We are IntechOpen, the world's leading publisher of Open Access books Built by scientists, for scientists



185,000

200M



Our authors are among the

TOP 1% most cited scientists





WEB OF SCIENCE

Selection of our books indexed in the Book Citation Index in Web of Science™ Core Collection (BKCI)

Interested in publishing with us? Contact book.department@intechopen.com

Numbers displayed above are based on latest data collected. For more information visit www.intechopen.com



Chapter

Halocins, Bacteriocin-Like Antimicrobials Produced by the Archaeal Domain: Occurrence and Phylogenetic Diversity in *Halobacteriales*

Afef Najjari, Hiba Mejri, Marwa Jabbari, Haitham Sghaier, Ameur Cherif and Hadda-Imene Ouzari

Abstract

Members of extremely halophilic archaea, currently consisting of more than 56 genera and 216 species, are known to produce their specific bacteriocin-like peptides and proteins called halocins, synthesized by the ribosomal pathway. Halocins are diverse in size, consisting of proteins as large as 35 kDa and peptide "microhalocins" as small as 3.6 kDa. Today, about fifteen halocins have been described and only three genes, halC8, halS8 and halH4, coding C8, S8 and H4 halocins respectively have been identified. In this study, a total of 1858 of complete and nearly complete genome sequences of Halobacteria class members were retrieved from the IMG and Genbank databases and then screened for halocin encoding gene content, based on the BLASTP algorithm. A total of 61 amino acid sequences belonging to three halocins classes (C8, HalH4 and S8) were identified within 15 genera with the abundance of C8 class. Phylogenetic analysis based on amino acids sequences showed a clear segregation of the three halocins classes. Halocin S8 was phylogenetically more close to HalH4. No clear segregation on species and genera levels was observed based on halocin C8 analysiscontrary to HalH4 based analysis. Collectively, these results give an overview on halocins diversity within halophilic archaea which can open new research topics that will shed light on halocins as marker for haloarchaeal phylogentic delineation.

Keywords: archaea, bioinformatics, diversity, halocins, phylogeny

1. Introduction

Microorganisms of the third domain of life, Archaea, have been cultivated and described for more than 100 years [1], however, they have been first assigned to the Bacteria domain because of their great phenotypic similarities. In the late 1970s, Carl Woese and his collaborators, recognized the Archaea as the third domain of life on earth based on molecular phylogenetic analyses [2]. The dichotomous (eukary-otic/prokaryotic) classification was no longer valid, leading to a reclassification of

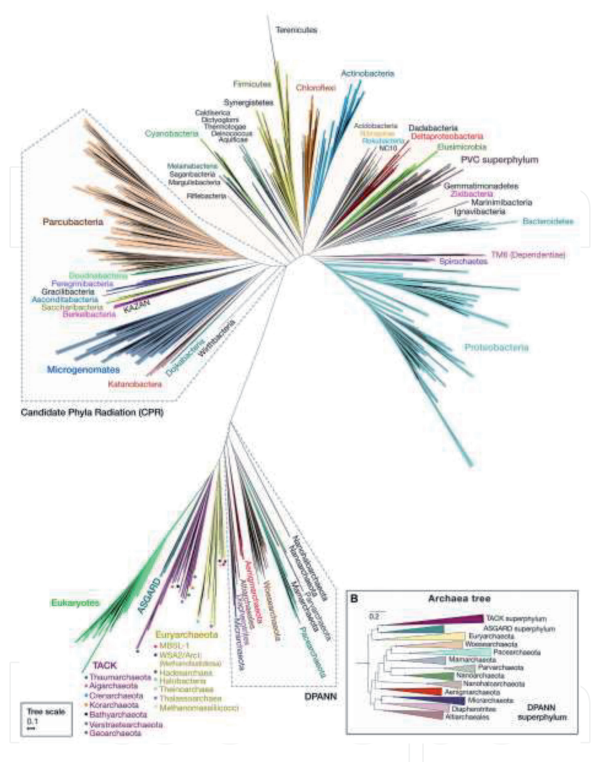
organisms as three separate domains: Eucarya (*Eukaryotes*), Archaea and Bacteria (bacteria) [3]. Archaea share several similarities with the other two domains of life. They are similar to size-level bacteria, organization of their chromosomes, absence of nucleus and organelles, presence of polycistronic transcription units and use of Shine-Dalgarno sequences for the initiation of the translation. In addition, it was shown that their metabolic proteins are essentially bacterial in nature following analysis of many complete genomes of Archaea [2]. Archaea also share similarities with the Eucarya domain, such as the proteins involved in key informational processes such as replication [4], transcription, translation [5, 6], DNA repair [7, 8], mRNA degradation and proteolysis. Translation in Archaea has eukaryotic initiation and elongation factors, and their transcription involves TATA binding protein and TFIIB [9].

The biotopes colonized by these microorganisms, are supposed to approach to the primitive terrestrial atmosphere (high salinity or pH, devoid of O₂, rich in H₂ and CO₂ constituting the raw materials for the production of methane) [10]. They present spectacular adaptations, especially in extreme environments. We distinguish: (i) Thermophilic Archaea: living at high temperatures (60–80°C) (ii) Hyperthermophilic Archaea: living at very high temperatures (up to 121°C); (iii) Psychrophilic Archaea: prefering low temperatures (below 15°C) [11]; (iv) Halophilic Archaea: colonizing very saline environments (3–5 M NaCl) such as the Dead Sea [12, 13]; (v) Acidophilic Archaea: living at low pH (as low as pH 1 and dying at pH 7) and Alkaliphilic Archaea: thriving at high pH (up to 9) [14].

2. Taxonomy of the archaeal domain

The first phylogenetic study based on the comparison of the 16S rDNA gene sequences coding for the small subunit, separated the first founding members of Archaea into two taxa, one grouping methanogenic species and those living under conditions of extreme salinity, the other containing species living at very high temperatures and at acidic pH [15]. Ten years later, analyses on a larger taxonomic group led to the division of the Archaea kingdom into two groups: (i) Crenarchaeota, which is composed exclusively of microorganisms living at very high temperatures, and (ii) *Euryarchaeota*, a heterogeneous group of species with different phenotypes (methanogenic species, species living at very high temperatures, moderate temperatures or at high salt concentrations) [3]. Fifteen years later and thanks to metagenomic analyses, two phyla, *Thaumarchaeota* and *Korarchaeota* were established based on the results of genomic comparison of two uncultivable strains, Candidatus cenarchaeum symbosium and Candidatus korarchaeum cryptofilum, with genomic traits belonging to both phyla *Crenarchaeota* and *Euryarchaeota* [16]. On the other hand, the symbiont Nanoarchaeum equitans, occupying cells of the host *Ignicoccus hospitalis*, showed even more genomic divergence with the other members of the *Crenarchaeota* and was therefore the first member of the phyla *Nanoarchaeota*.

