Genome sequencing of gut symbiotic Bacillus velezensis LC1 for bioethanol production from bamboo shoots

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Abstract
Background: Bamboo, a lignocellulosic feedstock, is considered as a potentially excellent raw material and evaluated for lignocellulose degradation and bioethanol production, with a focus on using physical and chemical pre-treatment. However, studies reporting the biodegradation of bamboo lignocellulose using microbes such as bacteria and fungi are scarce.

Results: In the present study, Bacillus velezensis LC1 was isolated from Cyrtotrachelus buqueti, in which the symbiotic bacteria exhibited lignocellulose degradation ability and cellulase activities. We performed genome sequencing of B. velezensis LC1, which has a 3,929,782-bp ring chromosome and 46.5% GC content. The total gene length was 3,502,596 bp using gene prediction, and the GC contents were 47.29% and 40.04% in the gene and intergene regions, respectively. The genome contains 4018 coding DNA sequences, and all have been assigned predicted functions. Carbohydrate-active enzyme annotation identified 136 genes annotated to CAZy families, including GH, GTs, CEs, PLs, AAs and CBMs. Genes involved in lignocellulose degradation were identified. After a 6-day treatment, the bamboo shoot cellulose degradation efficiency reached 39.32%, and the hydrolysate was subjected to ethanol fermentation with Saccharomyces cerevisiae and Escherichia coli KO11, yielding 7.2 g/L of ethanol at 96 h.

Conclusions: These findings provide an insight for B. velezensis strains in converting lignocellulose into ethanol. B. velezensis LC1, a symbiotic bacteria, can potentially degrade bamboo lignocellulose components and further transformation to ethanol, and expand the bamboo lignocellulosic bioethanol production.

Keywords: Bacillus velezensis LC1, Carbohydrate-active enzyme, Bamboo shoot, Cellulose, Ethanol

Background
Lignocellulose, a widely distributed, renewable and enormous biomass resource, is one of the most important raw materials for bioethanol production [1]. Bamboo, a lignocellulosic feedstock, is a regenerated biomass material with abundant resources, short growth cycle, high yield and similar chemical composition as wood, and it is considered as a potentially excellent raw material [2, 3]. Many studies have evaluated bamboo lignocellulose degradation and bioethanol production, with a focus on using physical and chemical pre-treatment [4, 5]. However, studies reporting the biodegradation of bamboo lignocellulose using microbes such as bacteria and fungi are scarce.

Lignocellulose hydrolysis, especially cellulose degradation, remains a considerable challenge in lignocellulosic bioethanol production [6]. In nature, numerous examples for lignocellulose degradation are present; of these, phytophagous insects are considered the most notable. In these insects, intestinal symbiotic microbes played
important roles in lignocellulose degradation [7]. Therefore, the intestines of phytophagous insects were considered as important locations for isolating lignocellulolytic microbes [8].

Microbial degradation of lignocellulose is a green biological refining method with advantages over physical and chemical methods [9]. The bacterial genus *Bacillus* is an excellent degrader that exhibits various abilities for degrading lignocellulose biomass, including cellulose, hemicellulose and lignin [10, 11]. Furthermore, genome sequencing, considered an efficient method for investigation of function, has been utilized in lignocellulose degradation research. However, the lignocellulose degradation of *Bacillus* is still unclear [12]. Dunlap et al. [13] reported that *B. oryzicola* and *B. methylotrophicus*, were classified into the *B. velezensis* group. Recently, the complete genome and genes associated with lignocellulose degradation of several *B. velezensis* strains were sequenced and are enriched in the genome [14–16]. However, its potential application in converting lignocellulose into bioethanol has received little attention.

In the present study, we isolated an endophytic bacteria from the gut of *Cyrtotrachelus buqueti* that showed a bamboo lignocellulose-degrading ability [17] and sequenced the whole genome of the bacteria *B. velezensis* LC1, determined the cellulase activities and analysed the ethanol production of bamboo shoot. CAZy genes involved in degradation of lignocellulose were identified through genomic analysis. The chemical changes of the cell wall components were investigated, as well as the hydrolytic and ethanol-fermenting properties of bamboo shoots.

**Results and discussion**

**Identification and cellulose-degrading potential of Bacillus velezensis LC1**

Five cellolytic strains, including PX9, PX10, PX11, PX12 and PX13, which produced clear zones around the colonies after Congo red staining, were isolated from the intestine of *C. buqueti* on CMC agar. Among the five strains, PX12 exhibited the highest cellulose hydrolysis capacity, with a higher hydrolysis capacity ratio (HCR: 4.71) than PX9 (HCR: 2.41), PX10 (HCR: 1.92), PX11 (HCR: 2.12) or PX13 (HCR: 2.56), as determined using the cellulose hydrolysis assay (Fig. 1a, b; Additional file 1: Figure S1). Based on the HCR ratio, many potent cellulosytic bacteria were previously screened from various regions, such as *Geobacillus* sp. from a hot spring and *Paenibacillus lautus* BHU3 from a landfill site [18, 19]. Similarly, PX12 was considered as a good cellulosolytic bacterium and was used for further study.

