Screening a bacterium and its effect on the biological degumming of ramie and kenaf

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Materials and Methods

The soil samples were collected from a continuously cultivated banana plantation in Lingshui, Sanya, Hainan Province in China (50 m above sea level in 18°28’ N and 110°1’ E), and then crushed and passed.
through a 30-mesh sieve to remove large lumps and impurities. Twenty grams of sieved soil samples were placed in Erlenmeyer flasks filled with 10 g glass beads (Φ 2 mm), and a 200 mL sterile physiological saline was mixed to prepare a soil suspension. The Erlenmeyer flasks were then shaken at 200 r min⁻¹ for 2 h. 20 mL of the solution above the suspension was inoculated into a 200 mL ramie degumming system (20 g: 200 mL ramie-to-water ratio) and cultured at 35 °C at 180 r min⁻¹ for 24 h. A 20-mL suspension taken from the previous step was inoculated into a 200 mL kenaf degumming system (20 g: 200 mL kenaf-to-water ratio) and cultured at 35 °C at 180 r min⁻¹ for 24 h. Ramie and kenaf were alternately enriched three times. The suspension was spread onto the Petri dish with a nutrient agar (NA) medium that contained 35 g L⁻¹ of nutrient agar and 5 g L⁻¹ of glucose after serial dilutions. The Petri dishes were incubated at 35 °C for 22-24 h and then the single colonies were obtained.

Active pectin-degrading microorganisms were screened using a halo-producing assay on a selective medium, placed in Petri dishes that contained 5 g L⁻¹ of sodium polygalacturonate (Sigma, P91350), 10 g L⁻¹ of tryptone, 5 g L⁻¹ of yeast extract, 5 g L⁻¹ of NaCl, and 20 g L⁻¹ agar. Only active isolates can form clear halos in the pectin plate. The above dominant isolate was cultured in a nutrient broth medium (NB) composed of 10 g L⁻¹ of glucose, 5 g L⁻¹ of beef extract, 5 g L⁻¹ of peptone, and 5 g L⁻¹ of NaCl at 35 °C for 6 h at 200 r min⁻¹. We dabbled 0.5 μL of the suspensions on the selective medium, and then inverted and incubated at 35 °C for 18-20 h. Lugol iodine solution turned the background dark brown; as a result, clear haloes appeared around the active pectin-degrading strains. Diameter ratios H/C of the pectinolytic bacteria were calculated using the data obtained by the diameters of the bacterial colonies (C) and their hydrolysis circles (H) [Merin et al., 2015].

Phenotypic and biochemical characteristics were performed according to the reference [Mani et al., 2015]. Briefly, the pure cultures isolated on the NA medium at 35 °C for 24 h were gram stained, then the morphology of the strain were observed under the optical microscope at a magnification of 40 × 100. Characteristics of the single colonies such as form, size, color, swelling and gloss, were described. Furthermore, temperature, pH and NaCl tolerance of the strain were tested.

Genomic DNA was extracted by MiniBEST Bacterial Genomic DNA Extraction Kit Ver.2.0, and the 16S rDNA sequence was amplified by the 16S rDNA PCR Amplification Kit with forward primer (Pf): 5'-GAGCGGAGATAAACATTTCCACAGG-3' and reverse primer (Pr): 5'-CGCCAGGGTTTTCCAGCTCAGG-3' (Hakovirta et al., 2016). PCR products were purified and sequenced by Sangon Biotech (Shanghai) Limited Company in China. The 16S rDNA sequence obtained from strain hnl-1 was submitted to GenBank to be assigned a gene accession number, and then compared with the sequence from the GenBank database by the Blastn method. The Neighbor-joining method of the MEGA6.0 software was used to construct the phylogenetic tree of the 16S rDNA of hnl-1 and target bacteria strains belonging to the same genus but different species of Bacillus spp.

