

## Isolation and identification of halophilic bacteria from Urmia Lake in Iran

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### Abstract

Halophiles are in all three domains of life: Archaea, Bacteria and Eucarya. Halophilic microorganisms in addition to form major part of life biodiversity can have many biotechnological applications. The objective of this research is isolation and identification of halophilic bacteria from Urmia Lake in Iran and the study of its bacterial biodiversity. After sampling of brines from Urmia Lake from 10 stations and depth of approximately 30-50 cm, in April 2011 and transfer to the laboratory in the sterile conditions, samples were enriched and cultured on defined media, and incubated. After appearance of colonies, selected strains were studied based on morphology, physiology and biochemical characteristics. For phylogenetic identification, their genomic DNA were extracted and amplified by PCR technique. Therefore their sequences were determined by genetic experiment based on 16S rRNA gene sequence and their similarity were analysed in GenBank of EzTaxon database. Finally the phylogenetic tree was constructed. Studied strains belonged to three genera: *Halomonas* 50% (including *H. andesensis* LC6(T) [12.5%], *H. gomseomensis* M12(T) [12.5%], *H. hydrothermalis* Slthf2(T) [12.5%], *H. boliviensis* LC1(T) [6.25%] and *H. janggokensis* M24(T) [6.25%]), *Salinivibrio* 25% (including *S.costicolasubsp. alcaliphilus* DSM 16359(T) [18.75%] and *S. sharmensis* BAG(T) [6.25%]) and *Idiomarina* 25% (including *I. loihiensis* L2TR(T) [25%]).

**Keywords:** Halophilic bacteria, Species diversity, Isolation, Phylogenetic, DNA analysis, Urmia Lake

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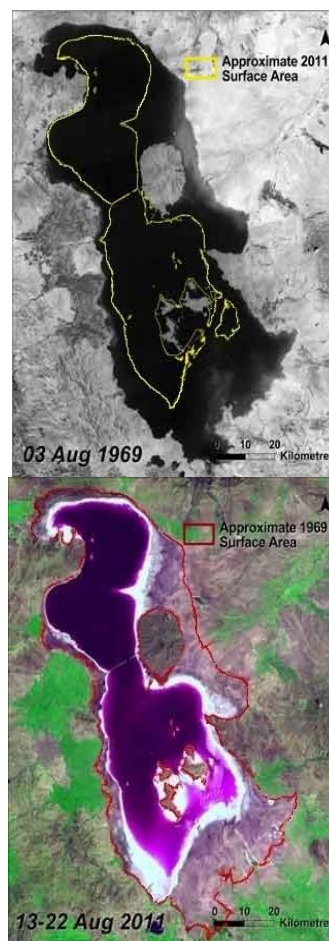
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## Introduction

Microorganisms requiring extreme environments for growth are called extremophiles (MacElroy, 1974). Extremophilic microorganisms can thrive in extreme environments such as unusual levels of salt, pH, pressure, temperature, *etc.*, and those which are adapted to live in hypersaline habitats are considered halophiles (Kumar and Gummadi, 2009). Halophiles are microorganisms that adapt to moderate and high salt concentrations. They are found in all three domains of life: Archaea, Bacteria and Eukarya. Halophilic bacteria grow over an extended range of salt concentrations (3–15% NaCl, w/v and above), unlike the truly halophilic archaea whose growth is restricted to high saline environments (Litchfield, 2002). Halophiles represent valuable sources of various biomolecules which can offer potential applications for biocatalysis and biotransformation (Schiraldi and Rosa, 2002). Hypersaline environments are those with salt concentrations above that of sea water (3.3% total dissolved salts) (Oren, 2002a). Urmia Lake (located in the northwest of Iran) can be considered as one of the largest permanent hypersaline lakes in the world and resembles the Great Salt Lake in the western USA in many respects of morphology, chemistry and sediments (Kelts and Shahrabi, 1986). The predominance of the Na<sup>+</sup> and Cl<sup>-</sup> ions illustrates the thalassohaline character of Urmia Lake (Sorgeloos, 1997). Therefore, Urmia Lake is an oligotrophic lake of thalassohaline origin (AzariTakami, 1993). Unfortunately, this lake has faced with drought problem in recent years. Fig. 1 shows the drought process of Urmia Lake

during 1969-2011, based on satellite images (Pengra, 2012).



