Cellulosic hydrocarbons production by engineering dual synthesis pathways in Corynebacterium glutamicum

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Abstract
Background: Lignocellulose provides the only practical carbohydrates feedstock for sustainable bioproduction of hydrocarbons as future alternative of fossil fuels. Production of hydrocarbons from lignocellulose is achieved by a biorefinery process chain including pretreatment to breakdown the crystalline structure for cellulase-catalyzed hydrolysis, detoxification of inhibitory compounds generated during pretreatment, enzymatic hydrolysis to fermentable monosaccharide sugars, and fermentation to hydrocarbon products. The major barriers on fermentative production of hydrocarbons from lignocellulose include two aspects: one is the inherent stress of pretreatment-derived inhibitors on microbial cells, the other is the toxicity of hydrocarbons to cell membranes. The microbial cell factory should be tolerant to both inhibitor stress and hydrocarbons toxicity.

Results: Corynebacterium glutamicum was selected as the starting strain of hydrocarbons synthesis since it is well adapted to lignocellulose hydrolysate environment. The dual hydrocarbon synthesis pathways were constructed in an industrial C. glutamicum S9114 strain. The first pathway was the regular one in microalgae composed of fatty acyl-acyl carrier protein (fatty acyl-ACP) reductase (AAR) and aldehyde deformylating oxygenase (ADO) with fatty acyl-ACP as precursor. The second pathway was the direct decarboxylation of free fatty acid by fatty acid decarboxylase (OleT) using the rich fatty acids from the disruption of the transcriptional regulator fasR gene. The transmembrane transportation of hydrocarbon products was avoided by secretively expressing the fatty acid decarboxylase (OleT) to the extracellular space. The hydrocarbons generation from glucose reached 29.2 mg/L, in which the direct decarboxylation pathway contributed more than 70% of the total hydrocarbons generation, and the AAR–ADO pathway contributed the rest 30%.

Conclusion: The dual hydrocarbon synthesis pathways (OleT and AAR–ADO pathways) were constructed in the inhibitors tolerant C. glutamicum S9114 strain for hydrocarbon production using lignocellulose feedstock as the starting feedstock. When corn stover was used for hydrocarbons production after dry acid pretreatment and biodetoxification, the hydrocarbons generation reached 16.0 mg/L. This study provided a new strategy for hydrocarbons synthesis using microbial cell factory suitable for lignocellulose feedstock.

Keywords: Hydrocarbons, Lignocellulose, Corynebacterium glutamicum, Dual synthesis pathways, Fatty acid decarboxylase OleT, Secretive expression

Background
Current hydrocarbon fuels are dominantly produced from petroleum refining [1, 2]. The coming carbon-neutral economy requires the production of future aviation fuel and diesel in a sustainable way from renewable
resources. Among all potential feedstocks, lignocellulose provides the only practical carbohydrates option for bio-production of biofuels by its abundance and availability [3, 4].

Various microorganisms had been tested and engineered as microbial cell factories for hydrocarbons synthesis [5, 6]. The most acknowledged pathway is the decarboxylation of fatty acyl-ACP to fatty aldehydes by fatty acyl-ACP reductase (AAR) or the reduction of fatty acid by fatty acid reductase (FAR) [1, 7], then the fatty aldehydes are converted to hydrocarbons by aldehyde deforming oxygenase (ADO) or aldehyde decarbonylase (ADS) [8, 9]. An alternative pathway is the direct decarboxylation of free fatty acids to terminal alkenes by fatty acid decarboxylase OleT [10], nonheme iron oxidase UndA [11], or fatty acid desaturase UndB [12] when abundant free fatty acids are available.

Pretreatment generally leads to the partial degradation of hemicellulose (mainly xylan) to furfural and cellulose to 5-hydroxymethyl furfural (HMF), and lignin to various phenolic compounds, along with other weak organic acids [13]. Biodetoxification fungus completely removes the most toxic inhibitors of furfural and HMF, as well as most of acetic acid and phenolics such as p-hydroxybenzaldehyde, vanillin, syringaldehyde, etc. However, the residual phenolics and acetic acid are not removed as completely as furfural and HMF because the extensive biodetoxification leads to the loss of xylose sugars. These residual inhibitors still show observable inhibitions in the cell growth and hydrocarbons synthesis metabolism of microorganisms such as Escherichia coli and most of the hydrocarbon synthesis strains. Furthermore, lipophilic hydrocarbons are toxic to cells due to the interference on cell membranes. The accumulation of hydrocarbons inside the cell increases the permeability and fluidity of the cell membrane and interferes with the function of membrane proteins, which in turn affects energy transfer and the stability of the cell membrane [14]. Previous studies showed that the microorganism C. glutamicum was unusually adaptive to lignocellulose hydrolysate environment by its strong tolerance to residual inhibitors of biodetoxifications and well utilization of rich vitamin B components in lignocellulose [15–20]. In this study, C. glutamicum was selected as the potential microbial cell factory [16, 18].