Today, we count more of 15 phyla in the reign of Archaea, some of them having been grouped in superphylum. One distinguishes the superphylum TACK, proposed in 2011 and of which the eukaryotes would have evolved according to the theory of the eocyte, grouping *Thaumarchaeota*, *Aigarchaeota*, *Crenarchaeota* and *Korarchaeota* phyla [17]. This superphylum has been joined by recently proposed phyla: *Bathyarchaeota*, *Geoarchaeota* and *Lokiarchaeota* [18]. Another superphylum, DPANN, was proposed in 2013 and includes *Diapherotrites*, *Parvarchaeota*, *Aenigmarchaeota*, *Nanoarchaeota* and *Nanohaloarchaeota* phyla (**Figure 1**) [19]. Finally, the phyla *Woesearchaeota* and *Pacearchaeota*, described in 2016, were grouped in the DPANN superphylum (**Figure 1**).



Halocins, Bacteriocin-Like Antimicrobials Produced by the Archaeal Domain: Occurrence... DOI: http://dx.doi.org/10.5772/intechopen.94765

Figure 1. *Representation of the tree of life based on SSU rDNA gene sequences of the three domain of life* [23].

3. Antimicrobial potential of extremely halophilic archaea

Halophilic archaea were the first members of archaea found to produce bacteriocins-like proteins known as halocins. The first studies date from the beginning of 1980s with experiments demonstrating the presence of antagonistic interactions between halophilic archaeal strains isolated from the Alicante salt in Spain [20]. Today, about fifteen halocins have been described and only three genes, *halC8*, *halS8* and *halH4*, coding C8, S8 and H4 halocins, respectively, have been identified [21, 22]. Yet, no three-dimensional structural data of archaeocins are available in public databases.

3.1 Halocins

Halocins, bacteriocins-like peptides and proteins produced by extremely halophilic archaea, were first discovered in 1982 by F. Rodriguez Valera [24, 25]. They are classified according to their size into two major classes: high molecular mass (protein, > 10 kDa) and low molecular mass (peptide, \leq 10 kDa) called microhalocins [26, 27]. It has been shown that halocins are effective against *Haloarchaea* and *Crenarchaea* such as *Sulfolobus* spp. and *Methanosarcina thermopila*, and thus act across the main subdivision of the archaeal domain. These compounds represent a general class of antiarchaeal toxins and there is no confirmation about the inhibition of bacteria [26, 27]. Although several halocins were identified, only some of them have been characterized and purified.

3.1.1 Microhalocins

These halocins are composed of a peptide with size below or in the range of 10 kDa. Seven halocins have been characterized including HalS8, HalR1, HalC8, HalU1, HalH6, Sech7a and Sech10. They are hydrophobic and retain their activity in the absence of salt and can be stored at 4°C. They are relatively insensitive to heat and organic solvents [28].

3.1.1.1 Halocin S8 (HalS8)

HalS8 is the first characterized microhalocin with 36 amino acids (3580 Da), it is synthesized by the uncharacterized S8a haloarchaea [29]. Halocin S8 showed a narrow inhibitory spectrum and can only inhibit *Halobacterium salinarum* NRC817, *Halobacterium* GRB and *Haloferax gibbonsii* [29]. It can be desalted and it is heat resistant. Its activity is resistant to trypsin but sensitive to proteinase K and is undetectable in the transition to the stationary phase [29, 30]. The *halS8 gene* is *encoded* on a ~200-kbp megaplasmid [29].

3.1.1.2 Halocin HalR1 (HalR1)

Halocin R1, the second characterized microhalocin, is produced by *Halobacterium salinarum* GN101, a strain isolated from solar salt marsh in Mexico [31]. Initially HalR1 was described with a molecular weight of 6.2 kDa [32] and later on, it was shown that the HalR1 peptide is composed of 38 amino acids [24, 29]. Like HalS8, the activity is not affected by desalting and is resistant to acids, bases, organic solvents DNase and RNase, and against some proteases such as papain, trypsin or thermolysin, but it is sensitive to proteinase K, pronase P and elastase [20, 32].

3.1.1.3 Halocin C8 (HalC8)

Halocin C8 is produced by *Natrinema* sp. AS7092, a strain isolated from the large Chaidan Salt Lake, China (7.44 kDa, 76 amino acids) [28]. It is a unique polypeptide with an isoelectric point of 4.4 [33]. Its activity is retained after desalting, boiling and frozing [33]. Halocin C8 has a very broad spectrum of activity against several species and genera of *Halobacteriales* members including *Natronobacterium gregoryi*, *Nbt*. comb. nov and *Natronomonas pharaonis* [28]. The *halC8* gene encodes both halocin C8 and its immunity protein HalI.

3.1.1.4 Halocin A4 (HalU1)

Halocin A4, also called also halocin U1, is produced by an uncharacterized haloarchaea strain isolated from a Tunisian saltern [34]. Its molecular weight is 7.435 Da, as determined by the spectrometric mass, and is both acidic (pH = 4.14) and hydrophobic (eluent at ~85% acetonitrile) [26]. Halocin A4 has been reported to inhibit the growth of crenarchaeal *Sulfolobus* sp. strains [26]. Gene encoding HalA4 is *located* on a 300 kpb megaplasmid, pHM300 (NC_017943) [29].

3.1.1.5 Halocin H6 (HalH6)

Halocin H6 is produced by *Haloferax gibbonsii* Ma 2.39 species [27]. Its activity is resistant to trypsin. Stabilities of this peptide were studied and have shown that HalH6 can be desalted and it retained its activity after heat treatment up to 10 min at 100°C [27]. Halocin H6 is considered as a bactericidal substance which causes cell lysis and the specific target of HalH6 is the Na+/H+ antiport [27, 35].

3.1.1.6 Halocin Sech7a

Halocin Sech7a was excreted by the extremely halophilic haloarchaeon Sech7a, isolated from brine samples of Secovlje solar salterns crystallizers in Slovenia [36]. Sech7a is about 11 kDa. It is stable over a wide pH range and is heat labile at temperatures above 80°C. Its optimal activity was observed in the early exponential phase growth at 45°C. It loses activity under low salt conditions, but its activity can be restored after dialysis against initial saline conditions [36].

3.1.1.7 Halocin SH10

Halocin SH10 is produced by *Natrinema* sp. BTSH10, a strain isolated from the Kanyakumari salt marsh, Tamil Nadu, India [37]. The optimal production of halocin SH10 is at 42°C, pH 8.0 and 3 M NaCl at the stationary phase. In this context, it was reported that the activity is lost under acidic conditions [37]. Production of SH10 is influenced by the carbon source composition of the medium — *Natrinema* sp. BTSH10 could produce maximal halocin in the presence of beef [37].

