time and then declined over time. In a batch reaction of 35 h, the cumulative electric charges generated from 50 mL of reaction solution were 58.34 C. The concentration of product α -ketoglutarate determined by HPLC was 3.19 mM, convertible to the yield of 46.6%, suggesting that the same amount of D-xylose was consumed and corresponding to 61.58 C of electricity. In consideration of a conversion efficiency of 97% from NADH to electrons determined in a previous study⁴⁹, the η_F value was calculated to be 97.7% (58.34÷61.58÷0.97×100%).

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361 Long-term electricity generation of MFCs fueled by hemicellulose fractions of corncob

As xylan is a predominant component of hemicellulose, we next examined the performance of 362 MFCs powered by real hemicellulose. Biomass corncob pretreated by heating in alkaline 363 solution after neutralizing was used as fuels for power generation⁵⁰. 1.06 g/L xylan from the 364 pretreated hydrolysates of corncob was fueled, in which D-xylose was not detectable by 365 HPLC. To supply constant fuels to MFC, the above pretreated hemicellulose was 366 supplemented once a day to roughly maintain the same level. Then we examined the 367 performance of the MFC using enzyme-engineered-bacterial-consortia-modified-bioanode. As 368 369 shown in Fig. 6, the P_{max} of enzyme-bacterial-consortia-bioanode based MFC was 162.73 \pm μ W·cm⁻² (Fig. 6, black line). The remarkable power output of the 3.69 370 enzyme-bacterial-consortia-bioanode based MFC is much excelled over microbial community 371 372 based MFC¹³, possibly because diverse biological properties of various microorganisms in 373 one system may result in the instability in bacterial community. In addition, the long-term generation of electricity was also evaluated. The bacterial-consortia-bioanode based MFC 374 375 maintained more than 95% of the P_{max} after 6 days' operation. On the contrary, the poor stability was observed in other biomass-based sugar-fed enzymatic fuel cells⁵¹. Besides, the 376 conversion of pretreated hemicellulose from corncob to a-ketoglutarate was also monitored 377 378 during 6 days. The yield of α-ketoglutarate was 44.3% during the 6 days' long-term electricity generation of MFC fueled by the pretreated hydrolysates of corncob (Fig. 6, red line). 379

Moreover, the morphology of the bacterial-consortia-bioanode was observed by scanning electron microscope (SEM). After generating power for 6 days, the *E. coli* consortia and MWCNTs were still closely bound to the CC, while the *E. coli* were intact (Supplementary Fig. 15). These results indicate that the cells survived and stayed stable on CC during 6 days' operation. These results suggest that the capabilities of engineered bacterial consortia can efficiently convert hemicellulose biomass into high-value products and electric power.

386

In summary, a synthetic pathway that can generate high value-added chemical of 387 α-ketoglutarate and 4 electrons from hemicellulose in a controllable manner was successfully 388 constructed by integrating six enzyme-displayed strains into a high-efficient bacterial 389 consortia through tuning the expressed enzyme molecules and adjusting ratios of consortia 390 members. The production of α -ketoglutarate with excellent yield of 0.47 g/g was realized 391 392 within 6 h by the enzyme-engineered bacterial consortia surface-displayed saccharification and oxidation pathway of xylan, the most abundant hemicellulose type. Then a 393 two-compartment xylan/O2 MFC was assembled using enzyme-bacterial consortia modified 394 395 bioanodes and a laccase-displayed strain modified biocathode, realizing the direct conversion of biomass into electricity and α -ketoglutarate. The optimized MFC realized 46.6% (w/w) 396 yield of α -ketoglutarate, the highest OCV and the largest P_{max} . The Faraday efficiency was 397 97.7%. Furthermore, this MFC exhibited a considerable power output and long-term stability 398 towards real biomass samples. To the best of our knowledge, this is the first example to use 399 engineered bacteria consortia as biocatalyst for efficient utilization of hemicellulose, 400 401 representing an important step toward further improving conversion of biomass to high 402 value-added chemical and powerful electric energy. Based on these results, we envision that

403 controllably enzyme-engineered bacterial consortia would find a wide range of applications in
404 the fields of metabolic engineering, synthetic biology, enzyme engineering, bioenergy and
405 enzymatic electrosynthesis.

