CELLULOLYTIC POTENTIAL OF A NOVEL STRAIN OF PAENIBACILLUS SP. ISOLATED FROM THE ARMORED CATFISH PAROTOCINCLUS MACULICAUDA GUT

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ABSTRACT

A cellulolytic bacterial strain, designated P118, isolated from the gut of the tropical fish Parotocinclus maculicauda was identified as belonging to the genus Paenibacillus based on phenotypic and chemotaxonomic characteristics and the 16S rRNA gene sequence. The novel strain was Gram-positive, spore-forming and rod-shaped. Catalase but not oxidase was produced. Carboxymethylcellulose was hydrolyzed but starch or gelatin was not. Acetoin production was negative whereas nitrate reduction and urease production were positive. Many carbohydrates served as carbon sources for growth. MK-7 was the predominant isoprenoid quinone. Anteiso-C15:0 (38.73%) and C16:0 (20.85%) were the dominant cellular fatty acids. Strain P118 was closely related to Paenibacillus amyloyticus NRRL NRS-290, P. pabuli HSCC 492, P. tundrace Ab10b, P. xylanexedens B22a, and P. tylopili MK2 with 98.3-98.8% 16S rRNA gene sequence similarity. The results presented here suggest that strain P118 represents a novel species of the genus Paenibacillus and it is a potential strain for further studies concerning its role in the production of industrially important products from cellululosic biomass.

Key words: Bioprospection; Cellulose degradation; Paenibacillus sp.; Parotocinclus maculicauda; Taxonomy

INTRODUCTION

The armored catfish Parotocinclus maculicauda is a member of the Loricariidae family and the Hypoptopomatinae sub-family, formed by 16 genera and 79 species (27). Fish belonging to the species P. maculicauda are detritivorous, live well in rivers with clean water with a neutral to slightly acidic pH and in lower tropical temperatures, and are popularly known as Red Fin Dwarf Pleco. They are usually found in the south of Brazil (5). However, Leitão et al. (16) have also

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observed the presence of *P. maculicauda* in different coastal rivers of the Atlantic Forest. *P. maculicauda* shows a behavior based on grazing, being found near to the bottom of the creek on rocks, logs and submerged plants, scraping the periphyton matrix (4). The bacterial community found associated with this fish is almost unknown. However, it is well established that the microflora present in water influences directly the microorganisms found in the interior of the digestive tract of fish (9). Based on the kind of diet imposed to *P. maculicauda*, which consists mainly of plant material from allochthonous and autochthonous origins, it is expected the presence of cellulolytic bacteria in the fish gut.

Bioprospection is the systematic search of new sources of chemical compounds, genes, proteins, microorganisms and other products that have economic potential and can be found in biodiversity. Less than 1% of the existent microorganisms have been cultivated and characterized so far. Novel microorganisms obtained from poorly studied environments can bring new insights on novel genes and enzymes. In this context, the bacterial community present in the digestive tract of *P. maculicauda* can be identified providing survival and competencies to its host. Among the different isolates obtained, specific bacterial populations can be selected for one specific characteristic of interest. In this study, among different bacterial isolates, a cellulolytic strain denoted P118 was isolated from the gut of *P. maculicauda*. Therefore, our goal was to identify and characterize this isolate and to demonstrate its potential for cellulose degradation. Biochemical and molecular data obtained suggest that the isolate belongs to a new species of *Paenibacillus*.

**MATERIALS AND METHODS**

**Fish samples**

Twenty fishes belonging to the *P. maculicauda* species were obtained from Mato Grosso river (Saquarema, Rio de Janeiro, Brazil) located at 22°53’32.5”S and 42°38’59.8”W. This river has 12 km of extension, 2 to 4 m of width and at least 20 inches of depth. The fishes were collected manually using trawls and immediately taken to the laboratory for dissection in aseptic conditions.

**Isolation of bacterial strains**

The food content from the digestive tract of the different fish samples was transferred to sterilized tubes and portions of 200 mg were mixed with 1.8 ml of saline. This mixture was diluted in saline and plated in triplicate onto Trypticase Soy Broth (TSB, Difco) - agar (1.2%). After 48 h of incubation at 32°C, different colonies were selected based on their morphotypes. The different isolates were cultivated and maintained in slants with the same medium at 32°C and room temperature, respectively.