3.1.2 Protein halocins

This class comprises halocins composed of proteins greater than 10 kDa in size. Currently, there are two characterized protein halocins, HalH1 and HalH4, in the range of 30 to 35 kDa [28].

3.1.2.1 Halocin H4

Halocin H4, produced by *Haloferax mediterranei* R4 (ATCC 33500) was isolated from a Spanish solar salt pond in Alicante. It is the first halocin that was studied [20]. The optimal activity was detected at the midpoint between exponential and stationary phases [20]. Halocin H4 is sensitive to proteases, high temperature and desalting. HalH4 has an antimicrobial activity against other haloarchaeons. It Interacts with the membrane of the target cells where it causes permeability changes that result in an ionic imbalance leading to cell lysis and death [21, 38]. The *halH4* gene, encoding halocin activity, is located on the pHM300 megaplasmid, a single polypeptide of 34.9 kDa.

3.1.2.2 Halocin H1

Halocin H1 is produced by *Haloferax mediterranei* M2a (previously known as *H. mediterranei* Xia3) isolated from salt ponds in Santa Pola (Alicante, Spain) [20]. Halocin H1 is a single 31 kDa polypeptide characterized by a broad inhibitory spectrum among *Halobacteriales* members. HalH1 activity is temperature and salt dependant. It is stable at 50°C only and requires a salt concentration of 1.5 M to maintain its activity [38, 39]. Optimum activity was observed at mid-exponential phase. The sensitivity to proteases and the gene encoding activity were not determined yet.

3.2 Applications of halocins

Some studies reported the role of halocins in a variety of environmental, industrial and biotechnological applications *** (REFERENCES?). However, this topic is poorly documented and somewhat controversial. One of these applications is the use of halocin producing strains in the textile industry during the tanning process characterized by high salinity concentration, halocins could inhibit the growth of pathogenic microbes affecting the quality of products. [7, 10]. Moreover, some halocins have also been reported for biomedical and therapeutic uses, for example, Halocin H7 has been shown to inhibit the Na+/H+ antiport in *Haloarchaea*, can be used as a treatment to reduce the injuries caused when ischemic organ transplantation is re-infused [35]. The therapeutic potential of halocins needs more research on their physical structures and their modes of action. On the other hand, halocins are known also to have a potential application in food industry as preservative agents by controlling the growth of haloarchaea in salted food products [40].

4. Materials and methods

Here, we evaluated the evolutionary relationship between bacteriocin- like-producing haloarchaea members based on comparisons of their amino acid sequences retrieved from annotated genomes sequences deposited in the IMG database [41].

4.1 Database search of halocin gene clusters

Schematic workflow of the methodology employed of amino acid sequences retrieving and phylogenetic assessment is illustrated in **Figure 2**. The methodology consisted of: first, complete and nearly complete genome sequences of *Halobacteriales* members were retrieved from IMG database. Then, in *silico* screening for gene sequences encoding halocins was done based on the BLASTP algorithm with default parameters [42]. All redundant and low-quality sequences were eliminated from datasets.

4.2 Phylogenetic reconstruction

Multiple sequences alignment of retrieved amino acid sequences were performed using ClustalW [43]. The evolutionary history was inferred using the Unweighted pair group method with arithmetic mean (UPGMA) method [44] implemented in MEGA X [45, 46]. The optimal tree with the sum of branch length = 18.99 is shown. Percentages of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) are shown next to the branches. The tree is drawn to scale, with branch lengths in the same units as those of the evolutionary distances used to infer the phylogenetic tree. The evolutionary distances were computed using the Poisson correction method [47] and are in the

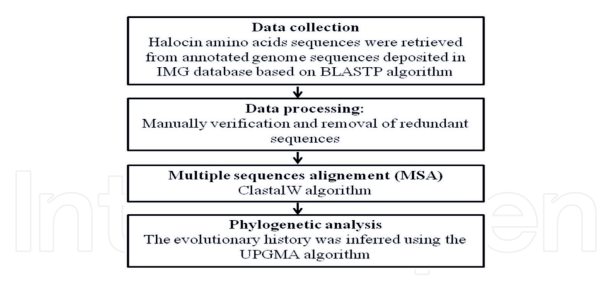


Figure 2.

Schematic workflow of the methodology employed for amino acids sequences retrieving and phylogenetic assessment.

units of the number of amino acid substitutions per site. All ambiguous positions were removed for each sequence pair (pairwise deletion option). In the final dataset, a total of 405 positions was obtained.

5. Results

5.1 Amino acid sequence of halocins

A total of 1858 of complete and nearly complete genome sequences of *Halobacteriales class* members were retrieved from the IMG database and creened for halocin encoding gene based on the BLASTP algorithm with default parameters [42].

A total of 61 amino acid sequences were retrieved from 15 genera belonging to Halobaceria class including Natrinema, Haloferax, Haloterrigena, Natronorubrum, Halobacterium, Haladaptatus, Halorubrum, Halococcus, Halopiger, Natrialba, Halolamina, Natronococcus, Haloarcula, Halapricum and Halorussus. Furthermore, some other unclassified halohilic archaea were present as well, including uncultured halophilic archaeon, halophilic archaeon sp. DL31 and Haloarchaeon S8 (**Table 1**).

Results showed that some species present more than one copy for halocin encoding genes. In fact, three (n = 3) classes of halocins were identified in this study (**Table 1**).

The first class is halocin C8-like bacteriocin domain (HalC8), the best known bacteriocin like sequences in archaea, it has been demonstrated to be produced from a ProC8 precursor, targeted to the membrane by the Tat pathway, and cleaved by an unknown mechanism to yield the active mature peptide HalC8 and an immunity protein HalI, protecting the producing strain against its own AMP [22]. HalC8 was identified in all species except *Natrialba aegyptia* DSM 13077. Indeed, *Natrinema* genus members appears to be more represented in terms of C8-like bacteriocin production. It's worth noting that among the six officially described species within the *Natrinema* genus, five are described in the current analysis. Several studies report the production of HalC8 and/or the presence of the *halC8* gene among *Natrinema* species isolated from different geographical origins like Chaidan Salt Lake in Qinghai province, China [33], Ichekaben salterns Chotts and sebkhas in algeria [48, 49].

The second class is halocin H4 (HalH4) identified in *Haloferax mediterranei* strain ATCC 33500, *Natrialba aegyptia and Natrinema gari* JCM 14663 species.

Extremophilic Microbes and Metabolites - Diversity, Bioprospecting and Biotechnological...