406

407 Methods

408 Strains and chemicals

All strains and plasmids used in this study are listed in Supplementary Table 1. The primers 409 used are listed in Supplementary Table 2. E. coli DH5a was used for gene recombinant 410 411 manipulation and E. coli BL21 (DE3) AxylAB was used as host to express recombinant 412 proteins. Strains of E. coli carrying recombinant plasmids were routinely grown in Luria-Bertani (LB) medium (10 g/L tryptone, 10 g/L NaCl, 5 g/L yeast extract) at 37 °C and 413 200 rpm. Whenever necessary, antibiotic (kanamycin, 50 µg/mL) was added. Taq DNA 414 415 Polymerase and all restriction endonucleases were purchased from Fermentas (St. Leon Rot, Germany). Kits used for molecular cloning were obtained from Sparkjade Biotech Co., Ltd 416 417 (Qingdao, China). Corncob xylan to represent hemicellulose were purchased from Meryer Co. 418 Ltd (Shanghai, China). 4-nitrophenyl-β-D-xylopyranoside (PNPX), ABTS, AQDS, D-xylose, L-arabinose, xylono-1,4-lactone, D-xylonic acid lithium salt and a-ketoglutarate were 419 420 purchased from Sigma-Aldrich (St. Louis, MO, USA). Platinum (Pt) wire, Ag/AgCl electrode and Nafion 117 membrane (N117) were purchased from Incole Union Technology Co. Ltd 421 (Tianjin, China). Multi-walled carbon nanotubes (MWCNTs) were purchased from Macklin 422 Biochem Technology Co. Ltd (Shanghai, China). Corncob was obtained from Corn Research 423 424 Center of Qingdao Agriculture University. Other chemicals were purchased from Solarbio Science & Technology Co., Ltd (Beijing, China). 425

426 Plasmids construction

427 Construction of the knockout plasmids

- 428 For genetic operations into the genome of *E. coli* BL21 (DE3), the pRE112 suicide vector was
- 429 utilized. To construct plasmids pRE112- $\Delta xy lAB$, approximately 862 bp upstream and 857 bp
- 430 downstream fragments of the *xylA* and *xylB* genes were amplified using the upstream primers
- 431 xylAB-Up-F/R and downstream primers xylAB-Down-F/R, respectively, and inserted into the
- 432 Kpn I site of pRE112. The plasmid pRE112- $\Delta xy lAB$ were transferred from *E. coli* χ 7213 to *E.*
- 433 *coli* BL21(DE3) by conjugation. Suitable recombinants were identified by antibiotic
- 434 resistance screening and sucrose reverse screening. The resulting strain *E. coli*- $\Delta xylAB$ was
- 435 identified by PCR via the primers xylAB-1-F/R.

436 Construction of anchoring motif expressing plasmid

- 437 The gene encoding N-terminal ice nucleation protein originating from *Pseudomonas borealis*
- 438 was amplified from vector pTInaPbN-Xdh by PCR using primers INP-F/INP-R, and inserted
- 439 into the *Nco* I/*Nde* I site of pET-28a (+) to generate plasmid pYJ-00.

440 Constructions of hemicellulose hydrolytic enzymes expressing plasmids

441 The gene encoding TtGH8 from Teredinibacter turnerae (GenBank No. CP001614.2), SXA

442 from *Selenominas ruminantium* (GenBank No. WP_026766185) were codon-optimized for *E*.

443 coli and synthesized by BGI Co., Ltd (Shenzhen, China). To construct intracellular protein

- 444 expression vectors pYJ-01 and pYJ-02, TtGH8 and SXA encoding gene and corresponding
- 445 cloning vector gene were amplified from above cloning plasmids and pET-28a by PCR
- 446 method using primers pET-28a-F(TtGH8)/pET-28a-R(TtGH8) & TtGH8-1-F/TtGH8-1-R,
- 447 pET-28a-F (SXA)/pET-28a-R (SXA) & SXA-1-F/SXA-1-R, respectively, and using the

ClonExpress Ultra One Step Cloning Kit (Vazyme Biotechnology, Nanjing, China). To 448 construct intracellular protein expression vectors pYJ-03, pYJ-01 and pYJ-02 vectors were 449 used as PCR templates and pET-28a-TtGH8-F/pET-28a-TtGH8-R & SXA-2-F/SXA-2-R as 450 primers, and two amplified gene fragments were ligated by in-fusion method using the 451 452 ClonExpress Ultra One Step Cloning Kit. For the construction of plasmids overexpressing 453 enzymes displayed on cell surface, TtGH8 and SXA encoding gene and corresponding cloning vector gene were amplified from above cloning plasmids and pYJ-00 by PCR using 454 primers pET-28a-INP-F (TtGH8)/pET-28a-INP-R (TtGH8) & TtGH8-2-F/TtGH8-2-R, 455 456 pET-28a-INP-F(SXA)/pET-28a-INP-R(SXA) & SXA-3-F/SXA-3-R, respectively, and using the ClonExpress Ultra One Step Cloning Kit to generate pYJ-04 and pYJ-05. Plasmid pYJ-06 457 458 constructed using pYJ-04 pYJ-05 PCR templates was and as and 459 pET-28a-INP-TtGH8-F/pET-28a-INP-TtGH8-R & SXA-4-F/SXA-4-R as primers by fusion PCR strategy. 460

461 Construction of pentose oxidative enzymes expressing plasmids

462 To construct intracellular protein expression vectors pYJ-07, pYJ-08, pYJ-09, pYJ-10 and pYJ-11, the genes encoding XDH, XylC, XylD, XylX, KGSADH were amplified from the 463 genomic DNA of *Caulobacter crescentus* NA1000³⁵ and corresponding cloning vector gene 464 amplified pET-28a by PCR 465 was from method using primers pET-28a-F (XDH)/pET-28a-R(XDH) & XDH-1-F/XDH-1-R, pET-28a-F (XylC)/PET-28a-R (XylC) & 466 XylC-1-F/XylC-1-R, pET-28a-F(XylD)/pET-28a-R(XylD) XylD-1-F/XylD-1-R, & 467 pET-28a-F(XylX)/pET-28a-R(XylX) 468 & XylX-1-F/XylX-1-R, pET-28a-F(KGSADH)/pET-28a-R(KGSADH) & KGSADH-1-F/KGSADH-1-R, respectively, 469