PX12 was identified and confirmed by 16S rRNA sequencing. Phylogenetic analysis of the obtained PX12 sequence revealed a 99% resemblance with *B. velezensis* strain JYP2 (NZ_CP020375.1) (Fig. 1c). Moreover, a house-keeping gene was used for phylogenetic analysis; it showed that the sequence of PX12 had a 98.6% similarity with *B. velezensis* strain S3-1 (NZ_CP016371.1) (Fig. 1d). Overall, the PX12 was identified as *B. velezensis* and was named as *B. velezensis* LC1.

Several *B. velezensis* strains have been noted for their lignocellulose-degrading abilities [14–16]. To investigate the cellulose-degrading ability, *B. velezensis* LC1 was cultured on CMC agar to determine cellulase activities by the dinitrosalicylic acid spectrophotometric (DNS) method for 6 days (Fig. 1e–g) [20]. The cellulase activities of strain LC1 were then determined. The endoglucanase activity was 0.689±0.011 U/ml at day 1 and increased to 0.752±0.013 U/ml at day 6, which was in accordance with the exoglucanase activity (from 0.359±0.016 U/ml to 0.385±0.022 U/ml), whereas the β-glucosidase activity decreased from day 6 to day 1. Previous studies have reported the cellulase activities of other lignocellulolytic *Bacillus* strains. For example, *Bacillus* sp. 275, *Bacillus* sp. R2, *B. velezensis* 157 and *B. velezensis* ZY-1-1 [10, 11, 21, 22] showed similar results as those achieved in our study. This indicated that the strain played a potential role in cellulose degradation.

**Genome sequencing and assembly of Bacillus velezensis LC1**

The genome contributes to a clear understanding of bacterial decomposition mechanisms of cellulose; thus, the genome of *B. velezensis* LC1 was analysed to decipher the genetic code involved in cellulose degradation. The complete genome sequence of *B. velezensis* LC1 was assembled into a ring chromosome with 3,929,782 bp and had a GC content of 46.5% (Fig. 2). A length of 3,502,596-bp genes was found based on gene prediction, and the ratio of gene length/genome was 89.13%. The intergene region/whole genome ratio was 7.19%, and the GC contents of the gene and intergene region were 47.29% and 40.04%, respectively. Furthermore, 4018 CDSs were contained in the genome, and all were assigned functions. CDSs were further annotated in NR, Swiss-Prot, COGs, KEGGs, GO and Pfam, and their numbers were 4018, 3520, 2996, 2186, 2718 and 3315, respectively (Table 1).

**COGs involved in carbohydrate metabolism**

In total, 3046 genes were classified into 2996 COGs, of which carbohydrate transport and metabolism, amino acid transport and metabolism and transcription were the most enriched COGs, which represented 9.46%, 7.62% and 7.29%, respectively (Fig. 3a). To elucidate the function of *B. velezensis* LC1 in cellulose degradation at the genetic level, specific COGs involved in carbohydrate
metabolism were analysed. A total of 222 genes were annotated into carbohydrate metabolism, including 130 COGs, of which the most abundant COGs were COG0366 (alpha-amylase), COG0524 (pfbk domain protein), COG2814 (Major facilitator), COG0726 (4-amino-4-deoxy-alpha-L-arabinopyranosyl undecaprenyl phosphate biosynthetic process), COG1263 (PTS System), COG0477 (major facilitator superfamily), COG1455 (pts system), COG1940 (ROK family) and COG2723 (beta-glucosidase) (Additional file 2: Table S1). COG366 encodes an alpha-amylase that acts on a bond between starch and glycogen, hydrolysing polysaccharides into glucose and maltose [23]. COG2814 is involved in cellular transport of some complexes, such as carbohydrates and amino acids. COG0477, a secondary active transporter, helps to catalyse the transport of various
substrates [24, 25]. Moreover, other important COGs in carbohydrate metabolism were annotated, e.g. COG0395 was reported to participate in carbohydrate uptake [26] and COG1109 catalysed the conversion of glucosamine-6-phosphate [27]. The high diversity of function annotations indicated that *B. velezensis* LC1 had a potent capability in lignocellulose degradation.

