

# ISOLATION AND MOLECULAR IDENTIFICATION OF A CELLULOTIC BACTERIUM FROM MUNICIPAL WASTE AND INVESTIGATION OF ITS CELLULASE PRODUCTION

## ISOLAMENTO E IDENTIFICAÇÃO MOLECULAR DE UMA BACTÉRIA CELULOLÍTICA DO LIXO MUNICIPAL E INVESTIGAÇÃO DE SUA PRODUÇÃO DE CELULASE

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**ABSTRACT:** Municipal waste is rich in lignocellulosic compounds which contain cellulose, lignin and hemicellulose. Microorganisms can break down such compounds and convert them into glucose and other carbohydrates. The current study was performed to isolate and identify cellulolytic bacteria in municipal waste. Municipal waste samples were collected and plated on Carboxymethyl cellulose (CMC) agar. Preliminary identification of the isolates was performed using standard biochemical assays. The activity of carboxymethyl cellulose (CMCase) was specified through measuring the release of reducing sugars from CMC. Different nitrogen sources at various concentrations and initial pH values were evaluated for their effect on enzyme production. Further the enzyme production was determined at different fermentation times. Molecular identification was then performed using bacterial 16S rRNA gene amplification and sequencing. A cellulolytic bacterium was isolated from municipal waste samples and identified based on morphological, physiological and biochemical characteristics along with 16S rRNA analysis. The isolated bacterium was identified as *Bacillus subtilis* (accession number: KU681044). Whose growth characteristics showed that its growth curve entered the logarithmic phase following 10–18 h with the stable growth phase ranging from 23 to 37 h. The optimal carbon source for fermentation was 1% rice hull, with the nitrogen source comprised of 2% peptone and yeast extract. The the minimum CMCase activity was observed at an initial medium pH of 4.0, while the maximum was observed at pH 7. The strain grew vigorously and the cellulase yield was high at 6–24 h fermentation time period. The isolated bacteria showed the degrading potential of cellulose which could be employed in local industrial process.

**KEYWORDS:** *Bacillus subtilis*. Carboxymethyl cellulose. KU681044. Lignocellulosic compounds. Municipal waste.

## INTRODUCTION

The drastic augment in world population and the pace of urbanization, has led to an increase in municipal waste production. Recycling municipal waste in a natural way is the most novel and appropriate alternative considering other available options. Municipal waste is rich in Lignin-cellulosic material that basically contains cellulose and lignin and a low proportion of hemicelluloses (RAHMAN, 2004). Microorganisms can break down lignin-cellulosic materials, converting them into valuable products such as organic acids and antibiotics.

Cellulose is a polysaccharide with a crystalline form, which consists of repeating units of D- glucose linked together via  $\beta$  1- 4 glycoside bonds (BHAT, 2000). The microbial degradation of cellulosic compounds is caused by an enzyme called cellulase, which hydrolyses the  $\beta$  1- 4 glycoside bonds of cellulose in order to release glucose units. Widely applied in different industries including

food, feed, textile and pulp, cellulases have gained great significance in present day biotechnology. The bioconversion of cellulosic materials is currently a subject of intensive research as a contribution to the development of large scale conversion process beneficial to mankind (BEHERA et al., 2014).

Several bacterial and fungal species, such as aerobic and anaerobic bacterial and fungal strains, researched and reported. Cellulolytic bacterial species include *Trichonympha*, *Clostridium*, *Actinomyces* and *Ruminococcus species* (GUPTA et al., 2012).

Despite the huge significance of cellulase in different industries, little is known as to the cellulolytic microbial species inhabiting municipal waste, hence the current research which was performed to isolate and identify bacterial species with cellulose degrading potential in municipal waste landfill located in Saravan region, Rasht city, Iran.

## MATERIAL AND METHODS

### Sample collection

The samples were collected from municipal waste landfill located in Saravan region, Rasht city, Iran. Sampling was performed at depths of 5-100 cm and the samples were collected in sterile container and stored at 4°C until used.