**Detection of cellulolytic activity**

The different isolates were inoculated as a 5 µl-spot on the surface of a carboxymethylcellulose (CMC) containing medium (CMC, 10g; KH₂PO₄, 4g; Na₂HPO₄, 4g; tryptone, 2g; MgSO₄.7H₂O, 0.2g; CaCl₂, 0.001g; FeSO₄.7H₂O, 0.004g; agar 15g, 1 l distilled water, pH 7.0; (3)). After incubation at 32°C for 48 h, the bacterial cells were flooded with 5 ml of congo red (0.1%) and washed with NaCl (1%). The cellulolytic strains were identified by the formation of a clear halo around the bacterial growth (3).

**Cellulase production**

A pre-inoculum of the selected strain (P118) was prepared in a 250 ml Erlenmeyer flask containing 50 ml of TSB. This flask was inoculated with a loop full of the strain grown in a TSB agar-containing plate and incubated at 32°C under agitation (rotator shaker 200 rpm) for 24 hours. Strain P118 was then inoculated (25 µl of the pre-inoculum) in different flasks containing 25 ml of CMC medium (without tryptone), pH 7.0, and was incubated at the same conditions for 5 days. Samples were collected daily and the whole content of each
flask (filtrate obtained after filtration through glass microfiber filter - Millipore) was used for determining the cellulase activity at different periods of time. Results were expressed as the average of two replicates.

Carboxymethylcellulase (CMCase) activity was assayed by measuring the release of reducing sugars in a reaction mixture of 1ml of the crude enzyme and 1 ml of 4% (w/v) CMC suspension in 50mM sodium citrate buffer (pH 4.8) incubated at 50°C for 10 min (6). Reducing sugars were assayed by the dinitrosalicylic acid (DNS) method (18). One unit (U) of CMCase activity corresponded to 1μmol of glucose equivalent released per minute under the assay conditions.

**Morphological, physiological and biochemical characterization**

Cellular morphology, form and position of spores and swelling of the sporangia were observed on crystal violet-stained smears by using a Zeiss phase-contrast microscope. Cellular motility was observed in fresh wet mounts of young (24-h) bacterial cultures in TSB.

Most biochemical tests were performed by using the methods and media described by Gordon *et al.* (7). For all tests that required complex media (such as temperature and pH range of growth, growth inhibition in the presence of NaCl and in pH 5.7, hydrolysis of starch, resistance to lysozyme and liquefaction of gelatin), TSB or TSB agar were employed. Growth under anaerobic conditions was determined after incubation for 7 days in anaerobic Gaspak jars (BBL) containing an atmosphere of 80% N₂, 10% CO₂ and 10% H₂. Catalase activity was determined by bubble production in a 3% (v/v) hydrogen peroxide solution.

Strains were also characterized by using API tests (API 50CH and 20E, bioMérieux) as described in Seldin and Penido (30) and in each manual. Data from API 50CH tests composed of 49 different carbohydrates were recorded as described previously (28).

Analysis of respiratory quinones and fatty acids were carried out by the Identification Service and Dr. Brian Tindall, DSMZ, Braunschweig, Germany.

**Preparation of genomic DNA, PCR amplification, 16S rRNA gene sequencing and phylogenetic analysis**

Total DNA was extracted from the selected strain (P118) as described in Seldin and Dubnau (29). The procedure described by Massol-Deya *et al.* (17) was employed for PCR amplification of the 16S rRNA coding gene using the pair of universal primers (pA and pH). The amplification conditions were: a hot start (2 min 10 s at 92°C) and 35 cycles of 92°C (1 min 10 s), 48°C (30 s) and 72°C (2 min 10 s). A final extension step was run for 6 min 10 s at 72°C and the reaction tubes were then cooled to 4°C. DNA preparation and PCR products were visualized after electrophoresis respectively in 0.8% and 1.2% agarose gel in 1x TBE buffer (26).