**GO terms annotations**

To explain the relevance of the genome of *B. velezensis* LC1, GO analysis was used to categorize genes into three categories according to matches with known sequences. In three categories, molecular function contained most numerous GO terms and gene number (3730), followed by biological process (Gene number: 3025) and cellular component (Gene number: 1637) (Fig. 3b). In molecular function, the most five pathway was ATP binding (GO:0005524; 314 genes), DNA binding (GO:0003677; 254 genes), transcription factor activity (GO:0003700; 115 genes), metal ion binding (GO:0046872; 114 genes) and hydrolase activity (GO:0016787; 84 genes). Oxidation–reduction process (GO:0055114) and regulation

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**Table 1**  **Genome features of *Bacillus velezensis* LC1**

| Features                     | Chromosome |
|------------------------------|------------|
| Genome size (bp)             | 3,929,782  |
| G+C content (%)              | 46.5       |
| GC content in gene region (%)| 47.29      |
| GC content in intergenic region (%) | 40.04    |
| Protein-coding genes (CDS)   | 4018       |
| Gene total len (bp)          | 3,502,596  |
| Gene/genome (%)              | 89.13      |
| Intergenic region len (bp)   | 427,187    |
| Intergenic len/genome (%)    | 10.87      |
| 5s rRNA                      | 9          |
| 16s rRNA                     | 9          |
| 23s rRNA                     | 9          |
| tRNA                         | 86         |
| NR annotation                | 4018       |
| Swiss-Prot annotation        | 3520       |
| Pfam annotation              | 3315       |
| COG annotation               | 2996       |
| GO annotation                | 2718       |
| KEGG annotation              | 2186       |
of transcription (GO:0006355) were most pathways in the biological process, and integral component of membrane (GO:0016021), cytoplasm (GO:0005737) and plasma membrane (GO:0005886) pathways in cellular component.

Furthermore, we analysed the GOs that are associated with carbohydrate metabolism. We identified 114 GO items associated with carbohydrate metabolism, including GO:0004553 (hydrolase activity that hydrolyses O-glycosyl compounds), GO:0005975 (carbohydrate metabolic processes) and GO:0016787 (hydrolase activity) (Additional file 3: Table S2).

**KEGG annotations**

The CDSs of *B. velezensis* LC1 were submitted to KAAS and KEGG pathways to identify metabolism pathways (Additional file 4: Table S3). As shown in Fig. 3c, of the six classification of KEGG pathways, metabolism contained the most numbers of genes, followed by environmental information processing. In KEGG metabolism annotations of *B. velezensis* LC1, carbohydrate metabolism and amino acid metabolism, which are considered its main functions, contained 392 and 285 genes, respectively. For these metabolisms, some pathways were dominant, such as sucrose and starch metabolism (ko00500), glycolysis/gluconeogenesis (ko00010), and amino and nucleotide sugar metabolism (ko00520). Forty-one genes were related to ko00500, and common enzyme endoglucanase (EC.3.2.1.4), present in ko00500, was involved in cellulose degradation (Fig. 3c). Starch and sucrose metabolic pathways occurred in *B. velezensis* LC1, indicating that cellulose could be hydrolysed into cellobiose and ultimately, β-d-glucose. In the genome, 39 genes were found in ko00010, in which d-glucose was phosphorylated into d-glucose-6-phosphate. In addition, ko00010 was linked with other pathways. For example, d-glucose-6-phosphate could be converted to pyruvate, which can be oxidized to acetyl-CoA, having an ability to enter the citrate cycle. Furthermore, ko00520 indicated that glucose from the ko00010 finally entered other pathways under various catalytic reactions. α-d-galactose can be transferred and isomerized in ko00520 and then entered into the
ascorbate and aldaraate metabolism pathways. Additionally, fructose, 1,4-β-D-xylan, and extracellular mannose were metabolized in ko00520.

The annotation involved in the degradation of lignin or aromatic compounds has also been identified. We identified various enzymes associated with lignin degradation, including oxidoreductase, reductases, dehydrogenases, esterases, thioesterases, transferases and hydrolas. Moreover, 13 monoxygenases, 12 dioxygenases, 2 peroxidases (including one DyP-type peroxidase) and 1 laccase were identified (Additional file 5: Table S4).

**Carbohydrate-active enzyme (CAZyme) annotation**

The bCAN carbohydrate-active enzymes (CAZy) annotation algorithm was used to analyse CAZy annotations to identify genes involved in lignocellulose degradation. The results showed that 136 genes were identified from CAZy families and distributed into five subfamilies. In *B. velezensis* LC1, glycoside hydrolases (GHs), which play key roles in carbohydrates degradation, contained 44 members [28]. Additionally, 38 glycosyl transferases (GTs), 30 carbohydrate esterases (CEs), 3 polysaccharide lyases (PLs), 6 enzymes with auxiliary activities (AAs) and 15 carbohydrate-binding modules (CBMs) were identified (Fig. 3d; Additional file 6: Table S5).