### Screening of cellulose producing bacteria

Soil samples were serially diluted in sterilized distilled water, of which 0.1 ml was plated on Carboxymethyl cellulose (CMC) medium which had the following composition (g/L) (Atlas R M 2010): carboxymethylcellulose (CMC), 10; K<sub>2</sub>HPO<sub>4</sub>, 1.0; KH<sub>2</sub>PO<sub>4</sub>, 1.0; MgSO<sub>4</sub>·7H<sub>2</sub>O, 0.2; NH<sub>4</sub>NO<sub>3</sub>, 1.0; FeCl<sub>3</sub>·6H<sub>2</sub>O, 0.05; CaCl<sub>2</sub>, 0.02; peptone, 2.0; Agar, 15.0. The incubation was performed at 37 °C for 3-5 days and the bacteria with cellulose degrading potential were screened by flooding of 0.1% Congo red solution on the plates and washing with 1.0M NaCl (APUN et al., 2000).

The formation of a clear zone around the bacterial colonies was regarded as cellulolytic activity. The cellulolytic activity was assessed through measuring the ratio of the clear zone diameter to colony diameter (ARIFFIN et al. 2006). The largest ratio was assumed to have the highest activity. The strain with high clearing zone was isolated for repeated screening. The strain (inoculum concentration of 1%, 50 mL in a 250 mL flask) was inoculated in liquid LB medium containing 1% CMC at 37°C and grown with shaking at 220 rpm for 24 h. Subsequently, the fermentation extract was centrifuged at 5000 rpm for 15 min, and the apparent supernatant was examined for enzyme assays under optimum reaction conditions to screen most cellulolytic bacterial strains. Strains demonstrating giant cellulase activity were used in subsequent experiments.

### Morphological and physiological characterization of cellulolytic isolates

Once the pure culture of the isolates was prepared, the bacterial isolates were initially identified by morphological examination and biochemical characterization. The parameters investigated were bacterial shape, Gram reactions, endospore formation, catalase production, Voges-Proskauer (V-P) reactions, starch hydrolysis, motility and citrate utilization.

### Effect of different fermentation conditions on cellulase production

In order to specify the optimal carbon and nitrogen source to intensify cellulase production, different carbon and nitrogen sources at various concentrations and initial pH values were assessed for their effect on enzyme production. Also simultaneously enzyme production at different fermentation times was dogged. Cultures (50 mL in 250 mL flasks) were incubated at 37°C on a rotary shaker at 220 rpm for 24 h, the fermentation extract was centrifuged at 5000 rpm for 15 min, and the apparent supernatant was employed in the enzyme assays.

### Molecular identification of the isolates

Bacterial genomic DNA was extracted using the previously described method with a minor modification (DAS; DASH, 2015). Briefly, 5 mL of Bacterial suspensions grown in Luria–Bertani (LB) broth were centrifuged (6000g /10 min) and the pellet was re-suspended in TE buffer. Then, 30 µL of 10% SDS and 3µL of 20 mg/mL proteinase-K were added to each tube and incubated for 60min at 37°C. Next, 600µL of 25:24:1 phenol/chloroform/isoamyl alcohol solution was added and centrifuged at 6000 g for 5 minutes. Finally, one volume of cold ethanol was added to the supernatant and the DNA was precipitated following centrifugation (10 min at 8,000 g). The DNA yield was re-suspended in TE buffer and stored at -20°C for subsequent analysis.

Polymerase Chain Reaction (PCR) was employed to amplify bacterial 16S rRNA gene. The master mix for the PCR was prepared as follows: 3µL of 10× PCR buffer, 1µL of 25mM MgCl<sub>2</sub>, 3µL of 10mM dNTP mix, 0.5µL of Taq DNA Polymerase, 12.5µL of MilliQ water and 1µL of the forward and reverse primers. Ultimately, 3µL of each DNA template was added in the corresponding tubes to make up the final reaction volume of 25µL.

Bacterial 16S rRNA gene was amplified using 27F (5' AGAGTTTGATCCTGGCTCAG 3') and 1107 R (5' AAGGTTACCTCACC GACTTC 3') universal primers (LANE, 1991). The thermal cycler was programmed as follows: 5 min at 94°C; 30 cycles of 45s at 94°C, 1 min at 58°C, 1 min at 72°C and 10 min at 72°C. The PCR product was sequenced for 16S rRNA (Macrogen, South Korea). The sequencing results were compared via applying the Basic Local Alignment Search Tool (BLAST) program to NCBI and 16S rRNA gene sequence homology analysis using GenBank data. A phylogenetic tree was constructed using the neighbor-joining model of the MEGA 5 program.