One of the uniqueness of C. glutamicum is that considerable free fatty acids are generated by C. glutamicum including oleic acid (C18:1), followed by palmitic acid (C16:0), and minor palmitoleic acid (C16:1) and stearic acid (C18:0) by the multi-enzyme complex fatty acid synthases (FASs) (one FAS-I modular and three FAS-II modules NCgI0281, NCgI0283 and NCgI0527) [21, 22]. The free fatty acids are accumulated and well maintained in extracellular space of C. glutamicum due to lack of β-oxidation pathway for fatty acid degradation, and lack of phosphatidic acid phosphatase (PAP) and diacylglycerol acyltransferase (DGAT) to triacylglyceride (microbial lipid) [23]. These advantages further suggest that C. glutamicum might be a favorable candidate of biorefinery hydrocarbons fermentation strains, though the cell membrane toxicity by hydrocarbons still exists.

We engineered an industrial C. glutamicum strain by constructing the dual hydrocarbon synthesis pathways in this study. One pathway was the regular algal pathway by fatty acyl-ACP reductase (AAR) and aldehyde deforming oxygenase (ADO) with fatty acyl-CoA as precursor; the other pathway was the direct decarboxylation of fatty acid by fatty acids decarboxylase (OleT). The toxicity of the intracellularly generated hydrocarbons by transmembrane transportation was lessened by the secretive expression of OleT guided by a signal peptide. To increase the fatty acids substrate supply, the transcriptional regulator fasR gene was disrupted to increase C16 and C18 fatty acids as the additional substrate supply [24]. The resulting engineered C. glutamicum was applied for hydrocarbons fermentation using corn stover as feedstock for hydrocarbons production. This study provided a new strategy of metabolic engineering for hydrocarbons production using lignocellulose feedstock.

Results

Construction of AAR–ADO hydrocarbon synthesis pathway in C. glutamicum

To construct the first AAR–ADO pathway from the fatty acyl-ACP substrate for hydrocarbons synthesis, the fragments of the lipoyl-ACP reductase gene aar (synpcc7942_1594) and the fatty aldehyde decarbonylase gene ado (synpcc7942_1593) from Synechococcus elongatus PCC7942 were synthesized and the expression plasmids were constructed; then the plasmid were introduced to C. glutamicum S9114 (Fig. 1a, Box 1). The two genes aar (synpcc7942_1594) and ado (synpcc7942_1593) were expressed under the control of promoter H36 with different alignments (C. glutamicum ZW1, ZW2, and ZW3 in Table 1). The gene combination of ado and aar with the consensus SD sequence aar-rbs-ado (ZW2) showed the optimal hydrocarbon generation among the gene alignments. Then this gene fragment aar-rbs-ado was integrated into the location of the TetR-type transcriptional regulator fasR to generate a recombinant C. glutamicum HW4. Figure 1b shows that the typical alkene components C12H16, C14H28, C16H32, C18H36, and C20H40, among other hydrocarbon components, were detected in the fermentation broth of C. glutamicum HW4, indicating that the stable AAR–ADO hydrocarbon synthesis pathway had been established in genome scale.
The fasR disruption by the aar-rbs-ado cluster also led to the excessive generation of C\textsubscript{16} and C\textsubscript{18} fatty acids. The fatty acids generation by the parental \textit{C. glutamicum} S9114 was about ~ 10 mg/L fermentation broth by GC–MS detection with dodecane as the standard. The disruption of fasR in \textit{C. glutamicum} HW4 resulted in the fatty acid production.