20 g raw material (shell-free ramie or kenaf) was stored in a 500 mL flask. The bacteria suspension was prepared according to inoculation of 2 %, and the temperature was adjusted to 35 °C. The raw material (shell-free ramie or kenaf) was soaked in a 200 mL bacteria suspension (20 g: 200 mL raw material-to-bacteria suspension bath ratio) and fermented at 35 °C at 180 r min⁻¹. Starting at 0.5 h, the fermentation solution was collected every 2 h, and labeled in order, 0, 2, 4, 6, 8, and 10 h. After fermenting at 10 h, the mixtures were rinsed with 90 °C hot water for 30 min. The bast fibers were washed in a washing machine for 20 min, and then dried in oven at 50 °C to obtain fine bast fibers.

The number of living strains was calculated by strain colonies in a solid culture medium. The chemical oxygen demand (COD) of the degumming solution was measured according to the standards for industrial circulating cooling water using the COD–potassium permanganate method (GB/T 15456-2008). The reducing sugar content was determined by the 3, 5-dinitrosalicylic acid method, and glucose was used as the standard for calculating the reducing sugar content [Hu et al., 2008].

Fiber production rate: With \( G_m \) representing the quantity of ramie or kenaf raw material, and \( G_f \) the quantity of fine fiber, fiber production rate \( r \) is calculated as follows [Biswa et al., 2016]:

\[
\text{r}(\%) = \frac{G_f}{G_m} \times 100
\]

Residual gum rate: With \( G_f \) representing the quantity of fine ramie or kenaf fiber and \( G_g \) the weight of fine fiber treated by alkali, residual gum rate \( w \) is calculated as follows [Zhou et al., 2017]:

\[
\text{w}(\%) = \frac{G_g - G_f}{G_f} \times 100
\]

Raw materials, degummed ramie and kenaf fibers were sampled. Sprayed with gold powder, the morphological characteristics of the longitudinal section of the samples were observed by scanning electron microscope (SEM).

A hundred individual fibers were randomly selected for each sample, and an XQ-2 fiber tensile tester was used to perform a stretch test with a maximum range of 100 cN, a pre-tension of 200 mg, a clamping distance of 20 mm, and a tensile speed of 10 mm min⁻¹. Then, the breaking strength of the fiber bundle was measured.

**Results**

153 single bacterial colonies on NA were isolated from soil samples [data not shown]. Seven bacterial strains producing pectin hydrolysis circles were obtained through a selective culture medium [Figure 1]. The H/C of the hydrolysis circle to the bacterial colony produced
by hn1-1 was the largest, reaching 2.4 (Table 1); therefore, hn1-1 was selected as the best pectinase producer. The hn1-1 colony was tested for extracellular pectinase production in selective culture medium without addition of agar, and it revealed pectinase activity 143.53 IU mL\(^{-1}\). Compared with the known Pectobacterium sp. CXJZU-120 (high yield 124.61 IU mL\(^{-1}\)) and the Bacillus subtilis T66 (low yield 33.23 IU mL\(^{-1}\)) measured under the same conditions, hn1-1 secreted high-yield of pectin-degrading enzyme, indicating a good ability of bast fibers bio-degumming [Liu et al., 2012].

The cells of hn1-1 were Gram-positive, motile, rods for aerobic growth at 35 °C (Figure 2). Colonies incubated on NA at 35 °C for 24 h were flat, yellowish red disks that had transparent edges and measured 2.0–2.5 mm in diameter. The temperature, pH, and NaCl concentration ranges for growth are 20–39 °C, 6.0-9.0, and 0-50 g L\(^{-1}\), respectively.

The full length of this sequence was 1,421 bp, as determined through the extraction of the genomic DNA of bacterial strain hn1-1 and the amplification and sequencing of the 16S rDNA [GenBank accession number: KX013542]. The 16S rDNA sequence of hn1-1 is 100 % consistent with that of B. cereus SE1, as indicated by the Blast comparison performed for the target sequence (GenBank accession number: KJ461699.1). The cluster analyses of the 16S rDNA sequence of the standard bacterial strains and the target bacterial strain belonging to Bacillus with different species were performed with MBGA6.0 software (Figure 3). Therefore, the bacterial strain can be preliminarily identified as the B. cereus hn1-1 by its morphological and molecular characteristics. However, the final taxonomic status of B. cereus should be determined by the DNA-DNA hybridization between the target bacterial strain and its closely related typical bacterial strains.