**Figure 1: The drought process of Urmia Lake during 1969-2011**

The aim of this research is the study of halophilic bacteria from Urmia Lake. These strains can form a small part of Urmia Lake biodiversity and be the base of wide applications in biotechnology too.

## Materials and methods

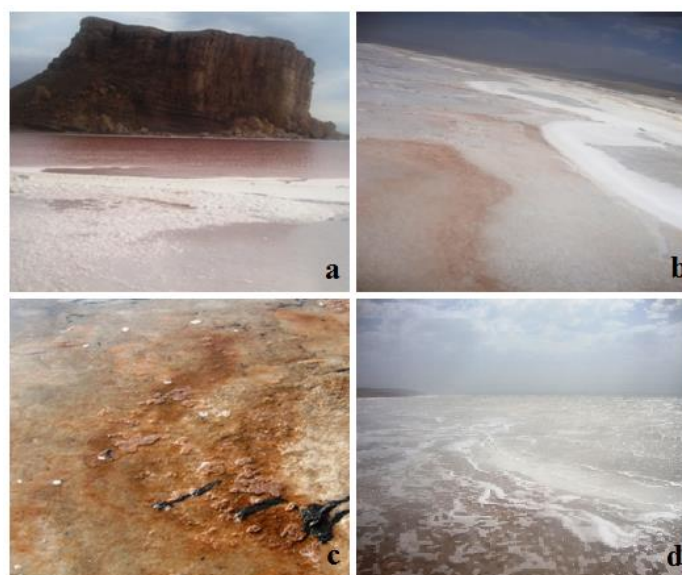
### Sampling

Urmia Lake in the northwestern corner of Iran is one of the largest permanent hypersaline lakes in the world. From among the regions with sampling ability, 10 stations were selected.

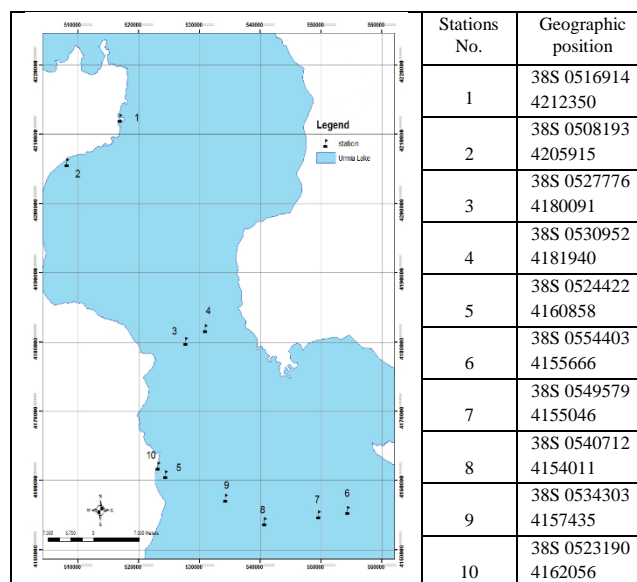
Water samples were collected in April 2011 from 10 stations (from depth of

approximately 30-50 cm) and transferred to the laboratory under sterile conditions.

Figs. 2 and 3 depict the images of sampling stations and their locations on the map.



**Figure 2:** The images of sampling stations, Urmia Lake, 2011  
**a:** station No. 1; **b:** station No. 2; **c:** station No. 5;  
**d:** station No. 3



**Figure 3:** The locations of sampling stations on the map

#### *Growth conditions and isolation*

Samples were added to Nutrient Broth (NB; MERCK) and Alkaline Peptone Water (APW; MERCK) media. Samples

were enriched by the methods of (Halako *et al.*, 2005; Amoozgar *et al.*, 2008) and incorporation of tow methods.

Enrichment in the Nutrient Broth was carried out in two forms:

1- First 10ml of water samples were centrifuged at 3000 rpm for 5 minute. After removing the top part of the sample, 1ml bottom sample of tube was added to 30ml Nutrient Broth (pH 7.2-7.4, supplemented with 5% (w/v) salt of lake).  
2- Water samples without centrifuge were added to 30ml Nutrient Broth (pH 7.2-7.4, supplemented with 5% (w/v) salt of lake). Then samples were incubated at 35°C on an orbital shaking incubator at 150 rpm for 48-72 hours.

