Test for the production and assay of the proteolytic activities of halophilic bacteria and archaea isolated from Algerian hypersaline environments

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Abstract: The present work was carried out on 133 halophilic strains, isolated on MGM medium at 12 and 23% (w/v) of salt. A screening of the extracellular proteolytic activities, carried out on the same medium supplemented with casein or gelatin at 1% (w/v), allowed us to select 24 bacterial and 21 Archaeal strains presenting a precipitate around the colonies for casein and/or a translucent halo (after addition of Frazier’s reagent) for the gelatin. The enzymatic test was done on liquid medium in micro-culture on a 2 mL Eppendorf tube. The assay of the proteolytic activity using Azocasein as substrate, following 2 protocols the first with PBS and the second with Tris HCl, with positive and negative controls, demonstrated interesting results for 10 strains among the 45 tested including 5 bacteria and 5 archaea. These have undergone morphological, physiological and molecular characterization based on amplification and sequencing of the 16S ribosomal RNA gene.

Keywords: halophilic bacteria; halophilic archaea; proteolytic activity; hypersaline environments

1. Introduction

Microbial life can be found over a wide range of extreme conditions (salinity, pH, temperature, pressure, light intensity, oxygen and nutrient conditions). Hypersaline environments constitute typical examples of environments with extreme conditions due to their high salinity, exposure to high and low temperatures, low oxygen conditions and in some cases, high pH values. Bacteria and Archaea are the most widely distributed organisms in these environments [1]. These unconventional conditions for life suggest that such microbes use original strategies (specific enzymes) to adapt to physico-chemical constraints that they have to face. Several of these enzymatic activities are nowadays good candidates to be used in biotechnological processes [2]. However, in comparison to other groups of extremophilic microorganisms such as the thermophiles and the alkaliphiles, the halophiles of all three domains have been relatively little exploited [3].

Proteases represent the largest and most important segment in the industrial enzyme market, used in detergents, food processing, and leather industry, as biocatalysts in organic synthesis, and, among many other applications, as therapeutics because their roles are involved in key decisions throughout an organism in several physiological and metabolic processes [4]. Therefore, obtaining
and preparing pure enzyme which is stable and active under multiple extreme conditions (alkaline pH, high salt concentrations and wide temperature) is scientifically and industrially significant [5].

2. Materiel and methods

2.1. Samples and strains isolation

Brine samples used in the present study were collected from two solar saltern on 2016, namely Ezzemoul, Ain M’lila east of Algeria and Betioua, Aarzew west of Algeria. Samples were spread on solid Modified Growth Medium [6], containing 5 g peptone, 1 g yeast extract with 12.5 and 23% (w/v) salt concentration, pH 7.5. Prokaryotic isolates were picked after 7–15 days of incubation at 37 °C, based on colony pigmentation, size, and margin and immediately streak-plated at least 3–4 times on fresh agar plates with the appropriate medium until purity was confirmed [7]. Beside this, 41 strains were recovered from previous work. A total of 133 strains (68 bacteria and 65 Archaea) were preserved on plats at 4°C.

2.2. morphology and physiology

The isolates were submitted to morphology examination using cells from exponentially growing cultures. The colonies’ aspect was examined. Cell morphology and Gram was observed. The proteolytic activity of the isolates was screened on solide MGM media (0.3 % peptone and 0.1% yeast extract) supplemented with 1% (w/v) casein [8], or 1% (w/v) gelatin [9], positive results was detected after, 5 days for bacteria and 10 days for archaea, incubating at 37 °C by the presence of a precipitate around the colonies for casein and/or a translucent halo (after addition of Frazier’s reagent) for the gelatin. Growth rates were estimated on MGM liquide medium at different NaCl concentrations (0-2.5- 5- 7.5- 10- 12.5- 15- 17.5- 20- 22.5- 25 and 27%). The culture was incubated in aerobic conditions at 37°C during 24h for bacteria and 4_h for archaea. The optical density at 660 nm was measured with a Synergy H1 hybrid multi-mode microplate reader [10].