PCR product of the 16S rRNA coding gene from P118 was sequenced by using an ABI Prism 3100 automatic sequencer. All sequences were identified using the Blastn facility (www.ncbi.nlm.nih.gov/blast) of the National Center for Biotechnology Information with GenBank non-redundant database. The 16S rRNA coding gene sequences of closely related strains were recovered from GenBank database and aligned with the sequences obtained in this study using the package software Clustal X (34). BioEdit version 7.0.5.3 (http://www.mbio.ncsu.edu/Bioedit/bioedit.html) was used for manual editing of the sequences.

Phylogenetic trees were constructed based on complete 16S rDNA sequences, using the Neighbor-Joining (NJ) method (25), whereas MEGA 4.1 software (15) was used to calculate pairwise p-distance values for 16S rRNA gene sequences among the different species studied here.

**RESULTS AND DISCUSSION**

The production of industrially important products from cellulosic biomass needs the bioconversion of the cellulosic components into fermentable sugars. A variety of microorganisms may have the ability to degrade the cellulosic
biomass to glucose monomers, and several efforts have been made to increase the production of cellulases through strain improvement. Some of these studies showed limited success (14) and the prospection of cellulolytic bacterial strains in singular environments has been cited in literature as an alternative approach for increasing cellulose production. For example, cellulose-degrading bacterial diversity was studied in an unusual environment like a weathered soil-like sample collected from the deep subsurface (1.5 km depth) of the Homestake gold mine in Lead, South Dakota, USA (22). Therefore, the prospection of cellulolytic bacterial strains in the gut of Parotocinclus maculicauda, used in this study, can be of great importance and utility. Because of the diet imposed to P. maculicauda, constituted mainly of plant material, the presence of cellulolytic bacteria was expected.

The food content from the digestive trait of the different fish samples was used for bacterial isolation. Different colonies were selected based on their morphotypes and were tested for cellulolytic activity. Although different isolates showed tiny halos around the bacterial growth in CMC plates, only one strain (designed P118) showed a clear and reliable halo around its spot growth. Because of its cellulolytic potential, strain P118 was further characterized taxonomically and the CMCase activity measured at different periods of time.

P118 cells are rod shaped, they grow and form spores aerobically, and they are catalase positive but oxidase negative. After 24 h in TSB, they occur either singly or in short chains and are motile. The cells are 2.9±0.4 x 0.9±0.1 μm. The spores are oval to ellipsoidal, predominantly subterminal to terminal and distend the sporangium. The minimum and maximum temperatures tested for P118 growth are 15 and 40°C, respectively (the optimum temperature for growth ranged from 25 to 32°C). No growth occurs at 10 or 45°C. P118 grows in TSB with pH varying from 5.0 to 9.5 (the optimum pH for growth ranged from 8-8.5). P118 grows in the presence of 2% NaCl but does not grow with 3% NaCl. Growth occurs in the presence of 0.001% lysozyme. The strain is facultative anaerobic and presents a Voges-Proskauer negative test. Nitrate is reduced to nitrite. Considering the results obtained using the API 50CH kit, acid is produced from L-arabinose, ribose, D-xylose, D-galactose, D-glucose, D-fructose, D-mannose, D-mannitol, amygdalin, arbutin, esculin, salicin, D-cellobiose, D-maltose, lactose, melibiose, sucrose, trehalose, raffinose, gentiobiose and D-turanose. Weak production of acid is observed when glycerol, D-arabinose and β methyl-D-xyloside are the carbohydrates tested. No acid is observed from erythritol, L-xylose, adonitol, L-sorbose, L-rhamnose, dulcitol, inositol, D-sorbitol, α methyl-D-mannoside, α methyl-D-glucoside, N-acetyl-glucosamine, inulin, melezitose, starch, glycogen, xylitol, D-lyxose, D-tagatose, D-fucose, L-fucose, D-arabitol, L-arabitol, gluconate, 2 keto-gluconate and 5 keto-gluconate. Strain P118 is able to hydrolyze 1% tween 80 and casein (weak test), but not gelatin. Does not utilize gluconate and caprate and produces urease.