Numbers of living strains in the biological degumming solution collected at different times were accounted for the strain colonies in the solid culture medium. During the fermentation process of the two raw materials, the living bacterial strain of B. cereus hn1-1 exhibited no change at 0–2 h and belonged to the delay period; the number of the living bacterial strain increased fast at 2–6 h, coming to the logarithmic growth phase; the number of the living bacterial strains have little change at 6–12 h, belonging to the stable phase (Figure 4A).

The reducing sugar content of the fermented solution sampled at regular intervals (Figure 4B) showed that during the fermentation process of kenaf by B. cereus hn1-1, the reducing sugar content increased rapidly at 0–2 h due to the dissolution of a large amount of water-soluble sugars in kenaf materials. The reducing sugar content then decreased rapidly at 2–6 h, thereby suggesting that the growth and reproduction of bacterial strains consumed a large amount of reducing sugars in the fer-

| Table 1 – H/C values by microorganism. |
|----------------------------------------|
| Strain No. | H/C | Strain No. | H/C | Strain No. | H/C |
|------------|-----|------------|-----|------------|-----|
| hn1-1      | 2.4 | hn 3-1     | 1.8 | hn 6-2     | 2.1 |
| hn 2-2     | 2.1 | hn 3-5     | 2.3 | hn 7-2     | 0   |
| hn 2-4     | 1.5 | hn 6-1     | 1.7 | hn 7-3     | 0   |
mentation liquid. The reducing sugar content stabilized at 6–10 h, thereby suggesting that the reproduction of *B. cereus* hn1-1 consumed carbohydrate substances with a comparable amount to oligosaccharides continuously produced by the extracellular, non-cellulolytic, enzyme-degrading gum. Meanwhile, reducing sugar content was stable throughout the fermentation process of ramie by *B. cereus* hn1-1.

The fermentation solution was sampled at regular intervals and tested for pH. The pH of the two raw materials by *B. cereus* hn1-1 in the fermentation process increased as the fermentation cycle increased (Figure 4C).

The fermented solution COD was measured using solution samples, which were collected at regular intervals (Figure 4D). The COD value of *B. cereus* hn1-1 in the fermentation process of ramie first decreased and then increased, thereby indicating that in the early stages of degumming (0–2 h), the growth and reproduction of bacterial strains consumed water-soluble organic compounds in ramie raw materials, decreasing the COD value. In the middle and late stages of degumming (2–10 h), the growth and reproduction of the bacterial strains consumed organic substances in the fermentation solution, and the degumming bacterial strains continued to break down the ramie colloid to produce organic substances. Meanwhile, production was greater than consumption, thereby resulting in a continuous increase in COD value. In the fermentation process of kenaf by *B. cereus* hn1-1, COD value increased rapidly at 0–2 h, possibly because of the dissolution of a large amount of organic substances in kenaf raw materials. COD value decreased at 2–10 h, leading to the conclusion that kenaf gum produces more organic substances than that consumed by *B. cereus* hn1-1.

As shown in Table 2, fiber production rates of ramie and kenaf were 72 % and 76 %, and their residual gum rates were 4 % and 5 %, respectively, after degumming by *B. cereus* hn1-1 for 10 h.

The surface morphologies of ramie and kenaf fibers before and after biological degumming were observed by a scanning electron microscope. Compared with the raw materials [Figures 5A and 5C], ramie and kenaf fibers easily decentralized after degumming with a smooth surface [Figures 5B and 5D], and thereby indicating good gum removal.