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470	and using the ClonExpress Ultra One Step Cloning K	it. To co	onstruct intracellular protein	
471	expression vectors pYJ-12, pYJ-09 and pYJ-11 vec	ctors we	re used as templates and	
472	pET-28a-XylD-F/pET-28a-XylD-R & KGSADH-2-F/F	KGSADH	I-2-R as primers, and two	
473	amplified gene fragments were ligated by in-fusion met	hod (the G	ClonExpress Ultra One Step	
474	Cloning Kit). For the construction of plasmids overex	pressing	enzymes displayed on cell	
475	surface, XDH, XylC, XylD, XylX, KGSADH encodir	ng genes	was amplified from above	
476	cloning plasmids and corresponding cloning vector gene	e was amp	plified from pYJ-00 by PCR	
477	method using primers pET-28a-INP-F(XDH)/pET-28a-	INP-R(X	DH) &XDH-2-F/XDH-2-R,	
478	pET-28a-INP-F(XylC)/pET-28a-INP-R(XylC)	&	XylC-2-F/XylC-2-R,	
479	pET-28a-INP-F(XylD)/pET-28a-INP-R(XylD)	&	XylD-2-F/XylD-2-R,	
480	pET-28a-INP-F(XylX)/pET-28a-INP-R(XylX)	&	XylX-2-F/XylX-2-R,	
481	pET-28a-INP-F(KGSADH)/pET-28a-INP-R(KGSADH)	& K(GSADH-2-F/KGSADH-2-R,	
482	respectively, and using the ClonExpress Ultra One Ste	ep Clonin	ng Kit to generate plasmids	
483	pYJ-13, pYJ-14, pYJ-15, pYJ-16 and pYJ-17. Plasmid p	oYJ-18 w	as constructed using pYJ-15	
484	and pYJ-16 as PCR templates and pET-28a-INP-XylD-I	F(XylX)/p	pET-28a-INP-XylD-R(XylX))
485	& XylX-3-F/XylX-3-R primers by fusion PCR strategy (ClonExp	ress Ultra One Step Cloning	
486	Kit). Plasmid pYJ-19 was constructed using pYJ-15	and pYJ	-17 as PCR templates and	
487	pET-28a-INP-XylD-F(KGSADH)/pET-28a-INP-XylD-F	R(KGSAI	DH) &	
488	KGSADH-3-F/KGSADH-3-R primers by fusion PCR s	trategy (C	ClonExpress Ultra One Step	
489	Cloning Kit). Plasmid pYJ-20 was constructed using p	YJ-19 and	d pYJ-16 as PCR templates	
490	and pET-28a-INP-XylD-KGSADH-F/pET-28a	-INP-Xy	ID-KGSADH-R &	
491	XylX-4-F/XylX-4-R primers by fusion PCR strategy (ClonExpr	ess Ultra One Step Cloning	

492 Kit).

493 Construction of lac-expressing plasmid

494 For the construction of plasmids overexpressing lac- displayed on cell surface, the gene
495 encoding laccase from *Bacillus subtilis* ⁵² was subcloned into pYJ-00 to generate pYJ-21 by
496 BGI Co., Ltd (Shenzhen, China).

497 **Purification of cytoplasmic enzymes**

498 Cells expressing seven intracellular enzymes, including TtGH8, SXA, TtGH8-SXA, XDH,

499 XylC, XylD, XylX, KGSADH and XylDK (strain TtGH8, SXA, TtGH8-SXA, XDH, XylC,

500 XylD, XylX, KGSADH and XylDK), were separately grown overnight in LB medium with kanamycin, and induced with 0.5 mM IPTG when cells had reached an OD₆₀₀ of 0.6. After 501 502 overnight growth at 16 °C, cells were harvested by centrifugation, resuspended in 40 mL 50 503 mM Tris-HCl (pH 7.0) buffer, broken by ultrasonication and centrifuged at 12000 g for 40 min at 4 °C to remove cell debris and unbroken cells. The soluble extract was applied to a 5 504 mL Ni-NTA column that had been equilibrated with 50 mM Tris-HCl (pH 7.0). Then the 505 506 column was washed consecutively with resuspended buffer and several washing buffer (20 507 mM Tris-HCl pH 7.0, 300 mM NaCl, 1 mM β-mercaptoethanol) containing increasing amounts of imidazole (10, 20, 30 and 50 mM). The bound protein was eluted with elution 508 buffer (20 mM Tris-HCl pH 7.0, 300 mM NaCl, 200 mM imidazole, 1 mM 509 β-mercaptoethanol). Protein dialysis was conducted in a buffer (10 mM Tris-HCl pH 7.0 510 containing 100 mM NaCl, 1 mM β-mercaptoethanol) at 4 °C. Protein purity was analyzed by 511 512 sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE), and protein 513 concentration was determined by Bradford assay.