The GH family contained various hydrolases that acted on the glycosidic bond. For example, endoglucanases (EC 3.2.1.4) from *Bacillus* that have cellulose degradation function usually belonged to the GH5 families [29] (Table 2). In the genome, six GH13s were obliged to hydrolyze starch, such as α-amylase, α-glucosidase and α-glycosidase [30]. Four GH4s, three GH1s and one GH16 exhibited potential cellulose degradation because of enzyme function. Three GH32s were found in the genomic annotations, which contained some hydrolases and levansases, thereby revealing their ability to hydrolyse sucrose [31]. Additionally, Four GH43s, two GH51s and one GH30, responsible for xylan degradation, were considered as other important members for hemicellulose degradation (Table 2). GH43 is an important component of xylan degradation system [32]. One GH5, GH30 and GH1 each were annotated as potential β-glucosidases to utilize cellobiose. One GH53, which hydrolyses (1→4)-β-D-galactosidic linkages, was reported as an endo-1,4-β-galactosidase [33]. β-mannosidase was classified into GH26, hydrolysing parts of mannan polysaccharides [34]. Additionally, maltose phosphorylase, which belongs to GH65, was reportedly involved in trehalase degradation [35].

CEs contributing to the decomposition of xylans were also identified in the genome, including two CE3s, one CE7s, three CE10s, and seven CE4s. CE3 as a potential acetyl xylan esterase enhanced xylan solubilization [36]. The acetylxylan esterase CE7 was considered as a capable xylan-degrading enzyme [37]. CE10 previously exhibited carboxylesterase and xylanase activities involved in hemicellulose degradation [38]. Polysaccharide deacetylases, which play a role in degrading polysaccharides and are classified as a CE4, were also identified. CE4 contained not only highly specific acetylxylan esterases, but also peptidoglycan N-deacetylases involved in chitin degradation [39]. Moreover, two PL1s and one PL9 were annotated to degrade pectin. Pectate lyase (EC 4.2.2.2) usually has (1→4)-α-D-galacturonic cleavage function, causing oligosaccharides present at the end [40] (Table 2). AA4, AA6, AA7 and AA10 were included in the genome. AA4 included vanillyl-alcohol oxidases, which could transform some phenols [41]. Additionally, AA7 enzymes were involved in biotransformation or detoxification of lignocelluloses [42]. Finally, we proposed a hypothetical cellulose-degrading and ethanol-producing pathway for *B. velezensis* LC1 (Table 3).

**Comparative genomic analysis of CAZymes with other *B. velezensis* strains**

The assembled genome of *B. velezensis* LC1 was compared to the genomes of other 10 *B. velezensis* strains, including, *B. velezensis* S3-1, *B. velezensis* LS69, *B. velezensis* JTYP2, *B. velezensis* DR-08, *B. velezensis* FZB42, *B. velezensis* LPL-K103, *B. velezensis* TB1501, *B. velezensis* UCMB5036, *B. velezensis* LB002 and *B. velezensis* 157. The result showed that GH, GT and PL family numbers were the same in these strains, while the strain LC1 contained more CE and AA family members and less CBM members (Table 4). The coexistence of these genes suggests that they play important roles in the enzymatic degradation of cellulose and hemicellulose. We consider these degradation enzymes in *B. velezensis* to have potential use for bioethanol production.

**Gene expression analysis for *B. velezensis* LC1 cultured in bamboo shoot powder or glucose medium**

Expression profiles of selected *B. velezensis* LC1 lignocellulolytic enzymes genes were monitored in cultures grown on bamboo shoot powder compared to cultures utilizing glucose. Quantitative real-time PCR (qRT-PCR) investigation results of genes encoding endoglucanase (gene1950), beta-gluconase (gene3931), 6-phospho-beta-galactosidase (gene1280), glucan endo-1,6-β-glucosidase (gene2084), alpha-glucosidase (gene2945), acetyl xylan esterase (gene0351), xylan 1,4-beta-xylolysidase (gene1870), arabinoylan arabinofuranohydrolase (gene1955), alpha-N-arabinofuranosidase (gene2785), and arabinan endo-1,5-alpha-L-arabinosidase (gene2795) are shown in Fig. 4. Transcript levels of all the genes of...
interest were significantly up-regulated (at least $P < 0.05$) in BSP cultures compared to glucose cultures. It indicated that these genes involved in BSP degradation by *B. velezensis* LC1.