**Fig. 1** Hydrocarbons synthesis pathway in \textit{Corynebacterium glutamicum} and construction of AAR–ADO pathway. **a** Overall hydrocarbons synthesis pathway in \textit{C. glutamicum} S9114; black lines represent native pathways. Red lines represent overexpressing heterologous genes. Red cross represents the deletion of the gene. ACP acyl carrier protein, AAR acyl-ACP reductase, ADH aldehyde reductase, ADO aldehyde deformylating oxygenase, ACC acetyl CoA carboxylase, FAS fatty acid synthase, FAT fatty acyl-ACP thioesterase, ACS fatty acyl-CoA synthetase, sGDH glucose dehydrogenase, FDH1 formic acid dehydrogenase, AasS fatty acyl-ACP synthetase, OleT fatty acid decarboxylase. **b** GC–MS chromatogram of hydrocarbons production in \textit{C. glutamicum} S9114 (control) and HW4 (with AAR–ADO pathway). Shake-flask fermentation, 30 °C, 200 rpm, pH 7.0 maintained by adding 5 M NaOH. Abundance represented the response intensity of chromatogram peaks. \textit{C. glutamicum} S9114 generated no hydrocarbons thus the peaks (black line) were not visible on the chromatogram; **c** metabolic modifications of \textit{C. glutamicum} for hydrocarbons production chart. Fermentation parameters were similar to **b**.
acid generation up to 103 mg/L, approximately one order of magnitude greater than that by the parental strain (Fig. 1c).

To increase the hydrocarbons production of \textit{C. glutamicum} HW4, a systematic metabolic engineering was performed (Fig. 1a) including (i) separately overexpressing \textit{aar} and \textit{ado} in plasmids pH36mob to increase the copy numbers (Fig. 1a, Box I); (ii) expression of NADPH reductase genes \textit{pntAB}, \textit{udhA} and \textit{sgdh} to increase the NADPH supply (Fig. 1a, Box II); (iii) expression of fatty acyl-ACP synthase AasS from \textit{Vibrio harveyi} B392 to convert free fatty acids to fatty acyl-ACP (Fig. 1a, Box III) [25], and (iv) expression of formic acid dehydrogenase FDH1 from \textit{S. cerevisiae} S288C to degrade the byproduct formic acid (Fig. 1a, Box IV). However, these efforts showed no improvements or even negative results (Fig. 1d), except that the enhancement of NADPH supply by overexpression of \textit{udhA} and \textit{sgdh} genes (encoding UdhA and sGDH, respectively) with ~8% increase of hydrocarbons production.

\textbf{Constructing the fatty acid decarboxylation pathway in \textit{C. glutamicum} and the secretive expression}

To increase the hydrocarbons production by \textit{C. glutamicum}, the second pathway was constructed by direct decarboxylation of free fatty acid. Free fatty acids accumulation is the unique phenomenon of \textit{C. glutamicum} because of the lack of lack of β-oxidation pathway for fatty acid degradation, as well as the lack of phosphatidic acid phosphatase (PAP) and diacylglycerol acyltransferase (DGAT) to produce triacylglyceride (microbial lipid) [21]. The direct decarboxylation pathway of free fatty acids was constructed by overexpression of the \textit{oleT} gene encoding the fatty acid decarboxylase OleT (Fig. 1a, Box V). Two fatty acid decarboxylase genes, \textit{oleT}_{JE} from \textit{Jeotgalicoccus} sp. ATCC 8456 and \textit{oleT}_{MC} from \textit{Macrococcus caseolyticus} WP_041635889.1 [10, 26], were selected, synthesized, and heterologously expressed in \textit{C. glutamicum} S9114. The di-alkenes of \textit{C_{12}H_{22}} and \textit{C_{14}H_{26}}, as well as the mono-alkenes of \textit{C_{15}H_{30}} and \textit{C_{17}H_{34}} were produced by the overexpression of the two \textit{oleT} genes (Fig. 2). The greater hydrocarbons generation of \textit{oleT}_{MC} expression indicates \textit{OleT}_{MC} was more adaptive for fatty acid decarboxylation in \textit{C. glutamicum} S9114.

To avoid the cell membrane damage by the transmembrane transportation of hydrocarbons, the extracellular decarboxylation of fatty acids into hydrocarbons were designed by the secretive expression of OleT (Fig. 3a). Two secretory pathways were tested, one was the Sec pathway to secret the unfolded proteins by the peptide \textit{Ncgl1289} from \textit{C. glutamicum} ATCC13032 [27], the other was the Tat pathway to secret the folded proteins by \textit{cgR}_{0494} from \textit{C. glutamicum} S9114 [28–30]. Each of the signal peptide genes was ligated with \textit{oleT}_{JE} and \textit{oleT}_{MC}. 
### Table 1 Strains and plasmids used