514 Cell-surface display of enzymes and cell-surface location analysis

For displaying enzymes on cell surface of E. coli, recombinant strains were cultured in LB 515 medium to an OD₆₀₀ of 0.6, 0.5 mM of IPTG was used to induce protein expression. For 516 laccase, cells were further cultured overnight at 16 °C, collected by centrifugation and 517 518 incubated in a buffer of Tris-HCl (50 mM, pH 7.0). The other cells were further cultured at 30 °C overnight, which were collected by centrifugation and incubated in Tris-HCl buffer (50 519 mM, pH 7.0). To determine the successful display of enzymes on the cell surface, outer 520 membrane proteins of recombinant strains and negative control strain were isolated⁵³. The 521 522 outer membrane protein fractions of related strains were analyzed by 10% (wt/vol) SDS-PAGE. 523

524 For Western-Blot assay, samples were transferred to PVDF membranes at 100 mA for 10 min. 525 Anti-6×HisTag mouse monoclonal antibody (catalog number D191001, Sangon Biotechnology, Shanghai, China) at 1:1000 was added and incubated for 3 h at room 526 temperature, then rinsed three times with phosphate buffered saline (PBS) with 0.05%(v/v)527 Tween-20 (PBST). It was then incubated with goat anti-mouse immunoglobulin G 528 (IgG)-horseradish peroxidase (HRP) conjugate (catalog number D110068, Sangon 529 530 Biotechnology, Shanghai, China) for 1 h at room temperature, washed three times with PBST and then once with PBS. The conjugation of antigen and antibody was detected with W-TMB 531 chromogenic kit (Sangon Biotechnology, Shanghai, China). 532

For Quantitative Immunoassay, outer membrane proteins of recombinant strains were
transferred to low fluorescent background PVDF membranes. Anti-6×HisTag mouse
monoclonal antibody at 1:1000 was added and incubated for 3 h at room temperature, then

rinsed three times with PBST. It was then incubated with goat anti-mouse IRDye 800CW 536 fluorescent secondary antibodies (catalog number 926-32210, LI-COR, Inc., Lincoln, NE) for 537 1 h at room temperature, and then washed with PBST, air dried, wrapped in aluminum foil, 538 and stored at 4 °C. LI-COR Odyssey CLx imaging system was used to detect bands on 539 540 membrane. The standard curve from known concentrations of purified recombinant XDH run on the same blot was obtained. Specific protein bands are quantified by fluorescent signals 541 (excitation wavelength, 785 nm; emission wavelength, 820 nm) and a linear regression 542 543 equation generated from recombinant XDH to estimate concentrations of proteins.

For fluorescence imaging, the induced cells were washed with PBS and blocked in PBS buffer containing 1% bovine serum protein (BSA) for 30 min at room temperature. Next anti-6xHis tag mouse monoclonal antibody was added (1:100) after blocking, incubated overnight at 4 °C, and incubated for 1 h the next day at room temperature. After three washes with PBS, the cells were incubated with FITC-conjugated Donkey anti-Mouse IgG (1:100) for 2 h at room temperature. The cells after PBS washing were observed with Laser Scanning Confocal Microscope (TCSsp5 II 03040101, Agilent, USA).

551 Enzymatic activity assay

Enzymatic activity was monitored by measuring the optical density at 600 nm (OD₆₀₀) for cells using a Cary-60 UV-VIS spectrophotometer (Agilent Technologies, Inc., Santa Clara, CA). Soluble commercial corncob xylan was prepared by water dissolution. TtGH8 activity was assessed on 1 g/L soluble commercial corncob xylan in 50 mM Tris–HCl buffer pH 7.0 with 10 mM CaCl₂ at 37 °C. The reducing sugars released by TtGH8 hydrolysis reaction was measured by DNS method with xylose as standard⁵⁴. SXA enzymatic activity was detected

558	using pNPX as the substrate at 37 °C. The product <i>p</i> -nitrophenol shows typical peak at 405
559	nm, which can be monitored by a spectrophotometer ²⁶ . Enzymatic activity of XDH was
560	assayed by detecting the generation of NADH at 340 nm and 37 °C using 10 mM D-xylose as
561	substrate as well as 1 mM NAD ⁺ as coenzyme. For XylC, it can accelerate the hydrolysis of
562	xylono-1,4-lactone into D-xylonic acid to generate D-xylonic acid, which can reduce the
563	absorbance at 405 nm of <i>p</i> -nitrophenol. Enzyme activity was quantified by measuring the
564	decrease in the absorbance at 405 nm ⁵⁵ . Spontaneous hydrolysis of xylono-1,4-lactone was
565	analyzed without catalysts. Enzymatic activity of XylD was assayed according to a modified
566	procedure. In a typical assay, the reaction cocktail contained a certain amount of purified or
567	cells and D-xylonic acid. After incubation at 37 °C for a certain period, the samples were
568	mixed with solution containing 1% semi-carbazide hydrochloride and 1.5% sodium acetate.
569	Finally after incubation at 30 °C for 10 min, the 2-keto-3-deoxyxylic acid produced was
570	quantified by detection of the absorbance at 250 nm typical of semicarbazone ³⁷ . Enzymatic
571	activity of XylX was assayed spectrophotometrically in a coupled assay with the
572	corresponding previous dehydratase XylD and KGSADH. ^{37, 56} The assay was performed in 50
573	mM Tris-HCl buffer (pH 7.0) with 10 mM MgCl ₂ containing 20 mM D-xylonic acid and 1
574	mM NAD ⁺ . After the addition of purified enzymes or enzyme-surface displayed strains, the
575	mixture was incubated at 37 °C. The increasing absorbance at 340 nm caused by NADH
576	produced in the reaction was monitored. Glutaraldehyde was used as the substitute substrate
577	to measure the activity of KGSADH by following the rate of NAD ⁺ reduction by measuring
578	the optical density at 340 nm ⁵⁷ . To determine kinetic constants of cell surface-displayed
579	enzymes, different amounts substrate was used to initiate a series of enzymatic assays. The