Domain: Archaea Natrinema Natrinema pellivubrum 157 Halocin C3-like Kingdom: ICM 10476 Halocin C3-like Bryarchaeota Natrinema aplikum DSM Jarrinema pallidum DSM Tortisema pallidum DSM JST3 Natrinema pallidum DSM Jonatinema pallidum DSM JST3 Natrinema pallidum DSM Jonatinema pallidum DSM JST3 Natrinema pallidum DSM JAP-DQR Natrinema pallidum DSM JST3 Natrinema altimense JCM JAP-DQR Natrinema altimense JCM JSS9 Natrinema aguri JCM 14663 Halocin C3-like Natrinema aguri JCM 14663 Halocin C4-like Halocin H4 Haloferax Haloferax mediterranei R-4 Halocin C4-like Haloferax kuestnewe DSM Halocin C4-like Halocin C4-like Haloferax kuest	Taxonomy	Genus	Species level	Class of Halocii
Phylamic Baryarchaeata Class: Halobacteria Cla	Kingdom:		-	
Class: Halobacteriales Smily: Halobacteriales Smily: Halobacteriales Haloferes Halobacteriales Haloferes H	-		Natrinema sp. J7-2	
Halobacteriaceae DSM 15624 Natrinema altunense Natrinema altunense 1A4 DCR Natrinema altunense JCM 12890 Natrinema altunense JCM Natrinema altunense JCM 12890 Natrinema altunense A1 Natrinema altunense 4.1 Natrinema agari JCM 14663 Halocin C8-like Jacobia Haloferax Haloferax Haloferax mediterranei Haloferax Haloferax mediterranei Haloferax Haloferax sep, ATCC BAA-646 Haloferax sep, ATCC BAA-645 Haloferax sep, ATCC BAA-646 Haloferax larsenii DS2 Haloferax larsenii DCM 13917 Haloferar larsenii CDM 5 Halocin C8-like Haloferrigena Halocin C8-like Haloferrigena Halocin C8-like Haloferax larsenii DCM 13917 Haloterrigena salifodimae Halocin C8-like KY19 Haloterrigena salindimae Haloterrigena salindimae Halocin C8-like KY19 Haloterrigena salindimae Haloterrigena salind JCM 13891 Haloterrigena salindimae Halocin C8-like </td <td>Class: Halobacteria</td> <td></td> <td>-</td>	Class: Halobacteria		-	
1A4-DGR Natrinema altanensa JCM 12890 Natrinema altanensa JCM 12890 Natrinema altanensa J2 Natrinema altanensa J2 Natrinema altanensa J1 Natrinema altanensa J2 Natrinema J2 Haloferax Haloferax Haloferax altanenii J2 Haloferax altanenii DS2 Haloferax altanenii DS2 Haloferax altanenii JCM J3917 Haloterrigena Haloterrigena altifodinae ZY19 Haloterrigena salifodinae ZY19 Haloterrigena salifodinae Haloterrigena salin JCM	Family:			
12890 Natrinema sp. J7-1 Natrinema altunense AJ2 Natrinema altunense AJ2 Natrinema altunense AJ2 Natrinema atunense 4.1 Natrinema atunense 4.1 Natrinema gari JCM 14663 Haloferax Haloferax mediterranei Haloferax Haloferax mediterranei Haloferax mediterranei Haloferax sp. ATCC BAA-646 Haloferax sp. ATCC BAA-646 Haloferax app. ATCC BAA-646 Haloferax sp. ATCC BAA-646 Haloferax app. ATCC BAA-645 Haloferay app. ATCC BAA-645 Haloferay app. ATCC BAA-645 Haloterrigena sulifodinae ZY19				
Natrinema altunense AJ2 Natrinema atunense 4.1 Natrinema gari JCM 14663 Halocin C8-like bacteriocin and Halocin H4 Haloferax Haloferax mediterranei A-1 Haloferax Haloferax mediterranei R-4 Haloferax mediterranei R-4 Halocin C8-like bacteriocin Haloferax lucentense DSM Halocin C8-like bacteriocin Haloferax sep. ATCC BAA-646 Haloferax alexandrinuss JCM 10717 Haloferax alexandrinuss JCM 10717 Haloferax alexandrinus JCM 10717 Haloferax larsenii JCM 13917 Halocin C8-like bacteriocin Haloterrigena Haloterrigena salifodinae ZV19 Haloterrigena jeotgali A29 Halocin C8-like bacteriocin Haloterrigena salina JCM 13891 Halocin C8-like bacteriocin Matronorubrum Natronorubrum tibetense DSM 13204 Halocin C8-like bacteriocin Natronorubrum Natronorubrum tibetense DSM 13204 Halocin C8-like bacteriocin				
Natrinema atunense 4.1 Natrinema gari JCM 14663 Halocin C8-like bacteriocin and Halocin 14 Haloferax Haloferax mediterranei Halocin 14 Haloferax Haloferax secondrinus Join 08-like Haloferax alexandrinus JCM 10717 Haloferax larsenii CDM 5 Haloferax larsenii CDM 5 Haloferax larsenii JCM 13917 Halocin 14 Haloterrigena Haloterrigena salifodinae Halocin 14 LZV19 Haloterrigena salifodinae Halocin 14 Haloterrigena solina JCM 13891 Halocin 08-like bacteriocin Haloterrigena solina JCM 13891 Haloterrigena solina JCM 13891 Haloterrigena solina JCM 13891 Haloterrigena NATOORUBUM Natronorubrum tibetense DSM 13204 Halocin 08-like bacteriocin Matronorubrum Natronorubrum tibetense DSM 13204 Halocin 03-like bacteriocin			Natrinema sp. J7-1	
Natrinema gari JCM 14663 Halocin C8-like bacteriocin and Halocin H4 Haloferax Haloferax mediterranei ATCC 33500 Halocin H4 Haloferax mediterranei R-4 Halocin C8-like bacteriocin Haloferax sp. ATCC BAA-646 Haloferax sp. ATCC BAA-645 Haloferax larsenii CDM 5 Haloferax larsenii CDM 5 Haloferax larsenii JCM 13917 Haloterrigena Halocin C8-like bacteriocin ZY19 Halocin C8-like bacteriocin Haloterrigena salifodinae Halocin C8-like bacteriocin J3917 Haloterrigena salifodinae Haloterrigena salifodinae Halocin C8-like bacteriocin Matomorubrum tibetense DSM 13204 Natronorubrum Natronorubrum tibetense GA33 Natronorubrum sediminis Halocin C8-like bacteriocin			Natrinema altunense AJ2	
bacteriocin and Halocin H4 Haloferax JCM 10717 Haloferax Haloferax JCM 10717 Haloferax Haloferax JCM 10717 Haloferax Haloferax JCM 10717 Haloferax Haloferax <t< td=""><td></td><td>Natrinema atunense 4.1</td></t<>			Natrinema atunense 4.1	
ATCC 33500 Haloferax mediterranei R-4 Haloferax lucentense DSM 14919 Haloferax sp. ATCC BAA-646 Haloferax volcanii DS2 Haloferax sp. ATCC BAA-646 Haloferax sp. ATCC BAA-645 Haloferax sp. ATCC BAA-645 Haloferax sp. ATCC BAA-645 Haloferax larsenii CDM 5 Haloferax larsenii JCM 13917 Haloterrigena Haloterrigena thermotolerans Haloterrigena salifodinae ZY19 Haloterrigena salifodinae Haloterrigena salifodinae ZY19 Haloterrigena salifodinae Haloterrigena salifodinae ZY19 Haloterrigena salina JCM 13891 Haloterrigena sp. P1A			Natrinema gari JCM 14663	bacteriocin and