## RESULTS AND DISCUSSION

In the present study, the isolated species were cultured at 37°C on CMC-Na solid media for 24 h, where the subsequent Congo red staining showed a remarkable clearing zone. Furthermore, by testing the enzyme activity, we selected the strain demonstrate the highest enzyme activity for upcoming research. According to preceding

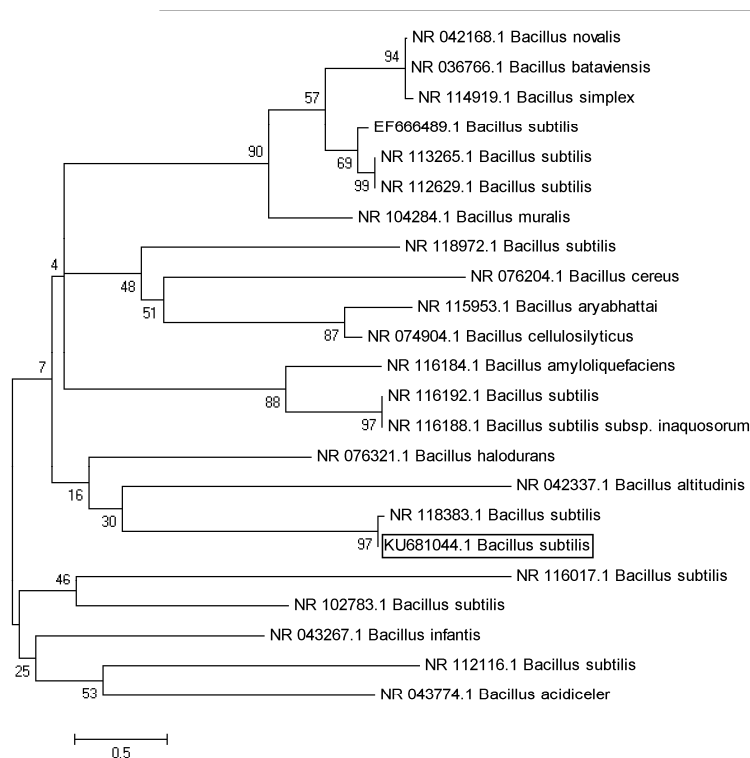
research, the enzyme activity of this strain (2.1 U/mL) was higher than that of other strains cultured for 24 h (SINGH et al., 2013; IMMANUEL et al., 2006; REDDY et al., 2017) suggesting that this strain exhibits steady inheritance and has strong CMCcase activity. The morphological and physiological characteristics of isolated strain were further determined (Table 1), according to which results, the strain was Gram positive rod.

**Table 1.** Morphological and physiological characterizations of the isolated strain.

| Characters | shape | Gram stain | spore | Catalase Test | VP | Aerobic growth | motility | Citrate utilization | starch hydrolysis |
|------------|-------|------------|-------|---------------|----|----------------|----------|---------------------|-------------------|
| CS1        | Rod   | +          | +     | +             | +  | +              | -        | +                   | +                 |

The 16S rRNA sequence of the cellulolytic bacterium was 1511 bp long. This sequence was compared with the database of known 16S rRNA sequences. Homology analysis showed that the sequence similarity of this strain to certain *Bacillus* species exceeded 95%. Thus, all characterization methods indicated that the isolated species was a *B. subtilis* strain (KIM et al., 2009). The 16S rRNA gene sequence of the isolate was submitted to the

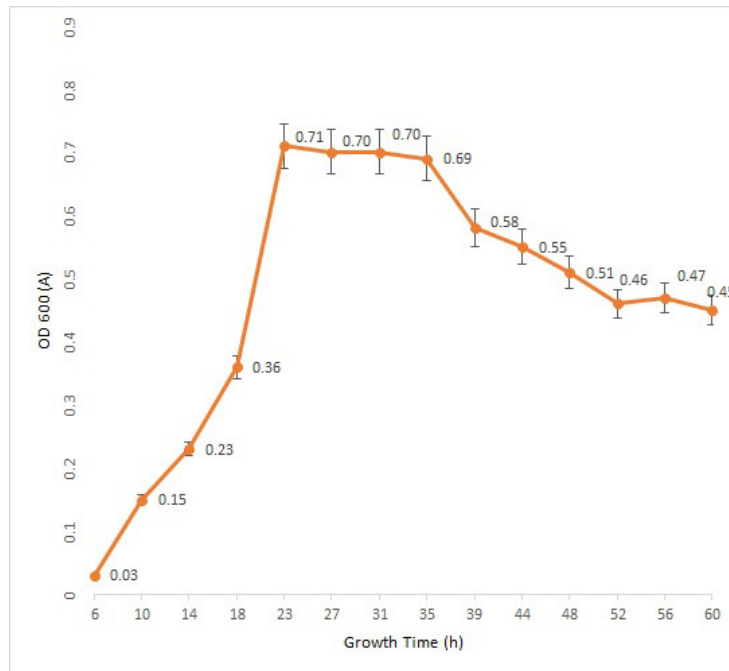
GenBank with KU681044 accession number. As shown in the phylogenetic tree constructed using MEGA5, the strain is related to *B. subtilis* (GenBank accession number: NR\_118383.1), with a similarity of 97 %. Therefore, all characterization methods showed that the isolated species was a *B. subtilis* strain. The phylogenetic tree of the isolated bacterium is illustrated in Figure 1.