data were applied to the Michaelis-Menten kinetic model using Graph-Pad Prism 5 software 580 (www.graphpad.com). To determine the optimum pH for the E. coli-Lac, all enzymatic 581 activities of the whole-cell catalyst were measured by adding ABTS to the final concentration 582 of 5 mM at 37 °C by varying the pH values of the buffer solution (50 mM), including HAc-583 NaAc buffer (pH 3-5.5), Na₂HPO₄-NaH₂PO₄ buffer (pH 6) and Tris-HCl buffer (pH 7-9). 584 The oxidation activity of E. coli-Lac towards ABTS was calibrated by measuring the 585 absorbance of the supernatant at 420 nm (ϵ =36 mM⁻¹cm⁻¹, referred to ABTS concentration).

586

Optimization of protein induced expression conditions 587

588 Recombinant strains were cultured in LB liquid medium at 37 °C and 200 rpm. To optimize inducer concentration for overexpressing INP-fused proteins, IPTG at different concentrations 589 of 0.1 mM, 0.5 mM, 1 mM and 1.5 mM were used when cells grew to an OD₆₀₀ value of 0.6. 590 591 Protein expression was induced overnight, and the activities of whole cell catalyst were determined as above described. In order to investigate the effects of induction temperatures on 592 intracellular and displayed proteins, 0.5 mM of IPTG were added in to cultures when cells 593 594 grew to an OD_{600} value of 0.6. Protein expression was induced overnight at different 595 temperatures of 16 °C, 25 °C, 30 °C and 37 °C, and the activities of crude extract or whole cell catalyst was determined as above described. 596

597

Bioproduction of α-ketoglutarate

598 Recombinant strains were cultured in LB medium to an OD₆₀₀ of 0.6, 0.5 mM of IPTG was used to induce protein expression. Cells were further cultured at 30 °C overnight, which were 599 600 collected by centrifugation and incubated in Tris-HCl buffer (50 mM, pH 7.0). To produce α-ketoglutarate, 100 mL of 50 mM Tris-HCl buffer (pH 7.0) with 10 mM MgCl₂, 10 mM 601

602 CaCl₂, 1.0 g/L commercial corncob xylan and 4 mM NAD⁺ were incubated with artificial 603 bacterial consortia (OD₆₀₀=10) in a flask, which was shaken in an incubator at 150 rpm and 604 37 °C for 6 h. The produced α -ketoglutarate was determined by HPLC.

605 **Preparation of bioanode and biocathode**

606 The thickness of CC (CeTech, Taiwan, China) is 0.32 ± 0.02 mm, which was cut into pieces of 1.5×1.5 cm² and washed by sonication in water and anhydrous ethanol, respectively. A 100 607 μ L of poly (acrylic acid)-MWCNTs dispersion (0.089 mg/cm²) was cast and dried in air to 608 acquire modified CC/MWCNTs. For the preparation of bioanode, 100 μ L of the prepared cells 609 610 (OD₆₀₀=5.0) were dropped onto the CC/MWCNTs and dried at room temperature to obtain CC/MWCNTs/cells modified electrodes. Then 50 µL of Nafion solution (0.1 wt%) was 611 612 syringed to the electrode surface to cover the electrode. For the modification of biocathode, 613 100 µL E. coli-Lac aqueous dispersion (OD₆₀₀=10.0) was coated on the CC/MWCNTs, and subsequently, 50 μ L of Nafion solution (0.1 wt%) was dropped onto the surface of the 614 resulting electrode, then dried at room temperature. The thus-prepared electrode was denoted 615 616 as CC/MWCNTs/E.coli-Lac. The onset potential is defined as the potential at which the current or current density goes above 1 μ A·cm⁻². 617