**Table 2** Annotated genes encoding lignocellulose-degrading enzymes of *B. velezensis* LC1, with a focus on enzymes degrading cellulose, hemicellulose and lignin

| Classification       | Gene ID   | Predicted function                                             |
|----------------------|-----------|----------------------------------------------------------------|
| Cellulose-related    | gene1950  | GH5 (endo-1,4-β-glucanase EC 3.2.1.4)                           |
|                      | gene2084  | GH30 (glucan endo-1,6-β-glucosidase EC 3.2.1.7)                 |
|                      | gene0865  | GH4 (6-phospho-β-glucosidase EC 3.2.1.86)                       |
|                      | gene2256  | C14 (glycerol-3-phosphate dehydrogenase)                        |
|                      | gene2945  | GH4 (alpha-glucosidase/alpha-galactosidase)                     |
|                      | gene3871  | GH4 (6-phospho-β-glucosidase EC 3.2.1.86)                       |
|                      | gene1280  | GH1 (6-phospho-beta-galactosidase)                              |
|                      | gene2085  | GH1 (aryl-phospho-beta-o-glucosidase)                           |
|                      | gene3887  | GH1 (6-phospho-β-galactosidase EC 3.2.1.85)                     |
|                      | gene3931  | GH16 (β-glucanase EC 3.2.1.7)                                   |
|                      | gene3442  | GH32 (sucrose-6-phosphate hydrolase EC 2.4.1.-)                 |
|                      | gene3822  | GH32 (sucrose-6-phosphate hydrolase EC 2.4.1.-)                 |
|                      | gene4083  | GH32 (levanase EC 3.2.1.65)                                     |
|                      | gene0317  | GH13 (α-glucosidase EC 3.2.1.20)                                |
|                      | gene3059  | GH13 (α-glucosidase EC 3.2.1.20)                                |
|                      | gene3448  | GH13 (α-glucosidase EC 3.2.1.20)                                |
|                      | gene0791  | PL1 (pectate lyase EC 4.2.2.-)                                  |
|                      | gene3935  | PL1 (pectate lyase EC 4.2.2.-)                                  |
|                      | gene3515  | PL9 (pectate lyase EC 4.2.2.2)                                  |
|                      | gene3891  | GH26 (β-mannosidase EC 3.2.1.78)                                |
|                      | gene3949  | GH43 (arabinan endo-1,5-α-L-arabinosidase EC 3.2.1.7)           |
|                      | gene2795  | GH43 (arabinan endo-1,5-α-L-arabinosidase EC 3.2.1.7)           |
|                      | gene1955  | GH43 (arabinofuranohydrolase EC 3.2.1.7)                        |
|                      | gene1870  | GH43 (1,4-β-xylidase EC 3.2.1.37)                               |
|                      | gene2761  | GH51 (α-N-arabinofuranosidase EC 3.2.1.55)                      |
|                      | gene2785  | GH51 (α-N-arabinofuranosidase EC 3.2.1.55)                      |
|                      | gene1954  | GH30 (glucuronoxylanase EC 3.2.1.7)                             |
|                      | gene0351  | CE 7 (acetylxyylan esterase EC 3.1.1.72)                        |
|                      | gene0450  | CE 3 (acetylxyylan esterase EC 3.1.1.72)                        |
|                      | gene2138  | CE 3 (acetylxyylan esterase EC 3.1.1.72)                        |
|                      | gene1274  | GH53 (arabinogalactan endo-1,4-β-galactosidase EC 3.2.1.89)     |
| Hemicellulose-related| gene1891  | AA10 (chitin-binding protein)                                  |
|                      | gene2781  | AA4 (glycolate oxidase)                                        |
|                      | gene0978  | AA6 (p-benzoquinone reductase)                                 |
|                      | gene0576  | AA7 (FAD-binding protein)                                      |
|                      | gene0946  | AA7 (FAD-dependent oxidase)                                    |
|                      | gene1182  | AA7 (FAD-binding protein)                                      |

Cellulose degradation efficiency and fermentation efficiency of bamboo shoots by *B. velezensis* LC1

Several previously reported genomes of *B. velezensis* strains contained genes encoding enzymes having lignocellulose-degrading potential [14–16]. However, lignocellulose-degrading abilities have not been completely verified in *B. velezensis* [12]. *B. velezensis* LC1, isolated from the intestine of *C. buqueti*, which was reported to be a microflora with lignocellulose-degrading ability, was prepared in a bamboo shoot powder (BSP) degradation assay [17]. We first determined the cellulose degradation efficiency to be 39.32% (Fig. 5a). Furthermore, we determined the glucose and xylose content in
the degradation products to be 55.30 ± 1.40 mg/L and 488.81 ± 45.06 mg/L, respectively (Fig. 5b). The reducing sugars in the culture medium was mainly derived from the hydrolysis of cellulose and hemicellulose in BSPs, the reducing sugar content was determined to reflect the degree of conversion of lignocellulose. It indicated that the cellulose and hemicellulose of BSP were degraded with incubation with *B. velezensis* LC1.