| Strains | Characteristics | Sources |
|---------|-----------------|---------|
| E. coli BL21 | Host for plasmid construction | Lab stock |
| Paecilomyces variotii FN89 | Biodetoxification fungus isolated by our lab | Lab stock |
| C. glutamicum 502-14 | Industrial strain | SIIL |
| C. glutamicum ZW1 | C. glutamicum 502-14 carrying pH36-ado-adr | This study |
| C. glutamicum ZW2 | C. glutamicum 502-14 carrying pH36-ado-rbs-adr | This study |
| C. glutamicum ZW3 | C. glutamicum 502-14 carrying pH36-ado-TacM-adr | This study |
| C. glutamicum HW4 | C. glutamicum 502-14 deleting fasR and carrying aar and ado | This study |
| C. glutamicum HW4-pH36-adr | C. glutamicum 502-14 carrying pH36-adr | This study |
| C. glutamicum HW4-pH36-adr | C. glutamicum 502-14 carrying pH36-adr | This study |
| C. glutamicum HW4-pH36-sgdh | C. glutamicum 502-14 carrying pH36-sgdh | This study |
| C. glutamicum HW4-pH36-asS | C. glutamicum 502-14 carrying pH36-asS | This study |
| C. glutamicum HW4-pH36-fdh1 | C. glutamicum 502-14 carrying pH36-fdh1 | This study |
| C. glutamicum pH36-adr | C. glutamicum 502-14 carrying pH36-adr | This study |
| C. glutamicum pH36-adr | C. glutamicum 502-14 carrying pH36-adr | This study |
| C. glutamicum pH36-sgdh | C. glutamicum 502-14 carrying pH36-sgdh | This study |
| C. glutamicum pH36-asS | C. glutamicum 502-14 carrying pH36-asS | This study |
| C. glutamicum pH36-fdh1 | C. glutamicum 502-14 carrying pH36-fdh1 | This study |
| C. glutamicum pH36-adr | C. glutamicum 502-14 carrying pH36-adr | This study |
| C. glutamicum pH36-adr | C. glutamicum 502-14 carrying pH36-adr | This study |
| C. glutamicum pH36-sgdh | C. glutamicum 502-14 carrying pH36-sgdh | This study |
| C. glutamicum pH36-asS | C. glutamicum 502-14 carrying pH36-asS | This study |
| C. glutamicum pH36-fdh1 | C. glutamicum 502-14 carrying pH36-fdh1 | This study |

**Plasmids**

| Plasmids | Characteristics | Sources |
|----------|-----------------|---------|
| pK18mobsacB | Mobilizable vector in C. glutamicum, kanamycin resistance, sacB | Wang et al. [17] |
| pEFtrumob | Insert promoter Pefu at the back of promoter Ptrc in pTRCmob | Lab stock |
| pH36-ado-adr | Overexpression vector, kanamycin resistance | Lab stock |
| pH36-ado-rbs-adr | pH36mob carrying rbs and ado under H36 control and ado under TacM control | This study |
| pH36-fdh1 | pH36mob carrying fdh1 | This study |
by overlap PCR, then inserted into the expression plasmids and introduced to *C. glutamicum* S9114 to obtain four recombinants, S9114-pH36-NsoleT<sub>IE</sub>, S9114-pH36-RsoleT<sub>IE</sub>, S9114-pH36-NsoleT<sub>MC</sub> and S9114-pH36-RsoleT<sub>MC</sub>. Both the SDS-PAGE and the Western blotting did not show the clear protein bands due to the high fatty acids content for protein extraction in the fermentation broth. Figure 3b shows that the secretive expression of OleT significantly improved the hydrocarbons generation. Among the hydrocarbons produced, the secretive expression of OleT<sub>MC</sub> by the Sec pathway (S9114-pH36-NsoleT<sub>MC</sub>) showed approximately fourfolds greater hydrocarbons (9.6 mg/L) than the intracellular expression (S9114-pH36-OleT<sub>MC</sub>, 2.4 mg/L), and approximately 36% more hydrocarbons than the secretive expression of OleT<sub>MC</sub> by the Tat pathway (S9114-pH36-RsoleT<sub>MC</sub>.

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**Table 1** (continued)