618 Fabrication of MFC

The dual-chamber hemicellulose/O₂ MFC was assembled with anodic compartment containing the artificial bacterial consortia modified CC/MWCNTs/cells and cathodic compartment containing CC/MWCNTs/*E.coli*-Lac, separated by a Nafion 117 proton exchange membrane (DuPont, USA) with a diameter of 1.6 cm. The anodic electrolyte consisted of 100 mM Tris–HCl buffer (pH 7.0) containing 100 mM NaCl, 10 mM MgCl₂, 10

mM CaCl₂, 4 mM NAD⁺ and xylan. 10 mM AQDS was used as mediator when necessary. 624 The cathodic electrolyte consisted of 100 mM Na₂HPO₄-citric acid buffer (pH 5.0) containing 625 626 0.5 mM ABTS. Polarization curves were obtained by performing linear sweep voltammetry (LSV) at the scan rate of 1 mV s⁻¹ and 37 °C. The specific current (I) was recorded in real 627 628 time. The voltage (V) between the anode and the cathode was set as the Y-axis of polarization curves. The output power (P) was derived via the relationship: $P = V \times I$. The specific current 629 and power were normalized to the geometric area of the anode $(1.5 \text{ cm} \times 1.5 \text{ cm} = 2.25 \text{ cm}^2)$ 630 to obtain the current density and power density, respectively. The calculations of current 631 632 density and power density refer to Eq.(1) and Eq.(2), respectively:

- 633 $j_{\text{max}=}$ I/2.25 (1)
- 634 $P_{\text{max}=}$ V ×I/2.25 (2)

635 Faradaic efficiency (η_F) assay

The η_F was determined through amperometry at 0.45 V. The generation of current was monitored during a reaction time in a volume of 50 mL at 37 °C. The reaction system composed of 100 mM NaCl, 10 mM MgCl₂, 10 mM CaCl₂ and 4 mM NAD⁺, 1 g/L commercial corncob xylan in 100 mM Tris–HCl buffer (pH 7.0). The production of a-ketoglutarate was detected by HPLC. The total charge (C) was calculated according to the generated current during the whole time. The η_F was calculated using the equation as follows⁵⁸:

643 $\eta_F = \int I \times dt / (C_{\alpha-KG} \times V \times n \times F)$ (3)

644 where I is the current generated, dt is the time to produce current, $C_{\alpha-KG}$ is the concentration of 645 produced α-ketoglutarate during the whole time, V is the reaction volume, n is the number of electrons generated per D-xylose consumed, and F is Faraday constant=96,485 C per moleelectron.

648 Long-term electricity generation of MFCs powered by pretreated hemicellulose

To obtain the hemicellulose from lignocellulose, biomass sample corncob was pretreated⁵⁰. 649 650 The milled corncob was pretreated by 2.5 M NaOH with a solid to liquid ratio of 1:30 (g/mL) at 115 °C for 1 h. After filtration, the filtrate was neutralized with 1 M acetic acid for use. The 651 amounts of xylan were determined by gradient precipitation with ethanol and then freeze 652 drying⁶. This pretreated hemicellulose was used as fuels to power MFC for monitoring 653 654 electricity and α -ketoglutarate generation. 100 µL of the prepared cells (OD₆₀₀=10.0) was dropped onto the 2 mg/mL of MWCNTs coated CC surface (2.25 cm²) and dried at room 655 temperature. The pretreated corncob was supplemented into anodic electrolyte daily to 656 657 maintain the constant concentration according to that of a-ketoglutarate. LSV was performed every 12 h incubating at 37 °C during 6 days. The produced α-ketoglutarate was determined 658 by HPLC. 659

660 Morphology observation of the bio-nanocomposite modified CC

To determine whether the *E. coli* consortia and MWCNT were successfully attached to the CC surface, scanning electron microscopic images of the modified CC were recorded using JSM-7500F scanning electron microscopy (JEOL, Tokyo, Japan). The CC was first washed three times with PBS, soaked with 2.5% glutaraldehyde, and fixed at 4 °C for 12 h. Next, the fixed CC was cleaned three times with PBS for 5 min each time. Then the cleaned CC was dehydrated with 30%, 50%, 70%, 90% and 100% ethanol for 5 min each time. Then the dehydrated CC was further dehydrated with 30%, 50%, 70%, 90%, 100% tert-butanol/ethanol for 5 min each time. After dehydration, the CC was frozen at -20 °C. Next, the frozen CC was
put into the freeze-drying machine (Songyuan Huaxing Biotechnology Co., Ltd, Beijing,
China) for freeze-drying. Then, the CC was pasted to the copper platform with conductive
glue, followed by gold spraying.

672 Analytical methods

Cell growth was monitored by measuring the optical density at 600 nm (OD₆₀₀) for cells using 673 a Cary-60 UV-VIS spectrophotometer. Concentrations of D-xylose, L-arabinose, 674 a-ketoglutarate were detected via Ultimate 3000 HPLC (ThermoFisher, USA) using an 675 676 Aminex HPX87H column. The mobile phase was 0.05 M H₂SO₄, and the flow rate was 0.6 mL min⁻¹ at a refractive-index detector at 50 °C⁵⁹. Cyclic voltammetry (CV) was performed 677 in a three-electrode configuration with CC/MWCNTs/cells as working electrode, an Ag/AgCl 678 679 reference electrode and Pt wire as auxiliary electrode connecting to a CHI 1000C potentiostat (CH Instrument, Shanghai, China). The electrochemical reactions were performed at 37 °C. 680

681

682 Statistics and reproducibility

Statistical analyses were mainly performed using Microsoft Excel software (version 2021).
Double-tailed t test or one-way ANOVA and a posteriori test were used for variance analysis.
The data were expressed as mean ± standard deviation (SD). Each group included at least
three independent biological samples. Compared to a reference sample, significance was
established with a *P*-value less than 0.05. No statistical method was used to predetermine
sample size. No data were excluded from the analyses. The experiments were not randomized.
The investigators were not blinded to allocation during experiments and outcome assessment.