Shimokawa et al. [43] reported that bamboo shoot was an excellent biomass stock for ethanol production.

### Table 3 Enzymes in cellulose-degrading and ethanol-producing pathways of *B. velezensis* LC1

| Gene ID | Enzymes definitions |
|---------|---------------------|
| 1 | gene1537; gene2613 | PTS system, fructose-specific IIA component [EC:2.7.1.69] |
| 2 | gene3441; gene3821 | Beta-fructofuranosidase [EC:3.2.1.26] |
| 3 | gene0676, gene3887; gene3888 | Fructokinase [EC:2.7.1.4] |
| 4 | gene3065 | Glucose-6-phosphate isomerase [EC:5.3.1.9] |
| 5 | gene1004 | Phosphoglucomutase [EC:5.4.2.2] |
| 6 | gene1974; gene1975; gene3557 | UTP-glucose-1-phosphate uridylyltransferase [EC:2.7.7.9] |
| 7 | gene1950 | Endoglucanase [EC:3.2.1.4] |
| 8 | gene2486 | Glucokinase [EC:2.7.1.2] |
| 9 | gene3065 | Glucose-6-phosphate isomerase [EC:5.3.1.9] |
| 10 | gene2837 | 6-Phosphofructokinase 1 [EC:2.7.1.11] |
| 11 | gene3722 | Fructose-bisphosphate aldolase, class II [EC:4.1.2.13] |
| 12 | gene3378 | Triosephosphate isomerase (TIM) [EC:5.3.1.1] |
| 13 | gene2815; gene3381 | Glyceraldehyde 3-phosphate dehydrogenase [EC:1.2.1.12] |
| 14 | gene3380 | Phosphoglycerate kinase [EC:2.7.2.3] |
| 15 | gene3377 | 2, 3-Bisphosphoglycerate-independent phosphoglycerate mutase [EC:5.4.2.12] |
| 16 | gene3377 | Enolase [EC:4.2.1.11] |
| 17 | gene2837 | Pyruvate kinase [EC:2.7.1.40] |
| 18 | gene1567 | Pyruvate dehydrogenase E1 component alpha subunit [EC:1.2.4.1] |
| 19 | gene1568 | Pyruvate dehydrogenase E1 component beta subunit [EC:1.2.4.1] |
| 20 | gene1570 | Pyruvate dehydrogenase E2 component (dihydrolipoamide acetyltransferase) [EC:2.3.1.12] |
| 21 | gene0862 | Dihydrolipoamide dehydrogenase [EC:1.8.1.4] |
| 22 | gene3777 | Phosphate acetyltransferase [EC:2.3.1.8] |
| 23 | gene2856 | Acetate kinase [EC:2.7.2.1] |
| 24 | gene0778; gene2879; gene4000 | Aldehyde dehydrogenase (NAD +) [EC:1.2.1.3] |
| 25 | gene1937; gene2005; gene3037; gene0198; gene0266; gene0591; gene0675; gene0788 | Alcohol dehydrogenase [EC:1.1.1.1] |

### Table 4 Comparative genomic analysis of CAZymes with other *B. velezensis* strains

| Strains | GHS | GT | PL | CE | AA | CBM |
|---------|-----|----|----|----|----|-----|
| *B. velezensis* LC1 | 43 | 38 | 3 | 30 | 6 | 15 |
| *B. velezensis* S3-1 | 40 | 36 | 3 | 10 | 1 | 27 |
| *B. velezensis* LS69 | 40 | 36 | 3 | 10 | 1 | 27 |
| *B. velezensis* JTYP2 | 39 | 36 | 3 | 10 | 1 | 28 |
| *B. velezensis* DR-08 | 40 | 36 | 3 | 10 | 1 | 27 |
| *B. velezensis* FZB42 | 37 | 36 | 3 | 10 | 1 | 27 |
| *B. velezensis* LPL-K103 | 36 | 32 | 3 | 10 | 1 | 29 |
| *B. velezensis* TB1501 | 40 | 36 | 3 | 10 | 1 | 28 |
| *B. velezensis* UCMB5036 | 40 | 36 | 3 | 10 | 1 | 26 |
| *B. velezensis* LB002 | 42 | 37 | 3 | 10 | 1 | 27 |
| *B. velezensis* 157 | 41 | 36 | 3 | 10 | 1 | 29 |
due to its high saccharification efficiency. To determine the ethanol production capacity of \textit{B. velezensis} LC1, a 6-day-treated hydrolysate was prepared for ethanol fermentation using glucose-fermenting \textit{S. cerevisiae} and xylose-fermenting \textit{E. coli} KO11. Ethanol production from BSP hydrolysate using \textit{B. velezensis} LC1 is illustrated in Fig. 5c. The result showed that the ethanol yield continually increased during 48–96 h, and reached $7.21 \pm 0.24$ g/L at 96 h, while the reducing sugar continually decreased after incubation 48 h. This indicated that \textit{B. velezensis} LC1 had a potent ability to convert lignocelluloses in bamboo shoot to ethanol.