| Characteristics | Sources |
|-----------------|---------|
| pH36-oleT<sub>IE</sub> | pH36mob carrying oleT<sub>IE</sub> | This study |
| pH36-oleT<sub>MC</sub> | pH36mob carrying oleT<sub>MC</sub> | This study |
| pH36-NsoleT<sub>IE</sub> | pH36mob carrying oleT<sub>IE</sub> with signal peptide of Ncgl1289 | This study |
| pH36-RsoleT<sub>IE</sub> | pH36mob carrying oleT<sub>IE</sub> with signal peptide of RS04950 | This study |
| pH36-NsoleT<sub>MC</sub> | pH36mob carrying oleT<sub>MC</sub> with signal peptide of Ncgl1289 | This study |
| pH36-RsoleT<sub>MC</sub> | pH36mob carrying oleT<sub>MC</sub> with signal peptide of RS04950 | This study |
| pH36-Ncgl1221-oleT<sub>MC</sub> | pH36mob carrying oleT<sub>MC</sub> with signal peptide of Ncgl1221 | This study |
| pH36-Ncgl1337-oleT<sub>MC</sub> | pH36mob carrying oleT<sub>MC</sub> with signal peptide of Ncgl1337 | This study |
| pH36-PorB-oleT<sub>MC</sub> | pH36mob carrying oleT<sub>MC</sub> with signal peptide of PorB | This study |
| pH36-PorC-oleT<sub>MC</sub> | pH36mob carrying oleT<sub>MC</sub> with signal peptide of PorC | This study |
| pEftu-Ncgl1221-oleT<sub>MC</sub> | pH36-Ncgl1221-oleT<sub>MC</sub> with H36 promoter replaced by Eftu promoter | This study |
| pEftu-Ncgl1337-oleT<sub>MC</sub> | pH36-Ncgl1337-oleT<sub>MC</sub> with H36 promoter replaced by Eftu promoter | This study |
| pEftu-PorB-oleT<sub>MC</sub> | pH36-PorB-oleT<sub>MC</sub> with H36 promoter replaced by Eftu promoter | This study |
| pEftu-PorC-oleT<sub>MC</sub> | pH36-PorC-oleT<sub>MC</sub> with H36 promoter replaced by Eftu promoter | This study |
| pH36-PorB-oleT<sub>MC</sub> with H36 promoter replaced by Eftu promoter | This study |
| pH36-PorC-oleT<sub>MC</sub> with H36 promoter replaced by Eftu promoter | This study |

**Fig. 2** GC–MS result of hydrocarbon production via different fatty acid decarboxylases in *C. glutamicum* S9114. Shake-flask fermentation at 30 °C, 200 rpm, pH 7.0 maintained by adding 5 M NaOH.
The results suggest that the cell damage was partially relieved by the secretion of decarboxylase and considerable hydrocarbons were generated in the extracellular space.

Cell surface display of the OleT expression was also tried by expressing the anchor proteins NCgl1221, NCgl1337, PorB and PorC from C. glutamicum S9114. Unfortunately, the results show no observable improvement or even suppression by the surface display expression of OleT under the control of the promotor Eftu and H36 (Fig. 4).

**Fig. 4** Hydrocarbons production of displaying fatty acid decarboxylase OleT<sub>MC</sub> on the cell surface under the control of H36 promote and Eftu promoter. Control represents C. glutamicum S9114 with the overexpression of oleT<sub>MC</sub>. Fermentations were carried out in shake flask, 30 °C, 200 rpm, pH 7.0 maintained by adding 5 M NaOH

Dual pathways construction for hydrocarbons synthesis in C. glutamicum

The two secretive overexpression of OleT<sub>MC</sub> by the Sec secretive pathway and the Tat secretive pathway were expressed in C. glutamicum HW4 (carrying the AAR–ADO pathway in genome scale) to give the two recombinants C. glutamicum recombinants, HW5 and HW6, respectively (Fig. 5a).

Figure 5b shows that both the recombinants C. glutamicum HW5 and HW6 significantly enhanced the generation of both the di-alkenes (C<sub>12</sub>H<sub>22</sub> and C<sub>14</sub>H<sub>26</sub>) and the mono-alkenes (C<sub>13</sub>H<sub>30</sub> and C<sub>17</sub>H<sub>34</sub>). The hydrocarbons generation by OleT pathway in C. glutamicum HW5 (Sec secretive pathway) (10.1 mg/L) were 1.9 times greater than the control (without the secretive expression HW4-pH36-OleT<sub>MC</sub> 3.5 mg/L), and 13% higher than that of C. glutamicum HW6 (Tat secretive pathway, 8.9 mg/L). The total hydrocarbons production by the dual pathways of AAR–ADO and the OleT<sub>MC</sub> in C. glutamicum HW5 reached 16.0 mg/L.

The optimal fermentation parameters of C. glutamicum HW5 with the dual hydrocarbons synthesis pathways were examined in bioreactors with automatic pH control and dissolved oxygen input using glucose as carbon resource (Fig. 5c). The results show that the hydrocarbons production was not affected by NADPH addition, and 30 °C and pH 7.0 were suitable for cell growth and hydrocarbons production; the hydrocarbons generation increased with increasing oxygen transfer rate by varying the stirring rate. At the proper fermentation conditions (30 °C, pH 7.0 and 750 rpm), totally 29.2 mg/L of hydrocarbons were produced, including 14.5 mg/L of di-alkenes and 7.3 mg/L of mono-alkenes by C. glutamicum HW5.