2.3. DNA extraction, 16S rRNA gene amplification and strains identification

The total genomic DNA of 10 proteolytic selected strains, was prepared from colonies resuspended in MilliQ water by heating at 98°C/10 min for bacteria and at 72°C/10 min for Archaea, followed by centrifugation at 13000 rpm for 5 min [6]; The tubes were immediately cooled on ice. Bacterial and archaeal 16S rRNA genes were amplified using primer sets 27F/1492R [11] and 21F/1492R [12], respectively. PCR was performed using a 50µL reaction mixture containing the following (per reaction): 5µL of PCR buffer 10x, 4 µL of deoxyribonucleotide triphosphate mixture (10mM each), 0.5 µL of each primer (10µM), 0.25µL (1.25 U) DreamTaq, and 1µL of template DNA. The following PCR conditions were used: 95 °C for30 s followed by 40 cycles of 95 °C for 30 s, 45,2 °C for 30 s, and 72 °C for 1 min, and finally an extension step of 8 min at 72 °C. The amplified DNA supplied with gel-Green was submitted to electrophoresis on 0.7% (w/v) agarose gels in (TE) buffer and then visualized under UV light. Identities with described taxa were investigated using the nBLAST tool against the EzBioCloud database of cultured organisms.

2.4. Enzymatic assay

The enzymatic assay was carried out using two methods based on azocasein (Sigma, St. Louis, MO, USA) as substrate. Absorbance was performed using a plate reader (using a Synergy H1 Hybrid Multi-Mode Microplate Reader).

Method 1: Cell-free culture supernatants were prepared by centrifugation for 5 min at 12,000 g. The reaction was performed in 50 mM Phosphate-buffered saline (PBS) solution pH 7.5 with 50 µl of azocasein 3% (w/v) and with 50 µl of culture supernatant for a final volume of 750 µl. The reaction was incubated at 37°C for 1 h and stopped by adding 125 µl of 20% (w/v) trichloroacetic acid. Positive control is prepared by using a solution of proteinase K, the blank assay was realized using the culture medium. After centrifugation at 15,000 g for 10 min, the absorbance of the supernatant was measured...
at OD366nm. One unit of protease activity was defined as the amount required to produce enough acid-soluble material from azocasein to yield an absorbance of 0.01 at 366 nm, following 1 h of incubation [13, 10].

Method 2: In this assay, 0.5 mL of reaction mixture consisting of 0.25 mL of crude enzyme (cell-free supernatant) and 0.25 mL of 0.8% (w/v) azocasein in 0.1 M Tris-HCl buffer (pH 8.0) containing NaCl at 12.5% (w/v) for bacteria and 23% (w/v) for archaea. The reaction mixture was incubated at 37°C for 2 h with mild shaking. The reaction was terminated by adding 0.5 mL of 10% (w/v) trichloroacetic acid and the mixture was allowed to stand at room temperature for 30 min. The precipitate was removed by centrifugation at 10,000 rpm for 10 min. Subsequently, a volume of 300 mL of the initial supernatant fluid was transferred into a microcentrifuge tube and then mixed with 350 mL of 1.0 N NaOH. The absorbance was measured at 440 nm. One unit (U) of halophilic protease activity was defined as the amount of enzyme activity that produces a change in absorbance of 0.01 at 440 nm in 2 h at 37°C under the standard assay conditions. The halophilic protease activity was calculated according to the following equation:

\[
\text{Halophilic protease activity (U/µl)} = \frac{(A-B) \cdot V_t}{V_e \cdot 0.01}
\]

Where A and B are the optical densities of the crude enzyme and the control, respectively, \(V_t\) is the total reaction volume and \(V_e\) is the volume of crude enzyme [14, 15].

3. Results

3.1. Morphology and physiology

Morphological aspect of cells for 45 proteolytic strains showed that most of archaea are Gram positive with cocci dominant, and for bacteria we find the two types of Gram with rod-shaped are the dominant as shown in table 1.

<table>
<thead>
<tr>
<th>Gram</th>
<th>Cell form</th>
<th>Protease</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Positive</td>
<td>Negative</td>
</tr>
<tr>
<td>Bacteria</td>
<td>8</td>
<td>16</td>
</tr>
<tr>
<td>Archaea</td>
<td>1</td>
<td>17</td>
</tr>
</tbody>
</table>

The growth rates of selected strains were estimated as described before. Results in figure 1, demonstrate that the bacteria strain 4 have a large salinity growth interval (2.5 – 17.5%, w/v), which mean that it is a moderately halophilic bacteria, in addition, the archaea strain N have a growth salt interval from 15 to 27% (w/v), suggesting that it is an extreme halophile [16].