Haloferax lucentense DSM 14919 Halocin C8-like bacteriocin Haloferax sp. ATCC BAA-646 Haloferax sp. ATCC BAA-646 Haloferax alexandrinus JCM 10717 Haloferax sp. ATCC BAA-645 Haloferax larsenii CDM 5 Haloferax larsenii CDM 5 Haloferrigena Haloterrigena Haloterrigena salifodinae ZY19 Haloterrigena anahii H13 Haloterrigena salifodinae ZY19 Haloterrigena sp. P1A Haloterrigena sp. P1A Haloterrigena turkmenica WANUI5 Halocin C8-like bacteriocin Natronorubrum Natronorubrum tibetense GA33 Halocin C8-like bacteriocin		Haloferax	-	Halocin H4
14919 bacteriocin Haloferax sp. ATCC BAA-646 Haloferax volcanii DS2 Haloferax volcanii DS2 Haloferax alexandrinus JCM 10717 Haloferax sp. ATCC BAA-645 Haloferax sp. ATCC BAA-645 Haloferax larsenii CDM 5 Haloferax larsenii JCM 13917 Haloterrigena Haloterrigena tursenii JCM 13917 Haloterrigena Haloterrigena salifodinae ZY19 Halocin C8-like ZY19 Haloterrigena salifodinae Haloterrigena salifodinae Halocin C8-like Bay1 Haloterrigena salina JCM Haloterrigena sp. P1A Halocin C8-like Haloterrigena turkmenica WANU15 Natronorubrum Natronorubrum tibetense GA33 Natronorubrum tubetense GA33 Natronorubrum sediminis			Haloferax mediterranei R-4	
BAA-646 Haloferax volcanii DS2 Haloferax volcanii DS2 Haloferax alexandrinus JCM 10717 Haloferax sp. ATCC BAA-645 Haloferax larsenii CDM 5 Haloferax larsenii JCM 13917 Haloterrigena Haloterrigena Haloterrigena salifodinae ZY19 Haloterrigena salifodinae ZY19 Haloterrigena salifodinae Katronorubrum Natronorubrum tibetense DSM 13204 Halocin C8-like bacteriocin Haloterrigena salifodinae Haloterrigena sp. P1A Haloterrigena sp. P1A Haloterrigena turkmenica WANU15 Natronorubrum Natronorubrum tibetense GA33 Natronorubrum sediminis			-	
Haloferax alexandrinus JCM 10717 Haloferax sp. ATCC BAA-645 Haloferax larsenii CDM 5 Haloferax larsenii JCM 13917 Haloterrigena Haloterrigena Haloterrigena jeotgali A29 Haloterrigena salifodinae ZY19 Haloterrigena salina JCM Haloterrigena salina JCM 13891 Haloterrigena sp. P1A Haloterrigena sp.			· ·	
JCM 10717 Haloferax sp. ATCC BAA-645 Haloferax larsenii CDM 5 Haloferax larsenii JCM 13917 Haloterrigena Haloterrigena Haloterrigena Haloterrigena Haloterrigena Haloterrigena Haloterrigena salifodinae ZY19 Haloterrigena pietgali A29 Haloterrigena salina JCM 13891 Haloterrigena salina JCM 13891 Haloterrigena salina JCM 13891 Haloterrigena turkmenica WANU15 Natronorubrum Natronorubrum tibetense GA33 Natronorubrum sediminis			Haloferax volcanii DS2	
BAA-645 Haloferax larsenii CDM 5 Haloferax larsenii JCM 13917 Haloterrigena Halocin H4 thermotolerans Halocin C8-like Date Haloterrigena salifodinae ZY19 Haloterrigena jeotgali A29 Haloterrigena solito JCM 13891 Haloterrigena solita JCM 13891 Haloterrigena sp. P1A Haloterrigena turkmenica WANU15 Natronorubrum Natronorubrum Natronorubrum tibetense GA33 Natronorubrum sediminis				
Haloferax larsenii JCM 13917Haloterrigena thermotoleransHalocin H4 thermotoleransHaloterrigenaHaloterrigena salifodinae ZY19Halocin C8-like bacteriocinHaloterrigena jeotgali A29 Haloterrigena mahii H13 Haloterrigena salina JCM 13891Haloterrigena salina JCM 13891Haloterrigena sp. P1A Haloterrigena turkmenica WANU15Halocin C8-like bacteriocinNatronorubrumNatronorubrum tibetense GA33 Natronorubrum sediminisHalocin C8-like bacteriocin				
13917 Haloterrigena Haloterrigena thermotolerans Halocin H4 Haloterrigena salifodinae Halocin C8-like ZY19 Halocin C8-like Haloterrigena jeotgali A29 Haloterrigena mahii H13 Haloterrigena salina JCM 13891 Haloterrigena sp. P1A Haloterrigena sp. P1A Haloterrigena turkmenica WANU15 Halocin C8-like Natronorubrum Natronorubrum tibetense Halocin C8-like GA33 Natronorubrum sediminis Halocin C8-like			Haloferax larsenii CDM 5	
thermotolerans Haloterrigena salifodinae Halocin C8-like ZY19 Haloterrigena salifodinae Halocin C8-like Haloterrigena jeotgali A29 Haloterrigena jeotgali A29 Haloterrigena mahii H13 Haloterrigena salina JCM 13891 Haloterrigena sp. P1A Haloterrigena turkmenica WANU15 Halocin C8-like Natronorubrum Natronorubrum tibetense Halocin C8-like DSM 13204 Halocin C8-like bacteriocin Natronorubrum tibetense GA33 Natronorubrum sediminis				
ZY19 bacteriocin Haloterrigena jeotgali A29 haloterrigena mahii H13 Haloterrigena salina JCM 13891 Haloterrigena sp. P1A haloterrigena sp. P1A Haloterrigena turkmenica WANU15 Natronorubrum Natronorubrum tibetense DSM 13204 Halocin C8-like bacteriocin Natronorubrum sediminis		Haloterrigena		Halocin H4
Haloterrigena mahii H13Haloterrigena salina JCM 13891Haloterrigena sp. P1AHaloterrigena sp. P1AHaloterrigena turkmenica WANU15NatronorubrumNatronorubrumNatronorubrum tibetense GA33Natronorubrum sediminis				
Haloterrigena salina JCM 13891 Haloterrigena sp. P1A Haloterrigena turkmenica WANU15 Natronorubrum Natronorubrum tibetense GA33 Natronorubrum sediminis			Haloterrigena jeotgali A29	
13891 Haloterrigena sp. P1A Haloterrigena turkmenica WANU15 Natronorubrum Natronorubrum tibetense DSM 13204 Natronorubrum tibetense GA33 Natronorubrum sediminis			Haloterrigena mahii H13	
Haloterrigena turkmenica WANU15NatronorubrumNatronorubrum tibetense DSM 13204Halocin C8-like bacteriocinNatronorubrum tibetense GA33Natronorubrum tibetense A33Halocin C8-like bacteriocin			e e	
WANU15 Natronorubrum Natronorubrum tibetense Halocin C8-like DSM 13204 bacteriocin Natronorubrum tibetense GA33 Natronorubrum sediminis Natronorubrum sediminis			Haloterrigena sp. P1A	
DSM 13204 bacteriocin Natronorubrum tibetense GA33 Natronorubrum sediminis				
GA33 Natronorubrum sediminis		Natronorubrum		