Data availability									
Data supporting the findings of this work are available within the paper and Supplementary									
Information files. A reporting summary for this Article is available as a Supplementar									
Information file. Source data are provided with this paper.									
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899 Author contributions

900	B.L., S	S.C.,	A.H.L.,	and J.M.Y.	conceived	and	coordinated	the	study	. B.L.,	A.H.L.,	S.C.	and
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- 901 J.M.Y. wrote the article with input from all other co-authors. B.L., J.Y., C.F.M., and L.W.
- 902 constructed the engineered strains, tested enzymatic activity and MFCs. Y.R.Z. and Z.C.L.
- 903 performed electrochemical testing. J.Y. conducted SEM characterizations. L.Z. and J.L.
- 904 performed biomass pretreatment. B.L., J.Y., C.F.M., and L.W. analyzed the experimental data.
- All authors contributed to the writing of the manuscript.

907 Competing interests

- 908 The authors declare no competing interest.

921 Table 1 The production of α-ketoglutarate by microbial cell factories or *in vitro* one-pot

	Organism	Substrate	Time (h)	Yield (g/g)	Reference
	Y. lipolytica H355A (PYC1-IDP1)	glycerol	117	40%	41
	Y. lipolytica WSH-Z06	glycerol	204	47%	39
	C. glutamicum WMB2 _{evo}	xylose	90	0.55%	42
	E. coli consortia	xylan	6	47%	this study
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922 reaction using different substrates.

939 Figure Legends

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motif. The final product is α -ketoglutarate. TtGH8, β -1,4 xylanase; SXA, β -D-xylosidase; 942 943 XDH, D-xylose dehydrogenase; XylC, xylonolactonase; XylD, xylonate dehydratase; XylX, 2-keto-3-deoxy-D-xylonate dehydratase; KGSADH, 2,5-dioxopentanoate dehydrogenase. 944 945 Fig. 2 The optimization of ratios and constituents of engineered bacterial consortia. a 946 Relative production of reducing sugars using corncob xylan (1g/L) as substrate catalyzed by 947 engineered bacterial consortia involving in the saccharification of xylan (up-stream pathway) with different cell density ratios of E. coli-TtGH8 to E. coli-SXA. 1 mL of 50 mM Tris-HCl 948 949 buffer (pH 7.0) with 10 mM CaCl₂ and 1.0 g/L commercial corncob xylan were incubated with artificial bacterial consortia (OD₆₀₀=10) at 37 °C (n = 3 biologically independent 950 951 experiments). The bacterial consortia with the ratio of 3:7 produced the highest level of reducing sugars compared to those with other ratios. b Relative production of NADH 952 catalyzed by different bacterial consortia using D-xylonic acid as substrate. When different 953 fusion proteins of XylD-KGSADH, XylD-XylX and XylD-XylX-KGSADH were separately 954 displayed on cell surface, the resultant strains were named as E. coli-XylDK, E. coli-XylDX 955 and E. coli-XylDXK, respectively. The total OD₆₀₀ of these four systems were the same and 956 the cell density ratios of different strains were the same in one system. 1 mL of 50 mM Tris-957 958 HCl buffer (pH 7.0) with 10 mM MgCl₂, 20 mM D-xylonic acid and 1 mM NAD⁺ were incubated with artificial bacterial consortia (OD₆₀₀=10) at 37 °C (n = 3 biologically 959 independent experiments). Strain E. coli-XylDXK as the control. The bacterial consortia 960 system composed of E. coli-XylDK and E. coli-XylX produced the highest level of NADH 961

Fig. 1 The saccharification and oxidative pathway of xylan catalyzed by bacterial

surface displayed enzymes using N-terminal region of ice nuclear protein as anchoring

962 compared to those with other system. c Relative production of α -ketoglutarate catalyzed by