Comparison of nearly complete 16S rRNA gene sequence of strain P118 with those in GenBank indicated that the novel strain was phylogenetically related to members of the genus Paenibacillus. P118 displayed highest levels of 16S rRNA gene sequence similarity with P. amyloyticus NRRL NRS-290 (98.8%), P. pabuli HSCC 492 (98.6%), P. tundrae Ab10b (98.5%), P. xylanexedens B22a (98.3%) and P. tylopili MK2 (98.3%). Phylogenetic tree constructed using the neighbor-joining (Fig. 1) and maximum-parsimony methods were concordant with one another (not shown). P118 formed a cluster with the five above mentioned species. When other cellulolytic Paenibacillus species were used for comparison (P. campinasensis – (12); P. phyllophaeraeae – (24); P. cellulosilyticus – (23); P. cellulositrophicus – (1); P. prosopidis – (35); P. piniihumi – (10); P. pini – (11); P. curdianolylticus – (21)) levels of 16S rRNA gene sequence similarities were below 96%. These species were included in the phylogenetic tree to demonstrate the distance between P118 and the recognized cellulolytic species (Fig. 1). No strain belonging to the five closest species to P118 has been described as able to degrade cellulose so far.
Profiles of the major fatty acids of strain P118 are shown in Table 1. Anteiso-C<sub>15:0</sub>, the major fatty acid in recognized members of the genus *Paenibacillus*, was also the major fatty acid component of strain P118 (38.73%) followed by C<sub>16:0</sub> (20.85%). For the menaquinone profile, the MK-7 was the predominant one and this is in agreement with the description of the *Paenibacillus* genus (2).

Table 2 shows the characteristics that differentiate strain P118 from the closest type strains of *Paenibacillus* chosen based on phylogenetic data (16S rRNA sequence). They clearly demonstrate that strain P118 does not match to any of these *Paenibacillus* species and may represent a novel species of the genus. It is important to highlight the pH range in which P118 grows and the resistance to lysozyme (0.001%), besides its cellulolytic property. However, further studies, such as DNA-DNA hybridization, are still necessary to define a new species.

Table 1. Cellular fatty acid profiles (%) of strain P118

| Fatty acid     |           |
|----------------|-----------|
| Iso-C<sub>13:0</sub> | 0.21      |
| Anteiso-C<sub>13:0</sub> | 0.25      |
| Iso-C<sub>14:0</sub> | 5.24      |
| C<sub>14:0</sub> | 12.36     |
| Iso-C<sub>15:0</sub> | 8.53      |
| Anteiso-C<sub>15:0</sub> | 38.73     |
| C<sub>15:0</sub> | 0.58      |
| Iso-C<sub>16:0</sub> | 3.93      |
| C<sub>16:1 w11c</sub> | 7.02      |
| C<sub>16:0</sub> | 20.85     |
| Iso-C<sub>17:0</sub> | 1.31      |
| Anteiso-C<sub>17:0</sub> | 0.99      |
Table 2. Some characteristics that differentiate strain P118 (isolated in this study) from the closest type strains of *Paenibacillus* chosen based on phylogenetic data (16S rRNA sequence)