![Figure 1](image)

Figure 1. salt optima for selected bacteria and archaea strains: (a) growth rates OD600nm of bacteria on different salt concentration liquid MGM medium; (b) growth rates OD600nm of archaea on different salt concentration liquid MGM medium.

3.2. Identification by 16S rRNA gene analysis
The 16S rRNA gene analysis related their grouping in seven genera; *Virgibacillus, Idiomarina, Halobacillus*, for bacteria; *Halorubrum, Halobacterium, Halogeometricum*, and *Natrinema* for archaea. Bacteria Gram negative isolates of *Idiomarina* genus belonged to *Gammaproteobacteria* Class, *Idiomarinaceae* family and Gram positive of *Virgibacillus* and *Halobacillus* belonged to *Bacilli* Class, *Bacillaceae* family [16, 17]. Archaea genera are all belonged to *Halobacteria* Class, *Halorubrum*, *Halobacterium*, *Halogeometricum*, and *Natrinema* for archaea. With high salt tolerance and generally pigment production [16].

### 3.3. Enzymatic assay

The enzymatic assay was first performed for the 45 proteolytic selected strains with azocasein using PBS buffer [13, 10], after production on casein and gelatin media. Results presented on figures 2 and 3 allowed us to select 5 bacteria and 5 archaea that shown interesting enzymatic activity: 1, 4, 11, 13 and 17 for bacteria; B, F, H, L and N for archaea.

![Figure 2](image1.png)

Figure 2. Bacterial proteasic activity assay tested on azocasein at 366nm: (a) OD366nm for bacterial activities on casein; (b) OD366nm for bacterial activities on gelatin.

![Figure 3](image2.png)

Figure 3. Archaeal proteasic activity assay tested on azocasein at 366nm: (a) OD366nm for archaenal activities on casein; (b) OD366nm for archaenal activities on casein.

The enzymatic assay was confirmed for archaeal strains using a second method based on azocasein also but in Tris- HCl buffer and including NaCl 23% (w/v) as described upper [15]. This time, results presented on figure 4 confirm a good proteolytic activity for only four strains: L, M, N and U.

For further tests, we combine the two methods results and select only four strains: 4 and 11 for bacteria, H and N for archaea.
The enzymatic activity (figure 5) for selected proteolytic bacteria and archaea isolates, estimated with the two methods showed a very good activity, in comparison with the positive control, especially with bacteria strain 4 (Idiomarina sp.) that conserve its proteolytic properties with the two methods, which means that in absent of NaCl (0%, w/v) or presence (12.5%, w/v) the proteolysis is well done.

For archaea, the strain N (Natrinema sp.) showed a good proteolytic activity with the second method in presence of NaCl (23%, w/v), which is reported by other studies [18, 19] however, the activity is completely lost in the first one probably because of salt missing.

If we combine this results with those of salt physiology, demonstrated in figure 1, we can easily see that Idiomarina sp. conserve it’s proteolytic activity due to the large salinity growth interval (2.5 – 17.5%, w/v), at the same time, the archaea Natronema sp. require a high salt concentration, which explain the proteolytic activity lost in the first enzymatic assay.

4. Conclusion

The proteolytic activity production from archaeal and bacterial microorganisms isolated from Algerian sites was investigated in this study. 45 Halophilic isolates were able to produce protease. Extremely halophilic archaeal isolates were affiliated to Natrinema, Halorubrum, Halobacterium, halogeomatricum and bacterial isolates to Virgibacillus, Idiomarina, and Halobacillus genera, of which several strains could produce hydrolytic enzymes. This study that the genera Idiomarina and Natrinema showed the most important proteolytic activity (Figure 5).

According to the results, it is suggested that Natrinema and Idiomarina genera are excellent candidates for production the proteolytic enzymes. Proteolytic activity produced by Natrinema under high salinity condition could made this strain an interesting candidate for future investigation. Those extremely halophilic isolates were selected for further studies for their great biotechnological applications with respect to their capacity to produce different hydrolases. It would be more
constructive if these enzymes are purified from the isolates and then characterized, which is the next step in the current work.

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Conflicts of Interest: The authors declare no conflict of interest.

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