Halocins, Bacteriocin-Like Antimicrobials Produced by the Archaeal Domain: Occurrence... DOI: http://dx.doi.org/10.5772/intechopen.94765

Taxonomy	Genus	Species level	Class of Halocin
Domain: Archaea	Halobacterium	Halobacterium sp. DL1	Halocin C8-like
Kingdom: <i>Euryarchaeota</i> Phylum: <i>Euryarchaeota</i>		Halobacterium salinarum DSM 670	bacteriocin
Class: <i>Halobacteria</i> Order: <i>Halobacteriales</i>	_	Halobacterium salinarum DSM 671	
Family: <i>Halobacteriaceae</i>	-	Halobacterium salinarum DSM 6692	
		Halobacterium salinarum DSM 3754	
	391	Halobacterium salinarum DSM 668	
-	Haladaptatus	Haladaptatus paucihalophilus DX253	
	_	Haladaptatus sp. R4	
	_	Haladaptatus paucihalophilus DSM 18195	
	Halorubrum	Halorubrum lacusprofundi R1S1	
	_	Halorubrum trapanicum CBA1232	
-	Halococcus	Halococcus sp. 197A	
	_	Halococcus salifodinae DSM 8989	
-	Halopiger	Halopiger sp. IIH3	
_	Natrialba	Natrialba aegyptia DSM 13077	
-	Halolamina	Halolamina pelagica CGMCC	
-	Natronococcus	Natronococcus occultus SP4	
-	Haloarcula	Haloarcula salaria H5-DGR	
	Halapricum	Halapricum salinum CBA1105	
	Halorussus	Halorussus amylolyticus YC93	
	Halophilic archaeon	halophilic archaeon sp. DL31	
	uncultured halophilic archaeon	uncultured halophilic archaon J07HX5	
	_	uncultured haloarchaeon J07ABHX67	
	_	Uncultured Halobacteriaceae archaea SG1_71_5	

Table 1. Classes of Halocins identified by in silico analysis of all genomes of halophilic archaea domain available in IMG database.

Extremophilic Microbes and Metabolites - Diversity, Bioprospecting and Biotechnological...

HalH4 was first characterized from *H. mediterranei* isolated from solar saltern lakes of Spain [50]. HalH4 is a 40 kDa protein with an N-terminal 46 aa leader peptide which is cleaved off leaving a 313 aa mature halocin [51].

The third class is halocin S8, a microhalocin of 36 amino acids (3580 Da) initially purified from an unidentified haloarchaeal strain S8a, isolated from the Great Salt Lake (Utah, 109 United States) [52].

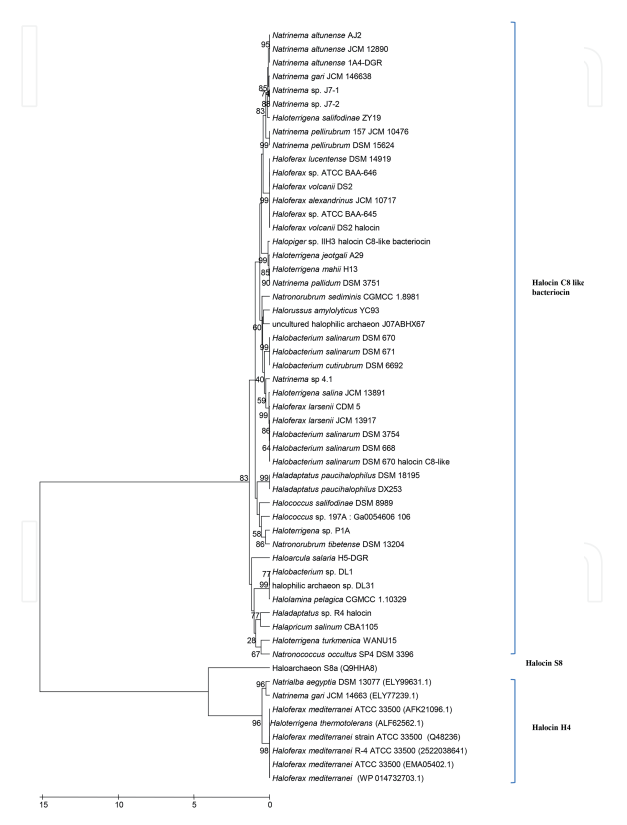


Figure 3.

Phylogenetic tree of halocin amino acid sequences of halophilic archaeal species. The evolutionary history was inferred using the UPGMA algorithm implemented in MEGA X software. Numbers at the nodes indicate the percentage of occurrence in 100 bootstrapped trees (bootstrap values > 50% are shown).

5.2 Phylogenetic analysis

Phylogenetic analysis of retrieved halocin peptide sequences was conducted and the result is illustrated in **Figure 3**. Results showed a clear segregation of the three halocins classes (C8, H4 and S8), where halocin S8 is phylogenetically more close to HalH4. Furthemore, no clear separation of species was observed based on HalC8 amino acids sequences analyses. HalC8 was detected in 12 genera belonging to three orders of *Halobacteria* class [53]: *Natrialbales* (*Natrinema*, *Haloterrigena*, *Natronorubrum*, *Halopiger*); *Haloferacales* (*Haloferax*, *Halolamina*) and *Halobacteriales* (*Halorussus*, *Halobacterium*, *Haladaptatus*, *Halococcus*, *Haloarcula*, *Halapricum*) and uncultured halophilic archaeon (J07ABHX67) phylogenetically related to species *Halorussus amylolyticus* YC93. The halocin S8 was detected only in the strain Haloarchaeon S8a (Q9HHA8). HalH4 is identified in *Natrialbales* (*Natrinema gari* JCM 14663 (ELY77239.1), *Haloterrigena thermotolerans*, *Natrialba aegyptia* DSM 13077) and *Haloferacales* (*Haloferax*).