Conclusions
The genome of a gut symbiotic bacteria, \textit{B. velezensis} LC1, was sequenced and analysed. The genome comprises a ring chromosome of 3,929,782 bp and has a GC content of 46.5%. A total of 136 CAZyme genes involved in lignocellulose degradation were annotated in the genome, and cellulose-degrading and ethanol-producing pathways were proposed. Moreover, the expression level of some CAZyme genes involved in cellulose and hemicellulose degradation, were up-regulated in bamboo shoot powder substrate. Moreover, a transcriptomic study would perform to identify the role of CAZyme genes in lignocellulose degradation as a future study. The cellulose degradation of bamboo shoot by the strain was 39.32%, and the hydrolysate was subjected to ethanol fermentation; the ethanol yield was 7.2 g/L at 96 h. This study suggests that \textit{B. velezensis} LC1 could be used for bamboo lignocellulose degradation and bioconversion of lignocelluloses to ethanol.

Methods
Insect sample, isolation and identification of cellulolytic bacteria
\textit{Cyrtotrachelus buqueti} specimens were sampled from the Muchuan County (E 103° 98′, N 28° 96′), China. Gut was extracted from individual insects and stored at 4 °C for isolation of bacteria. The gut was blended, homogenized and serially diluted (10$^{-1}$ to 10$^{-9}$), and inoculums of 10$^{-7}$ to 10$^{-9}$ dilution were plated on carboxymethyl cellulose (CMC) agar [44] for cellulolytic bacteria screening. Congo red dye was used to screen the cellulose-degrading bacteria as described by Teather and Wood [45].

Molecular characterization of bacterial isolate
The 16sRNA V3–V4 region was considered for amplification for bacterial identification and amplified using bacterial primers (27F 5′-AGA GTT TGATCMTGG CTC AG-3′ and 1492R 5′-TAC GGY TAC CTT GTA CGA CTT-3′); moreover, F 5′-GCCCATATTTCATTTCTCC-3′ and R 5′-GTCGTGTTATGGAAATAGG-3′ were selected for amplification of the house-keeping gene \textit{rpoB}. Thermocycling conditions were as follows: initial denaturation at 94 °C (2 min), followed by 30 cycles of denaturation at 94 °C (30 s), annealing at 55 °C (30 s) and extension at 72 °C (100 s), ultimately extending at 72 °C (2 min). The amplicons were checked by electrophoresis on a 1% agarose gel. MEGA5 was used to establish phylogenetic relationships among the obtained sequence and reference genes that were retrieved in NCBI GenBank through the neighbour-joining method.

Genomic DNA extraction and genome sequencing
Genomic DNA of \textit{B. velezensis} LC1 was extracted according to the CTAB method based on the protocol described by Lin [46], and it was purified using the Wizard Genomic DNA Purification Kit (Vazyme Biotech Co., Ltd, Nanjing, China). Whole-genome sequencing, performed by Shanghai Majorbio Bio-pharm Technology Co., Ltd (Shanghai, China), was fulfilled through PacBio systems with an average genome coverage of 100 x for the raw data [47]. HGAP 2.0 was used to filter and assemble reads to the scaffold.
Function annotation of *B. velezensis* LC1

Glimmer 3.02 was used to predict coding DNA sequences (CDSs). A BLAST search was then conducted for CDSs in some widely used databases: NCBI non-redundant (NR) database, Swiss-Prot, COGs, Gene Ontology (GO) and KEGGs [48, 49]. GO, an important bioinformatics tool, unified expressions of gene and genetic products in all species [50]. KEGGs, a database resource to understand high-level functions, also could analyse metabolic pathways. Additionally, the CAZymes were identified, classified and annotated using CAZymes database (CAZyDB: http://www.cazy.org/).

Quantitative real-time PCR

The primers used for qRT-PCR in this study were performed in Additional file 7: Table S6. PCR was performed under following conditions: 10 min initial denaturation at 95 °C, 45 cycles of 5 s denaturation at 95 °C, 50–65 °C anneal for 30 s, and 30 s extension at 55 °C, finally 10 s extension at 95 °C. All experiments were performed three times and analysed by $2^{-\Delta\Delta CT}$ Method. 16S rRNA was used as reference gene.