Corn stover was used as carbohydrates feedstock for hydrocarbon production by C. glutamicum HW5 (Fig. 6a). Corn stover hydrolysate was prepared by enzymatically hydrolyzing 15% (w/w) of the dry acid pretreated and biodetoxified corn stover with the addition of 5 g/L (NH₄)₂SO₄, 1 g/L KH₂PO₄, 1 g/L K₂HPO₄ and 0.25 g/L MgSO₄. The total hydrocarbons production by the dual pathways in C. glutamicum HW5 reached 10.8 mg/L, in which the OleT<sub>MC</sub> pathway generated 7.8 mg/L and the AAR–ADO pathway generated 3.0 mg/L of hydrocarbons.

The hydrocarbons generated from the two pathways shared different carbon chain lengths. The carbon
number of hydrocarbons from the OleT pathway was less than 20 and the retention time on GC–MS chromatograph was between 7–8 min. The carbon number from the AAR–ADO pathway was greater than 20 and the retention time on GC–MS was greater than 10 min (Fig. 6b). The hydrocarbons generation could be translated to 0.205 mg hydrocarbons/g cellulose in corn stover.

Hydrocarbons production from biomass has been the focus of metabolic engineering and various microorganisms had been engineered as microbial cell factories for hydrocarbons synthesis. This study used C. glutamicum as the hydrocarbons producing strain by its two inherent properties: inhibitors tolerance and free fatty acid generation. The dual hydrocarbons synthesis pathways were constructed and 10.8 mg/L of hydrocarbons was generated when corn stover was used, in which the OleT pathway contributed more than 70% of the total generation of hydrocarbons. The high residual fatty acids level indicated a further upgrading potential of hydrocarbons production by further metabolic engineering.

One of the uncertainties of hydrocarbons production is the less accurate calculation of hydrocarbons content based on the peak areas of hydrocarbons generation of GC–MS chromatogram due to the high and irregular fatty acids peaks on the baseline of GC–MS chromatogram. Due to the high boiling point of fatty acid and non-volatility, the accuracy of hydrocarbons measurement was negatively affected. Highly possible, the hydrocarbons contents were under-estimated because of the merger of hydrocarbons peaks with fatty acids peaks. We tried to measure the fatty acid content using GC–MS and eliminate the fatty acid content from hydrocarbon components for an accurate measurement. However, the high boiling points of fatty acids led to the incomplete volatilization of fatty acids and carbon residues accumulation in GC column (thus damaged the column). We also tried the esterification of the fatty acids in the samples, but this procedure caused a heavy loss of hydrocarbons and the accuracy of hydrocarbon measurement was even worse. A more accurate analysis method is under investigation to determine hydrocarbons contents with the existence of high free fatty acids in the fermentation broth.

Conclusion
The dual hydrocarbon synthesis pathways (OleT and AAR–ADO pathways) were constructed in the inhibitors tolerant C. glutamicum strain for hydrocarbon production using lignocellulose feedstock as the starting feedstock. The first one is the regular AAR–ADO pathway and the second is the fatty acid decarboxylation pathway from fatty acids by taking advantage of free fatty acid generation of C. glutamicum. The fatty acid decarboxylation pathway was further enhanced significantly by secretive expression of fatty acid decarboxylase OleT and performed the extracellular catalysis by secreted OleT enzyme. The hydrocarbons generation from glucose reached 29.2 mg/L, in which the direct decarboxylation pathway contributed more than 70% of the total hydrocarbons generation, and the AAR–ADO pathway contributed the rest 30%. When corn stover was used for hydrocarbons production after dry acid pretreatment and biodetoxification, the hydrocarbons generation reached 16.0 mg/L. This study provided a new strategy for hydrocarbons synthesis using microbial cell factory suitable for lignocellulose feedstock. The high residual fatty acids level indicated a further upgrading potential of hydrocarbons production by further metabolic engineering.

Discussion
Although the major inhibitors (furfural, 5-hydroxymethylfurfural, acetic acid) generated during pretreatment were removed by biodetoxification, the residual phenolic compounds still showed considerable stress on fermentation strains. In this study, we selected a high robust C. glutamicum as microbial cell factory for synthesis of hydrocarbons and the hydrocarbon products were successfully synthesized by the engineered C. glutamicum using corn stover feedstock after dry biorefinery process. To improve the low hydrocarbons synthesis efficiency of C. glutamicum, two hydrocarbon synthesis pathways were constructed, the first hydrocarbon synthesis pathway of AAR–ADO originates from cyanobacteria, and the second fatty acid decarboxylation pathway utilizes the unique and rich free fatty acid substrates.