It's worth noting that HalH4/HalC8 halocins were identified in *Haloferax*, *Haloterrigena* and *Natrinema* genera with only the species *Natrinema gari* JCM 14663 (ELY77239.1) being able to produce the two classes in the same time. Earlier studies reported that several described halocins, with broad inhibitory properties, are derived from *Haloferax* and *Natrinema* strains [24, 28, 45, 51, 54] and it has been suggested that halocin production may explain their dominance in some saline ecosystems [54, 55].

6. Conclusion

On the basis of our *in silico* analyses, we can conclude that halocin production is considered as a general feature of some members of halophilic archaea, particularly members of *Natrialbales* and *Haloferacales* orders with the occurrence of Halocin C8-like production. This group can thrive in saline ecosystems in which several other microorganisms are not able to live. Thus, the dominance of certain species isolated in some saline ecosystems could be attributed to halocin production as a mechanism of competition between microrganisms. This chapter will open new research lines that will shed light on halocins as marker for haloarchaeal phylogentic delineation.

Conflict of interest

We have no conflict of interest to declare.

Intechopen

Author details

Afef Najjari^{1*}, Hiba Mejri¹, Marwa Jabbari^{2,3}, Haitham Sghaier^{2,4}, Ameur Cherif⁴ and Hadda-Imene Ouzari¹

1 Faculté des Sciences de Tunis, Université de Tunis El Manar, LR03ES03 Microorganismes et Biomolécules Actives, 2092, Tunis, Tunisia

2 Laboratory "Energy and Matter for Development of Nuclear Sciences" (LR16CNSTN02), National Center for Nuclear Sciences and Technology (CNSTN), Sidi Thabet Technopark, 2020, Tunisia

3 Biochemistry and Molecular Biology, Code UR13ES34 Research Unit, Faculty of Science of Bizerte, University of Carthage, 7021 Zarzouna, Tunisia

4 ISBST, BVBGR-LR11ES31, Biotechpole of Sidi Thabet, Univ. Manouba, 2020 Ariana, Tunisia

*Address all correspondence to: najjariafef@gmail.com; afef.najjari@fst.utm.tn

IntechOpen

© 2020 The Author(s). Licensee IntechOpen. This chapter is distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

References

[1] Cavicchioli R. *archaea* — timeline of the third domain. (2011). Nature Reviews Microbiology. 9(1):51-61.

[2] Woese CR, Fox GE. (1977). The concept of cellular evolution. J Mol Evol. 10(1):1-6.

[3] Woese CR, Kandler O, Wheelis ML (1990). Towards a natural system of organisms: proposal for the domains Archaea, Bacteria, and Eucarya. Proc Natl Acad Sci U S A. 87(12):4576-9.

[4] Barry ER, Bell SD. DNA Replication in the Archaea (2006). Microbiol Mol Biol Rev. 70(4):876-87.

[5] Londei P (2005). Evolution of translational initiation: new insights from the archaea. FEMS Microbiol Rev. 29(2):185-200.

[6] Steitz TA (2008). A structural understanding of the dynamic ribosome machine. Nature Reviews Molecular Cell Biology. 9(3):242-53.

[7] Fujikane R, Ishino S, Ishino Y, Forterre P (2010). Genetic analysis of DNA repair in the hyperthermophilic archaeon, Thermococcus kodakaraensis. Genes Genet Syst. 85(4):243-57.

[8] Kelman Z, White MF (2005). Archaeal DNA replication and repair. Curr Opin Microbiol. 8(6):669-76.

[9] Soppa J (2006). From genomes to function: haloarchaea as model organisms. Microbiology. 152(Pt 3):585-90.

[10] Karr J (2006). Seven foundations of biological monitoring and assessment. Biologia Ambientale. 20:7-18.

[11] Van de Vossenberg JL, Driessen AJ, Konings WN (1998). The essence of being extremophilic: the role of the unique archaeal membrane lipids. Extremophiles. 2(3):163-70. [12] Rothschild LJ, Mancinelli RL(2001). Life in extreme environments.Nature. 409(6823):1092-101.

[13] Margesin R, Schinner F (2001). Potential of halotolerant and halophilic microorganisms for biotechnology. Extremophiles.5(2):73-83.

[14] Quatrini R, Johnson DB (2016).Acidophiles: Life in Extremely AcidicEnvironments. Caister Academic Press;2016. 300 p.

[15] Fox GE, Stackebrandt E,Hespell RB, Gibson J, Maniloff J,Dyer TA, et al (1980). The phylogeny ofprokaryotes. Science. 209(4455):457-63.

[16] Brochier-Armanet C, Boussau B, Gribaldo S, Forterre P (2008).
Mesophilic *Crenarchaeota*: proposal for a third archaeal phylum, the *Thaumarchaeota*. Nat Rev Microbiol. 6(3):245-52.

[17] Guy L, Ettema TJG (2011). The archaeal 'TACK' superphylum and the origin of eukaryotes. Trends in Microbiology. 19(12):580-7.

[18] Castelle CJ, Wrighton KC,
Thomas BC, Hug LA, Brown CT,
Wilkins MJ, et al. (2015). Genomic
Expansion of Domain Archaea
Highlights Roles for Organisms from
New Phyla in Anaerobic Carbon
Cycling. Current Biology. 25(6):690-701.

[19] MacLeod G, Bozek DA, Rajakulendran N, Monteiro V, Ahmadi M, Steinhart Z, et al (2019). Genome-Wide CRISPR-Cas9 Screens Expose Genetic Vulnerabilities and Mechanisms of Temozolomide Sensitivity in Glioblastoma Stem Cells. Cell Rep. 27(3):971-986.

[20] Rodriguez-Valera F, Juez G, Kushner D (2011). Halocins: salt-dependent bacteriocins produced by extremely halophilic rods. Canadian Journal of Microbiology. 28:151-4.

[21] Cheung J, Danna KJ, O'Connor EM, Price LB, Shand RF (1997). Isolation, sequence, and expression of the gene encoding halocin H4, a bacteriocin from the halophilic archaeon Haloferax mediterranei R4. J Bacteriol. 179(2):548-51.

[22] Sun C, Li Y, Mei S, Lu Q, Zhou L, Xiang H (2005). A single gene directs both production and immunity of halocin C8 in a haloarchaeal strain AS7092. Mol Microbiol. 57(2):537-49.