**Figure 1.** Phylogenetic analysis of cellulolytic strain isolated in this study (Accession number: KU681044). Bootstrap values based on 500 replications are given at the branching points.

The strain attained the logarithmic phase at 10–18 h after inoculation, after which its growth hit

a plateau and proceeded for approximately 20 h (Figure 2).

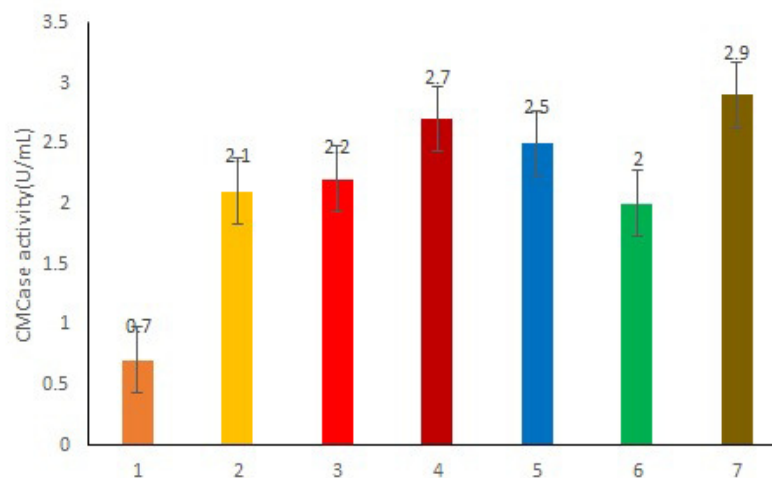


**Figure 2.** Growth curve of isolated bacteria. Error bars indicate standard deviation.

According to the strain growth characteristics, we concluded that the optimum enzyme production time was 18–35 h. Considering the fact that bacterial enzyme is most active in the logarithmic phase, we organized fermentation cultivation for 24 h in a follow-up experiment in order to investigate the enzyme features.

Since cellulase is an inducible enzyme, the medium for cellulose production in fermentation usually contains cellulose-rich substrates for a carbon source (LEE et al., 2008; DORSAM et al. 2017; ZHAO). In this medium, different carbon sources were tested at different concentrations in order to study their effects on cellulase production

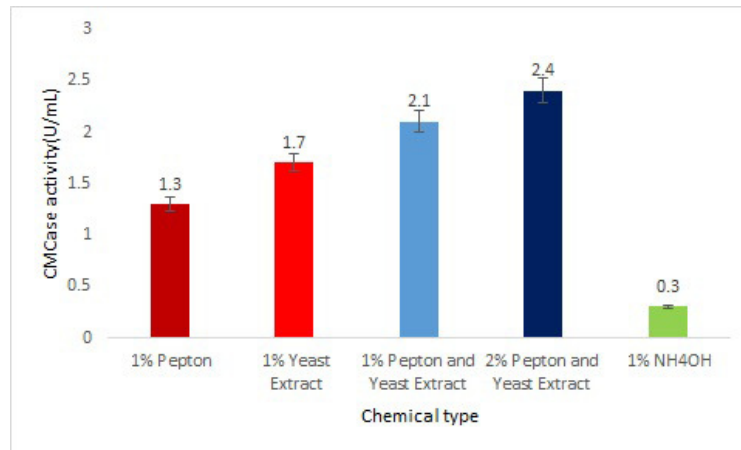
under uniform conditions (incubation time, 24 h; rotation speed, 220 rpm; temperature, 37°C; seeded culture concentration, 1%). The results showed that the strain could employ varied carbon sources, and the maximal CMCase activity (2.9 U/mL) was detected in a mixture of 1% rice hull and 1% CMC utilized as the sole carbon source. In accordance to Yang et al., however, when MCC was used as a sole carbon source, slight CMCase activity was observed (YANG et al., 2014). Consequently, the cheap and easily accessible rice hull was found to be an optimal carbon source applicable in subsequent experiments (Figure 3).