Due to the knockout of the fask gene, the engineered C. glutamicum accumulated considerably high fatty acids in the extracellular environment. We expressed the fatty acid decarboxylase and secreted the enzyme into

(See figure on next page.)

**Fig. 5**  Hydrocarbons production by dual synthesis pathways in C. glutamicum. a Secretive expression of fatty acid decarboxylase OleT<sub>MC</sub> in C. glutamicum HW4. b Hydrocarbons production by dual synthesis pathways in C. glutamicum. HWS indicates C. glutamicum HW4 with the secretive expression of oleT<sub>MC</sub> through Sec pathway and HW6 indicates C. glutamicum HW4 with the secretive expression of oleT<sub>MC</sub> through Tat pathway. Fermentation was carried out in shake flask, 30 °C, 200 rpm. pH was maintained at 7.0 by adding 5 M NaOH. c Improved hydrocarbons production by optimizing the culture conditions. Fermentation was carried out in a 1-L fermentor. The medium was mentioned in “Materials and methods”. The basic fermentation conditions were 30 °C, pH 7 and 600 rpm. When one of the fermentation parameters was changed for optimization, the others remained the same.
(a) Dual pathways of secretive fatty acid decarboxylase OleT and AAR-ADO

(b) Hydrocarbons production by dual hydrocarbons pathways in flask

(c) Total hydrocarbons production by dual hydrocarbons pathways in fermentor

Fig. 5 (See legend on previous page.)
S9114 was used as the starting strain and the genome of C. glutamicum was engineered to contain the AAR-ADO pathway and the OleT pathway. The genome was constructed by homologous recombination and verified by colony PCR. Escherichia coli BL21 was used for plasmid construction and cultured in LB medium. C. glutamicum S9114 was used as the starting strain and the genome sequence referred in NCBI with the accession number NZ_AFYA01000018. C. glutamicum was cultured in CM2B medium (yeast extract 10 g/L, peptone 10 g/L, NaCl 10 g/L). The CGXII-NL medium for hydrocarbons fermentation contained 60 g/L glucose, 1.0 g/L (NH4)2SO4, 2.5 g/L urea, 1.0 g/L KH2PO4, 1.0 g/L K2HPO4, 42 g/L MgSO4, 0.25 g/L MgSO4, 0.01 g/L CaCl2, 0.01 g/L FeSO4·7H2O, 0.01 g/L MnSO4·H2O, 0.001 g/L ZnSO4·7H2O, 0.0002 g/L CuSO4·5H2O, 0.0002 g/L NiCl2·6H2O, 0.0002 g/L biotin, 0.0005 g/L thiamin, 0.03 g/L PCA. 50 μg/mL of kanamycin was added into the media if needed.

The primers used for plasmids construction are shown in Additional file 1: Table S1. The fragments of aar, ado, sgdh, aasS, fdh1, oleTJE and oleTMC genes were synthesized by Shanghai Generay Biotech, Shanghai, China. The pntAB and udhA genes were amplified from E. coli BL21 genome. These fragments were then constructed into the expression vectors pH36mob and pEftumob separately by digestion-ligation or in-fusion cloning. The fragments of aar-ado, ado-rbs-aar and ado-TacM-aar were obtained by overlapping the corresponding fragments and then inserted into the expression vector pH36mob, resulting in several plasmids pH36-aar-ado, pH36-ado-rbs-aar, and pH36-ado-TacM-aar.

The signal peptide sequences of Ncgl1208 and cgR_0949 were amplified from C. glutamicum ATCC 13032 and C. glutamicum S9114. Then the signal peptide sequences of Ncgl1208 and the fragment of oleTJE gene were overlapped together and inserted into pH36mob, resulting pH36-NsoleTJE. Plasmids pH36-NsoleTMC, pH36-RsoleTJE and pH36-RsoleTMC were obtained in the same way. Plasmids were constructed similarly by fusing membrane protein sequences of Ncgl1337, Ncgl1221, porB, and porC in front of oleTMC for expression of oleTMC on the cell surface under the control of the promoter Eftu on pH36. All the above overexpression plasmids were verified via sequencing analysis and then transformed into C. glutamicum by electroporation. The recombinant strains grown on plates with kanamycin resistance were verified by colony PCR.