Bamboo shoots degradation by *B. velezensis* LC1

To obtain fermentable sugar from bamboo shoots, *B. velezensis* LC1 was used to degrade the bamboo shoot powder (BSP), which was prepared as described by Luo et al. [17]. *B. velezensis* LC1 was cultured in liquid medium at a pH of 7.2, temperature of 37 °C and 200 rpm for 6 days. The culture medium comprised BSP 10 g/L, (NH₄)₂SO₄ 2 g/L, K₂HPO₄ 1 g/L, KH₂PO₄ 1 g/L, MgSO₄ 0.2 g/L, CaCl₂ 0.1 g/L, FeSO₄·7H₂O 0.05 g/L and MnSO₄·H₂O 0.02 g/L. The reaction mixture was incubated at 100 °C for 30 min to terminate the reaction and centrifuged at 13,000 rpm for 10 min, and the hydrolysate-containing supernatant and deposit were collected separately. The obtained deposit was dried and weighed to determine the cellulose levels using the Van Soest method [51]. The hydrolysate-containing supernatant was used to determine the glucose and xylose contents according to the NREL methods [52].

**Fig. 5** Cellulose-degrading efficiency and fermentation performance of BSP by *Bacillus velezensis* LC1. **a** Cellulose degradation efficiency. **b** Glucose and xylose contents of the hydrolysate. **c** Ethanol production at 48 h, 60 h, 72 h, 84 h and 96 h
The BSP hydrolysate supernatant was sterilized and stored at –20 °C for ethanol fermentation.

Fermentation

_Saccharomyces cerevisiae_, a glucose-fermenting strain, was pre-cultured in YPD at 30 °C for 24 h, and _E. coli_ KO11, a xylose-fermenting strain, was cultivated in LB at 37 °C for 24 h. _S. cerevisiae_ (50 g/L) and _E. coli_ KO11 (100 g/L) were prepared after centrifugation of pre-cultured cells. The initial cell concentrations were 0.33 g/L (_S. cerevisiae_) and 1.0 g/L (_E. coli_ KO11) at the beginning of fermentation. 100 ml hydrolysate was used for ethanol fermentation in 250-ml serum bottles under anaerobic conditions. The fermentation was performed at 37 °C and 200 rpm for 96 h. After 48 h fermentation, ethanol production was monitored every 12 h. The ethanol concentration was determined via HPLC. All actions were repeated three times.

Supplementary information

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s13068-020-1671-9.

**Additional file 1: Figure S1.** Cellulolytic activities of 4 isolates, PX9, PX10, PX11, and PX13, cultured on the CMC agar plate with congo red.  
**Additional file 2: Table S1.** GO annotations of carbohydrate metabolism in genome of _B. velezensis_ LC1.  
**Additional file 3: Table S2.** COG annotations of carbohydrate metabolism in genome of _B. velezensis_ LC1.  
**Additional file 4: Table S3.** KEGG Pathway classification of _B. velezensis_ LC1.  
**Additional file 5: Table S4.** Annotation of dioxygenase, monooxygenase, peroxidase, laccase and oxidoreductase genes identified in the genome of _B. velezensis_ LC1.  
**Additional file 6: Table S5.** CAZy annotation in the genome of _B. velezensis_ LC1.  
**Additional file 7: Table S6.** Primers used for RT-qPCR analysis.

**Abbreviations**

_B. velezensis_; _Bacillus velezensis_; GH: Glycoside hydrolase; GT: Glycosyl transferase; CE: Carbohydrate esterase; CBM: Carbohydrate-binding domain; PL: Polysaccharide lyase; AA: Auxiliary activities; CAZyme: Carbohydrate-active enzyme; NCBI: The National Center for Biotechnology Information; BSP: Bamboo shoot powder; YPD: Yeast extract peptone dextrose medium; DNS: 3,5-Dinitrosalicylic acid; CDS: Sequence coding for amino acids in protein.

**Acknowledgements**

The data were analysed on the free online platform of Majorbio Cloud Platform (www.majorbio.com). We also thank other members of the laboratory for suggestions and discussion regarding this work and revision of the manuscript.

**Authors’ contributions**

CL and YL designed and performed the experiments, CL, YL and LL wrote the manuscript; LZ, LL, YL, HT and XX analysed the data. All authors read and approved the final manuscript.

**Funding**

This work was supported by Sichuan science and technology program (2019YFG0139).

**Availability of data and materials**

The sequence reads from this article have been deposited at the NCBI Sequence Read Archive under the accession PRJNA74012. The assembly data set supporting the results of this article has been deposited at GenBank under the accession CP044349. The version described in this paper is CP044349.

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

The authors have consented for publication.

**Competing interests**

The authors declare that they have no competing interests.

**Received** 4 October 2019 **Accepted** 29 January 2020 **Published online** 28 February 2020

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