**Figure 3.** Effect of different carbon sources on cellulase production(1: 1% MCC; 2: 1% CMC+1% Wheat Bran; 3: 1% Wheat Bran; 4: Rice hull; 5: 1% CMC; 6: 2% CMC; 7: 1% CMC+Rice hull) . Error bars indicate standard deviation.

Different nitrogen sources (1% yeast extract, 1% peptone, a 1:1 mixture of 1% peptone and yeast extract, a 1:1 mixture of 2% peptone and yeast extract, and 1% NH<sub>4</sub>OH) were examined. The results showed that the strain can efficiently exploit organic nitrogen sources. Moreover, the maximum

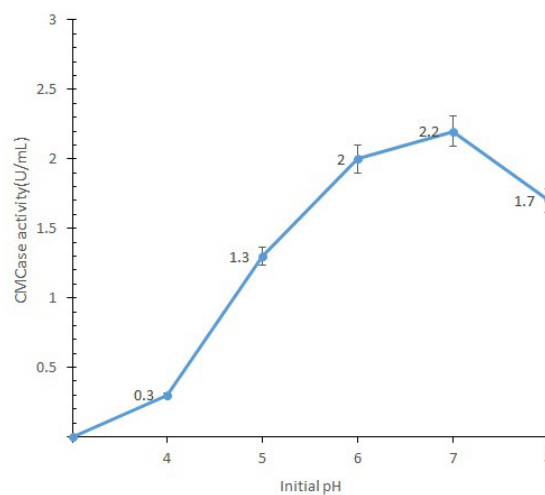
CMCase activity (2.4 U/mL) was observed when a 1:1 mixture of 2% peptone and yeast extract was used as the only nitrogen source. Nevertheless, the CMCase activity was nearly zero with inorganic nitrogen sources (NH<sub>4</sub>OH) (Figure 4).



**Figure 4.** Effect of different nitrogen sources on cellulase production. Error bars indicate standard deviation.

The reason is that the metabolism of inorganic nitrogen conduces to medium acidification, which in turn affects cellulase production. The *Bacillus* species isolated by Rajoka (2014) and Ray et al. (2007) showed the same capability with regards to nitrogen sources. In addition, the *B. subtilis* strain isolated by Acharya and Chaudhary (2011) was not capable of employing inorganic nitrogen sources when CMC was used as the carbon source; it was, however, able to employ these sources when wheat straw and rice hull were used as the carbon source.

To investigate the effect of pH on cellulase production, the pH of the medium was adjusted to 3.0, 4.0, 5.0, 6.0, 7.0, and 8.0. The CMCase activity in each pH-adjusted medium was then measured under identical conditions (rotation speed, 220 rpm; temperature, 37°C; seeded culture concentration, 1%; incubation time, 24 h). The minimum CMCase activity (0.3 U/mL), based on the results, was seen at an initial medium pH of 4.0, while the maximum (2.2 U/mL) was observed at pH 7 (Figure 5).