The up- and down-fragments of fasR gene were cloned from C. glutamicum S9114, and then inserted into pK18mobsacB. The fragment aar-rbs-ado was inserted between the up- and down-fragments of fasR gene, resulting in the pK18-AfasR-aar-rbs-ado plasmid. This plasmid was verified by sequencing analysis and transformed into C. glutamicum by electroporation. The correct recombinant mutant was isolated through two rounds of homologous recombination and verified by colony PCR [31].
Lignocellulose feedstock and biorefinery processing

Corn stover was harvested from Nanyang, Henan, China, in fall 2020. The raw biomass was air dried and milled, and then pretreated using the dry acid pretreatment method [32, 33]. Acid pretreatment was operated according to the protocols in [34–38]. A 20-L helical ribbon impeller-driven reactor was fed with 1200 g of corn stover (dry base) and 500–600 g of sulfuric acid solution to the dry solid weight to the acid liquid weight of 2:1. The corn stover and acid solution were co-currently fed into the reactor and stirred for 3 min at 50 rpm. The hot steam was then jetted into the reactor and maintained at 175 °C for 5 min. The pretreated corn stover solids were discharged from the bottom outlet port of the reactor without free wastewater generation, then briefly milled to move the extra-long fibers. The acid catalyst usage was adjusted according to the method previously described [39].

The solid-state biodetoxification was conducted in a 15-L bioreactor. The spore suspension of Amorphotheca resinae ZN1 was inoculated to the freshly pretreated corn stover solids and cultured at 30 °C for 48 h. Then the seed was inoculated into pretreated corn stover solids at 10% (w/w) mass ratio, and incubated at 30 °C for 36–48 h with the aeration rate of 1 vvm (air volume per culture volume per min). The brief stirring was conducted at 50 rpm every 12 h [33, 40]. The corn stover hydrolysate (CSH) was prepared by hydrolyzing the biodetoxified corn stover [16]. The duration of all fermentations was 72 h. All fermentations were carried out in triplicate, and the error bars were indicated by the standard derivations of three biological replicates.

Hydrocarbon fermentation

The seed culture was prepared as described in our previous study [16]. The shake-flask fermentation was conducted by inoculating the seed culture at 5% (v/v) inoculum ratio in 250-mL shake flasks containing 30 mL CGXII medium at 30 °C, 200 rpm. The pH was maintained at 7.0 by adding 5 M NaOH.

The bioreactor fermentation was conducted in a 1-L fermenter at 30 °C, 600 rpm and 1.4 vvm aeration, pH 7.0 by adding 2 M H2SO4 and 5 M NaOH automatically. The seed culture was inoculated into 600 mL of the fermentation medium at 10% (v/v) inoculum ratio. The cellulosic hydrocarbons fermentation was carried out in a 3-L fermenter containing 800 mL 15% (w/w) solids content corn stover hydrolysate. The other fermentation conditions were the same as mentioned above.

Hydrocarbon extraction and quantification

Hydrocarbons were extracted from 60 mL fermentation broth using 30 mL the mixture of methanol and chloroform (the ratio of methanol/chloroform was 2:1) for 24 h and then centrifuged at 10,000 rpm for 10 min. The solvent layer on the bottom of the mixture was rotary-evaporated to remove the solvent and the hydrocarbons obtained were re-dissolved by adding 1 mL of chloroform to obtain the samples for hydrocarbons measurement.

Samples were analyzed by Agilent 6890 GC–MS (Agilent Technologies, Santa Clara, CA, USA) with HP-5-MS column. The initial temperature was 50 °C and maintained for 2 min, then ramped up to 80 °C at a rate of 15 °C/min and held at 80 °C for 3 min, after that the temperature was ramped up to 280 °C at a rate of 15 °C/min and held at 280 °C for 8 min. The flow rate of the carrier gas helium was 1 mL/min. NIST MS SEARCH 2.0 library was used for qualitative analysis, and the matching degree of samples and standard products reached more than 95%. The internal calibration was 100 mg/L dodecane (C12H26) and the concentration of the hydrocarbons was calculated according to the ratio of the chromatographic peak area.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s13068-022-02129-7.

Additional file 1: Table S1. Primers used in this study.

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Authors’ contributions

YYX, KJH and ZH conducted the metabolic engineering, hydrocarbons fermentation and GC–MS analysis. PPZ conducted the screening of relevant genes. JBW provided secretive expression suggestions. CJ provided genetic manipulation and revision suggestion. JB conceived the research and experiment YYX, KJH and JB wrote the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

All data generated or analyzed during this study are included in this article. If additional information is needed, please contact the corresponding author.

Declarations

Ethics approval and consent to participate

Not applicable.
Consent for publication
Not applicable.

Competing interests
The authors declare at competing financial interest.

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