[23] Castelle CJ, Banfield JF. Major New Microbial Groups Expand Diversity and Alter our Understanding of the Tree of Life (2018). Cell. 172(6):1181-97.

[24] O'Connor E, Shand R (2002). Halocins and sulfolobicins: The emerging story of archaeal protein and peptide antibiotics. Journal of industrial microbiology & biotechnology. 28:23-31.

[25] Li Y, Xiang H, Tan H (2002). Halocin: protein antibiotics produced by extremely halophilic archaea.

[26] Haseltine C, Hill T, Montalvo-Rodriguez R, Kemper SK, Shand RF, Blum P (2001). Secreted euryarchaeal microhalocins kill hyperthermophilic crenarchaea. J Bacteriol. 183(1):287-91.

[27] Torreblanca M, Meseguer I, Rodriguez-Valera F (1989). Halocin H6, a Bacteriocin from Haloferax gibbonsii. Microbiology-sgm. 135:2655-61.

[28] Shand R, Leyva K (2008). Archaeal Antimicrobials: An Undiscovered Country. Archaea: New Models for Prokaryotic Biology. Caister Academic Press.

[29] Price LB, Shand RF (2000). Halocin S8: a 36-Amino-Acid Microhalocin from

the Haloarchaeal Strain S8a. J Bacteriol. 182(17):4951-8.

[30] Shand R, Price LB, O'Connor E (1999). Halocins: Protein antibiotics from hypersaline environments. In: Oren A (Ed), Microbiology and Biogeochemistry of Hypersaline Environments.CRC Press, Boca Raton, FL. 295-306.

[31] Ebert K, Goebel W, Rdest U, Surek B (1986). Genes and genome structures in the archaebacteria. Syst Appl Microbiol. 7, 30-35. 1986.

[32] Rdest U, Sturm M (1987). Bacteriocins from halobacteria. In: Burgess R, editor. Protein purification: micro to macro. New York, N.Y: Alan R. Liss, Inc.; pp. 271-278.

[33] Li Y, Xiang H, Liu J, Zhou M, Tan H (2003). Purification and biological characterization of halocin C8, a novel peptide antibiotic from Halobacterium strain AS7092. Extremophiles. 7(5):401-7.

[34] Shand R (2006). 29 Detection, Quantification and Purification of Halocins: Peptide Antibiotics from Haloarchaeal Extremophiles. Methods in Microbiology. 35:703-18.

[35] Meseguer I, Torreblanca M, Konishi T (1995). Specific inhibition of the halobacterial Na+/H+ antiporter by halocin H6. J Biol Chem. 270(12):6450-5.

[36] Pasić L, Velikonja BH, Ulrih NP (2008). Optimization of the culture conditions for the production of a bacteriocin from halophilic archaeon Sech7a. Prep Biochem Biotechnol. 38(3):229-45.

[37] Karthikeyan P, Bhat SG, Chandrasekaran M (2013). Halocin SH10 production by an extreme haloarchaeon Natrinema sp. BTSH10

isolated from salt pans of South India. Saudi Journal of Biological Sciences. 20(2):205-12.

[38] Perez AM (2000). Growth Physiology of Haloferax Mediterranei R4 and Purification of Halocin H4. Northern Arizona University; 156 p.

[39] Meseguer I, Rodriguez-Valera F (1986). Effect of Halocin H4 on Cells of Halobacterium halobium. Microbiology-sgm. 132:3061-8.

[40] Charlesworth J, Burns BP (2016).Extremophilic adaptations and biotechnological applications in diverse environments. AIMS Microbiology.2(3):251.

[41] Markowitz EM, Goldberg LR, Ashton MC, Lee K (2012). Profiling the "Pro-Environmental Individual": A Personality Perspective. Journal of Personality. 80(1):81-111.

[42] Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ (1990). Basic local alignment search tool. Journal of Molecular Biology. 215(3):403-10.

[43] Thompson JD, Higgins DG, Gibson TJ (1994). CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. Nucleic Acids Res. 22(22):4673-80.

[44] Sokal RR, Michener CD (1958). A statistical method for evaluating systematic relationships. 28, 1409-1438.

[45] Kumar S, Stecher G, Li M, Knyaz C, Tamura K (2018). MEGA X: Molecular Evolutionary Genetics Analysis across Computing Platforms. Mol Biol Evol. 35(6):1547-9.

[46] Sneath PHA, Sokal RR(1973). Numerical Taxonomy: The Principles and Practice of Numerical Classification. San Francisco: W.H.Freeman & Co Ltd; 1973. 588 p.

[47] Zuckerkandl E, Pauling L (1965). Molecules as documents of evolutionary history. J Theor Biol. 8(2):357-66.

[48] Imadalou-Idres N, Carré-Mlouka A, Vandervennet M, Yahiaoui H, Jean P, Rebuffat S (2013). Diversity and Antimicrobial Activity of Cultivable Halophilic Archaea from Three Algerian Sites. 7:1057-69.

[49] Quadri I, Hassani I, L'haridon S, Chalopin M, Hacene H, Jebbar M (2016). Characterization and antimicrobial potential of extremely halophilic archaea isolated from hypersaline environments of the Algerian Sahara. Microbiological Research. 186.

[50] Quesada E, Ventosa A, Rodriguez-Valera F, Ramos-Cormenzana A (1982). Types and properties of some bacteria isolated from hypersaline soils. Journal of Applied Bacteriology. 53(2):155-61.

[51] Besse A, Jean P, Rebuffat S, Carré-Mlouka A (2015). Antimicrobial peptides and proteins in the face of extremes: Lessons from archaeocins. Biochimie. 118.

[52] Rinke C, Schwientek P, Sczyrba A, Ivanova NN, Anderson IJ, Cheng J-F, et al (2013). Insights into the phylogeny and coding potential of microbial dark matter. Nature.499(7459):431-7.

[53] Gupta RS, Naushad S, Baker S (2015). Phylogenomic analyses and molecular signatures for the class Halobacteria and its two major clades: a proposal for division of the class *Halobacteria* into an emended order *Halobacteriales* and two new orders, *Haloferacales* ord. nov. and Natrialbales ord. nov., containing the novel families *Haloferacaceae* fam. nov. and Natrialbaceae fam. nov. Int J Syst Evol Microbiol. 65(Pt 3):1050-69. Extremophilic Microbes and Metabolites - Diversity, Bioprospecting and Biotechnological...

[54] Atanasova N, Pietilä M, Oksanen H (2013). Diverse antimicrobial interactions of halophilic archaea and bacteria extend over geographical distances and cross the domain barrier. MicrobiologyOpen. 2.

[55] Kis-Papo T, Oren A (2000). Halocins: are they involved in the competition between halobacteria in saltern ponds? Extremophiles. 4(1):35-41.