963 different bacterial consortia involving in oxidation of pentose monosaccharides (down-stream pathway) with various cell density ratios among E. coli-XDH, E. coli-XylC, E. coli-XylDK 964 and E. coli-XylX using D-xylose as substrate. 1 mL of 50 mM Tris-HCl buffer (pH 7.0) with 965 10 mM MgCl₂, 10 mM D-xylose and 1 mM NAD⁺ were incubated with artificial bacterial 966 consortia (OD₆₀₀=10) at 37 °C (n = 3 biologically independent experiments). The bacterial 967 consortia with the ratio of 1:5:20:25 produced the highest level of α -ketoglutarate compared 968 to those with other ratios. **d** Relative production of α -ketoglutarate catalyzed by different 969 970 bacterial consortia with various cell density ratios of up-stream pathway to down-stream 971 using corncob xylan (1g/L) as substrate at 37 °C for 6 h. The up-stream pathway included E. 972 coli-TtGH8 and E. coli-SXA with the optimal ratio of 3:7. The down-stream pathway 973 included E. coli-XDH, E. coli-XylC, E. coli-XylDK and E. coli-XylX with the optimal ratio 974 of 1:5:20:25. The total OD₆₀₀₀=10 of the bacterial consortia was applied. n = 3 biologically 975 independent experiments. The bacterial consortia with the ratio of 3:7 produced the highest level of α -ketoglutarate compared to those with other ratios. Data are presented as mean \pm SD. 976 The statistical significance is determined by a two-sided t test, and ***, **, * indicate 977 *P*<0.001, 0.01, and 0.05, respectively. Source data are provided as a Source Data file. 978

979 Fig. 3 Schematic drawing of electron transfer route and catalytic reactions in the
980 proposed two-compartment xylan/O₂ MFC. The system was composed of the
981 enzyme-engineered bacterial consortia based bioanode and *E. coli*-Lac based biocathode.

Fig. 4 The integration of the enzyme-engineered bacterial consortia for generating electricity in MFC. a CVs of the CC/MWCNTs/*E.coli*-Lac biocathode in 100 mM Na₂HPO₄-citric acid buffer (pH 5.0) under N₂-saturated atmosphere without ABTS (black line), and in the presence of 0.5 mM ABTS under N₂-saturated (red line) and under oxygen-saturated atmosphere (blue line). Scan rate: 10 mV s⁻¹. **b** CVs of CC in electrolyte

solution (black line); CC/bacterial consortia in electrolyte solution containing xylan (1 g/L) 987 and NAD⁺ (4 mM) (red line); CC/MWCNTs/bacterial-consortia in electrolyte solution 988 989 containing xylan (1 g/L) and NAD⁺ (4 mM) (blue line); CC/MWCNTs in electrolyte solution containing AQDS (10 mM) (green line). CC/MWCNTs/bacterial-consortia in electrolyte 990 991 solution containing xylan (1 g/L), NAD⁺ (4 mM) and AQDS (10 mM) (purple line). The electrolyte solution is 100 mM Tris-HCl buffer (pH 7.0) containing 100 mM NaCl, 10 mM 992 MgCl₂ and 10 mM CaCl₂. Scan rate: 10 mV s⁻¹. c Profiles of potential versus current density 993 MFC consisting of CC/MWCNTs/bacterial-consortia bioanode 994 (*j*). and 995 CC/MWCNTs/E.coli-Lac cathode, which contain 10 mM of AQDS in the anodic chamber (peach pink line). MFC consisting of CC/MWCNTs bioanode and CC/MWCNTs/E.coli-Lac 996 cathode, which contained bacterial consortia and 1 mM of AQDS in the anodic chamber (blue 997 998 line). MFC consisting of CC/MWCNTs/bacterial-consortia and CC/MWCNTs/E.coli-Lac biocathode, which contained 1 mM of AQDS in the anodic chamber (light green line). d 999 Profiles of power density dependent on different bioanodes, which are the same as c. Scan 1000 rate: 1 mV s⁻¹. Abbreviation: CC: carbon cloth. Source data are provided as a Source Data 1001 1002 file.

Fig. 5 The optimization of MFC performance. a Effect of loading amounts of bacterial consortia onto the CC/MWCNTs/bacterial-consortia bioanode on the power output. The bioanodes were prepared by dropping 100 μ L of the prepared cells with different OD₆₀₀ values onto the CC/MWCNTs and dried at room temperature. The xylan concentration is 1.0 g/L. The amounts of bacterial consortia were 5 OD (blue line), 10 OD (pink line) and 20 OD (green line), respectively. **b** Effect of concentrations of commercial corncob xylan on the 1010 10.0 OD_{600} . The concentrations of commercial corncob xylan were 0.5 g/L (blue line), 1.0 g/L (pink line) and 2.0 g/L (green line), respectively. Abbreviation: CC: carbon cloth. Source data 1011 are provided as a Source Data file. 1012 Fig. 6 Long-term operation stability of MFC fueled by hemicellulose fractions of 1013 1014 **corncob.** The time-dependent power density curve (black line) and α -ketoglutarate titers (red 1015 line) of enzyme-bacterial-consortia-modified-bioanode based MFC. Electrical outputs (power density) from engineered bacterial consortia were shown. The production of a-ketoglutarate 1016 in MFC was measured every 12 h during 6 days. LSV was recorded every 12 h during 6 days. 1017 1018 The systems absence of pretreated hemicellulose from corncob were used as negative controls 1019 (blue line representing power density curve and green line representing α -ketoglutarate titers). n = 3 biologically independent experiments. Data are presented as mean \pm SD. Source data 1020 1021 are provided as a Source Data file.

power output. The bioanodes were prepared by dropping 100 µL of the prepared cells with

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