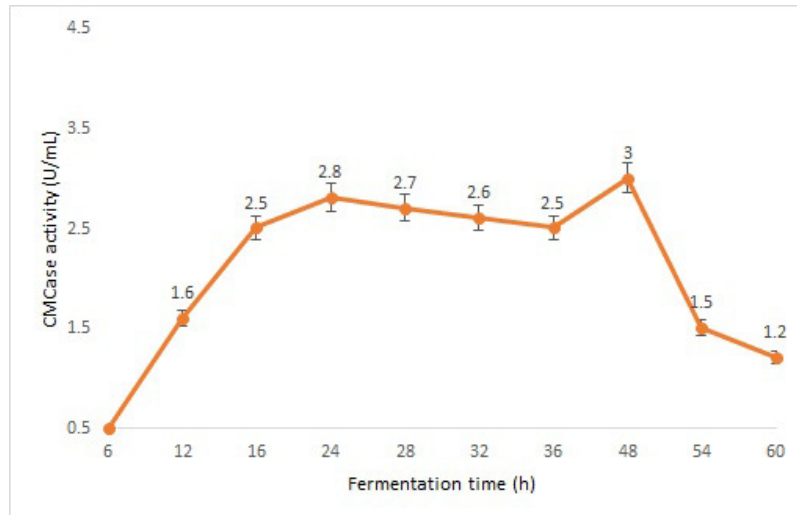


**Figure 5.** The effect of initial pH on cellulase production. Error bars indicate standard deviation.

The optimal initial pH of strains varies depending on the source: for instance, the optimal initial pH for cellulase production by *Bacillus* isolated from a hot spring (ACHARYA; CHAUDHARY, 2011) was 9.0. Accordingly, suitable pH conditions can be conducive to the growth of strains and the increase in cellulase yield, a fact probably associated with the negative feedback mechanism of enzymes.

At various fermentation times the CMCase activity was measured at 37°C, 5.5 pH and 220 rpm

with the use of optimal carbon and nitrogen sources. The findings showed significant variations in cellulase production at different times. The strain grew strongly and the cellulase yield was high when the fermentation time fell between 6–24 h. Suitable CMCase activity (2.8 U/mL) was perceived following 24 h cultivation; a reduction, nevertheless, was observed in this activity from 24 h to 36 h. The CMCase activity reached its maximum (3.0 U/mL) when the fermentation time was 48 h, following which time, it decreased sharply (Figure 6).



**Figure 6.** The effect of fermentation time on cellulase production. Error bars indicate standard deviation.

The fermentation time of the strain concerning the maximum cellulase yield was shorter than that of the *Bacillus* strains isolated by Rastogi et al. (2010), Sadhu et al. (2013) and comparable to that of *Bacillus* species isolated by Acharya and Chaudhary (2011).

Cellulose, as the main structural part of plants, is the most abundant organic carbon on earth. The organic carbon in the environment is mainly

recycled by microorganisms (CLEMMER; TSENG, 1986; SI et al., 2017). On the other hand, the degradation of cellulose is a complex process and requires the cooperation of cellulolytic microbial enzymes. Successful biological conversion of cellulose-containing compounds depends on the nature of the cellulose, cellulolytic enzyme source, and the optimum conditions for catalytic activity and enzyme production.

**RESUMO:** Resíduos urbanos são ricos em compostos lignocelulósicos que contêm celulose, lignina e hemicelulose. Microrganismos podem quebrar esses compostos e convertê-los em glicose e outros carboidratos. O presente estudo foi realizado para isolar e identificar bactérias celulolíticas em resíduos urbanos. Amostras de resíduos municipais foram coletadas e plaqueadas em ágar Carboximetilcelulose (CMC). A identificação preliminar dos isolados foi realizada utilizando ensaios bioquímicos padrão. A atividade da carboximetilcelulose (CMCase) foi especificada através da medição da liberação de açúcares redutores da CMC. Diferentes fontes de nitrogênio em várias concentrações e valores iniciais de pH foram avaliados quanto ao seu efeito na produção de enzimas. Além disso, a produção de enzima foi determinada em diferentes tempos de fermentação. A identificação molecular foi então realizada utilizando amplificação e sequenciamento do gene bacteriano 16s rRNA. Uma bactéria celulolítica foi isolada de amostras de resíduos urbanos e identificada com base em características morfológicas, fisiológicas e bioquímicas, juntamente com a análise 16S rRNA. A bactéria isolada foi identificada como *Bacillus subtilis* (número de acesso: KU681044). Cujas características de crescimento mostraram que sua curva de crescimento entrou na fase logarítmica após 10-18 h com a fase de crescimento estável variando de 23 a 37 h. A fonte de carbono ótima para a fermentação foi 1% de casca de arroz, com a fonte de nitrogênio composta de 2% de peptona e extrato de levedura. A atividade mínima de CMCase foi observada em um pH médio inicial de 4,0, enquanto a máxima foi observada em pH 7. A linhagem cresceu vigorosamente e o rendimento de celulase foi alto no período de 6 a

24 horas de fermentação. As bactérias isoladas mostraram o potencial de degradação da celulose que poderia ser empregada no processo industrial local.

PALAVRAS-CHAVE: *Bacillus subtilis*; Carboximetilcelulose; KU681044; Compostos lignocelulósicos; Resíduos municipais

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