| Characteristic          | P118   | *P. amylolyticus* \(a\) | *P. tundrae* \(b\) | *P. xylanexedens* \(c\) | *P. pabuli* \(d\) | *P. tylopili* \(e\) |
|-------------------------|--------|--------------------------|---------------------|--------------------------|-------------------|---------------------|
| Nitrate reduction       | +      | +                        | +                   | -                        | -                 | -                   |
| Hydrolysis of:          |        |                          |                     |                          |                   |                     |
| Casein                  | +w     | +w                       | nd                  | nd                       | v\(\ast\)          | -                   |
| Starch                  | -      | +                        | nd                  | nd                       | +                 | +                   |
| Gelatin                 | -      | +                        | nd                  | nd                       | nd                | -                   |
| Tween 20 (1% v/v)       | -      | +                        | nd                  | nd                       | nd                | nd                  |
| Growth in the presence of: |        |                          |                     |                          |                   |                     |
| 5% NaCl                 | -      | -                        | nd                  | nd                       | v\(\ast\)          | v                   |
| 0.001% lysozyme         | +      | -                        | nd                  | nd                       | -                 | nd                  |
| Conditions for growth:  |        |                          |                     |                          |                   |                     |
| pH range                | 5 – 9.5| 4.5 – 9                   | 5.2 – 8.8           | 5.7 – 8.8                | nd                | 6 – 9               |
| pH optimum              | 8 – 8.5| 7                        | 6.4                 | 6                        | nd                | 8                   |
| Temperature range (\(^\circ\)C) | 15 – 40 | 10 – 40                  | 13 – 37             | 13 – 32                  | 5 – 40            | 9 – 40 \(v\)        |
| Temperature optimum (\(^\circ\)C) | 25 – 30 | 37                       | 27                  | 23                       | 28 – 30           | 25                  |
| Urease                  | +      | -                        | nd                  | nd                       | -                 | -                   |
| Acid production from:   |        |                          |                     |                          |                   |                     |
| L-arabinose             | +      | +                        | -                   | -                        | +                 | +                   |
| D-xylose                | +      | +                        | +                   | -                        | +                 | +                   |
| Inositol                | -      | +                        | -                   | -                        | nd                | -                   |
| D-sorbitol              | -      | -                        | +                   | -                        | -                 | nd                  |
| \(\alpha\)-methyl D-glucoside | -      | +                        | +                   | -                        | +                 | nd                  |
| Lactose                 | +      | -                        | +                   | +                        | +                 | +                   |
| Inulin                  | -      | +                        | +                   | -                        | +                 | nd                  |
| Melezitose              | -      | +                        | +                   | +                        | nd                | -                   |
| Raffinose               | +      | -                        | +                   | +                        | +                 | +                   |
| DNA G+C content (mol%)  | 45.8\(\ast\) | 46.3 – 46.6           | 50.3                | 46.4                     | 48 – 50           | 44.3                |
| Major cellular fatty acids | anteiso-C\(_{15:0}\) | anteiso-C\(_{15:0}\) and C\(_{16:0}\) | anteiso-C\(_{15:0}\) and C\(_{16:0}\) | anteiso-C\(_{15:0}\) and C\(_{16:0}\) | anteiso-C\(_{15:0}\) and C\(_{16:0}\) |

\(\text{nd, not determined; } v, \text{ variable result; data from (a) Shida et al. (32), (b) and (c) Nelson et al. (20), (d) Nakamura (19) and Kuisiene et al. (13), (e) Kuisiene et al. (13). }\)\(\ast\) type strain hydrolyzes casein and is inhibited by 5% NaCl; \(\ast\) Dr. Peter Schumann, DSMZ, Braunschweig, Germany determined the DNA base composition of P118 by HPLC (33).

Strain P118 showed to be a cellulolytic bacterium, able to produce CMCase (Fig. 2). The time-course fermentation profile, when CMC was the sole carbon source for P118 growth, indicates a maximum activity at the first day (0.154 U/ml). Data concerning cellulase activity in *Paenibacillus* are scarce in literature, and very often those available cannot be used for comparison since different conditions, methodology for enzymatic activity assays and units are used. However, our results were comparable to those obtained by Ko *et al.* (12) for *P. campinasensis* BL-11, which presented an activity of 0.1 U/ml in LB medium plus xylan oat spelts. Furthermore, P118 showed a higher cellulase activity than *P. curdlanolyticus* B-6 grown in Berg medium containing 0.5% oat spelts (21). In fact, considering the biotechnological potential of P118, higher values for cellulase production can be found in literature for other bacterial genera, fungi and actinomycetes. However, it is important to consider that these values can be improved. When Semêdo *et al.* (31) first described the cellulolytic strain *Streptomyces drozdowiczii* M7a, its cellulase activity showed to be around 0.1U/ml. Changing the medium used and the culture conditions, an increase of 6 times in the enzyme production was observed (8). Moreover, it is important to state that P118 showed its maximal cellulase activity in the very first day of the experiment, which can be considered a very important characteristic for an economical point of view.

In conclusion, a novel and promising *Paenibacillus* strain isolated from *P. maculicauda* gut is now available for further studies aiming the optimization of its cellulase production, facilitating its potential use in industrial applications.
Figure 2. Carboxymethylcellulase (CMCase) activity at different periods of time. Results were expressed as the average of two replicates. One unit (U) of CMCase activity corresponded to 1 µmol of glucose equivalent released per minute under the assay conditions.

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