As shown in Table 3, the ramie and the kenaf fibers bio-degummed by *B. cereus* hn1-1 were stronger than the chemically degummed fibers. Therefore, biological degumming caused less damage to fibers than chemical degumming, thereby obtaining a better quality of fine ramie and kenaf fibers.

**Discussion**

*B. cereus* is an advantageous bacterial strain of flax retting, which played a key role in the first 72 h of the...
biological flax degumming (Zhao et al., 2016). This study is the first to explore the biological degumming effects of *B. cereus* on ramie and kenaf. *B. cereus* achieved the same results as the ramie degumming technology currently used in the industry (such as CXJZ-120). The results were close to the effects of applying that bacterial strain for the degumming of kenaf (Chiliveri et al., 2016; Liu et al., 2012). Compared with other degumming bacterial strains (Basu et al., 2009; Zheng et al., 2001), *B. cereus* has a shorter degumming cycle (10 h) and ideal gum removal capabilities (residual gum rate is less than 6 %) and produces fiber of good quality (breaking strength is more than 850 cN dtex\(^{-1}\)).

The biological degumming of bast fibers is divided into bacterial degumming and enzymatic degumming, depending on the initial additives, and their essence is the degradation of gum catalyzed by non-cellulolytic enzymes (Bruhlmann et al., 2000). Large-scale enzymatic degumming is difficult because the complicated gum complexes of the bast fiber materials make building degumming enzyme systems that match the given gum components challenging. Moreover, compounded enzyme preparation is expensive. A viable bacteria degumming process used the “culture of bacteria by gum production of enzyme by bacteria enzyme degumming bast fibers” cycle to biologically degum bast fibers. Bacterial strains consume part of the gum hydrolysates. On the one hand, this consumption pattern can produce a degumming enzymatic reaction that is not refined by the inhibitory role of product feedback and promote an ideal biological degumming of bast fibers. On the other hand, this consumption can effectively reduce the COD value of sewage and emission of organic pollutants. *B. cereus* h1n1-1 is advantageous because it consumes the oligosaccharides produced by ramie and kenaf gum degradation in different degrees. The different results yielded by this bacterial strain in fermentation of the two raw materials (Figure 4B) possibly due to differences between the soluble sugar contents and properties of the raw ramie and kenaf materials. It may be also related to the nutritional characteristics of the bacterial strain (whether the bacterial strain is

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**Table 2** – Comparison on fiber production rates and residue gum rates.

|          | Biological degumming | Chemical degumming | Biological degumming | Chemical degumming |
|----------|----------------------|--------------------|----------------------|--------------------|
| Ramie    | 72 ± 1.6             | 60 ± 1.1           | 4 ± 0.09             | 10 ± 0.1           |
| Kenaf    | 76 ± 2.1             | 65 ± 1.2           | 5 ± 0.08             | 18 ± 0.2           |

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**Table 3** – Comparison on mechanical characteristics of degummed bast fibers.

|          | Biological degumming | Chemical degumming | Biological degumming | Chemical degumming |
|----------|----------------------|--------------------|----------------------|--------------------|
| Fiber fineness (dtex) |                         |                    | Breaking strength (cN dtex\(^{-1}\)) |                        |
| Ramie    | 6.25 ± 0.18          | 8.2 ± 0.24         | 850 ± 14             | 750 ± 10           |
| Kenaf    | 12.25 ± 0.32         | 18.2 ± 0.38        | 969 ± 11             | 785 ± 9            |

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**Figure 5** – Fiber morphology scanned by scanning electron microscope (SEM). Note: A Raw ramie material; B Bio-degummed ramie fiber; C Raw kenaf material; D Bio-degummed kenaf fiber.
preferred to the gum of the given raw material) [Fan et al., 2015]. The change regulation of pH (Figure 4C) was almost consistent with that of viable count (Figure 4A), which may be because the degumming bacterial strains consumed a large amount of acidic nutrients in the fermentation solution or secreted a large amount of alkaline extracellular enzymes that led to the increase in pH; nonetheless, this area deserves further study [Kalim and Ali, 2016].

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