Enrichment in the Alkaline Peptone Water was carried out following on:

10ml of water samples were centrifuged at 3000 rpm for 5 minute. After removing the top part of the sample, 1ml bottom sample of tube was added to 9ml of Alkaline Peptone Water (pH 8.6±0.2, supplemented with 5% and 10% (w/v) salt of lake) and incubated at 37°C for 24-48 hours. Therefore samples were cultured on Nutrient Agar (NA; MERCK) (Amoozegar *et al.*, 2008) and MacConkey Agar w/0.15% Bile Salts, CV and NaCl (Mac A; HIMEDIA) media (This medium was selected for the isolation of Gram-negative strains), and incubated at 35-37 °C for 24-48 hours.

All of the media were contained with 5% and 10% (w/v) salt (the salt of lake). Repeated cultures were carried out to achieve pure cultures.

#### *Phenotypic characteristics and phylogenetic analysis*

Phenotypic characteristics including morphological, physiological and biochemical tests were determined for each strain. Colonial morphology was described by using standard

microbiological criteria, such as pigmentation, form, colonial elevation and opacity. Cell morphology was examined by light microscopy. Gram staining was performed (Murray *et al.*, 1994) and the result was confirmed by the KOH test (Baron and Finegold, 1990). Motility was analysed by the wet-mount method (Murray *et al.*, 1994). Other phenotypic and biochemical characteristics were checked by using standard procedures and as recommended by Smibert and Krieg (1994).

Because of genotypic and phylogenetic studies, 16 isolates were selected for genetic experiment based on 16S rRNA gene sequence. First genomic DNA of the selected strains was extracted by IBRC genomic DNA extraction kit on following the manufacturer's recommended procedure and amplified by PCR technique. The 16S rRNA gene of the selected strains was amplified using the universal primers 27F (5'-AGAGTTTGATCCTGGCTCAG-3') and 1492R (5'-GGTTACCTTGTTACGACTT-3').

The amplification program was done by initial denaturation at 94°C for 3min followed by 30 cycles of denaturation at 94 °C for 1min, annealing at 57-58°C for 1min, extension 72°C for 1.5min, last extension 72°C for 10min and hold time 25°C for 10sec. Then amplified PCR products were sent to MacroGen Co. (South Korea) for 16S rRNA gene sequencing. Data obtained from sequencing were edited by bioinformatic software of Chromas Pro. and saved in FASTA format. At last sequences similarity of these strains were analysed on comparison with registered strains in

GenBank of EzTaxon database (EzTaxon server 2.1) (Chun *et al.*, 2007). After identification of closest strains to studied strains, data were aligned by Clustal\_X software (Larkin *et al.*, 2007).

Finally their phylogenetic tree was constructed by MEGA5 software and the neighbor-joining method (Saitou and Nei, 1987; Tamura *et al.*, 2011).

## Results

The average of temperature at sampling stations, pH and total salinity of samples were 13.7 °C, 7.02 and 32.4%.

Colonies were round with entire edges, convex, shiny to opaque or translucent. Isolated strains produced creamy colonies on Nutrient Agar and pink colonies on MacConkey Agar. All cells were Gram-negative, motile and rod-shaped which curved rods cells were seen between of them. The results of morphologic, physiologic and biochemical tests were listed in Table 1. Also Table 2 shows the comparison of 16S rRNA gene sequences in ExTaxon.

**Table 1: Morphologic, physiologic and biochemical characteristics.**

Tests		URM 1	URM 3	URM 5	URM 7
Isolation media	Centrifuge of sample	-	-	-	Centrifuge
	Enrichment medium	NB	NB	NB	NB
	Culture medium	NA	NA	NA	NA
	Salinity (w/v)	5%	5%	5%	5%
Colony pigmentation	Cream	Cream	Cream-Yellowish	Cream	
Colony morphology	Round, Convex, Entire edges, Translucent	Round, Convex, Entire edges, Translucent	Round, Convex, Entire edges, Bright	Round, Convex, Entire edges, Opaque	
Cell morphology	Short rod	Short rod	Short curved rod	Short curved rod	
Gram staining	Gram-negative	Gram-negative	Gram-negative	Gram-negative	
KOH	+	+	+	+	
Motility	+	+	+	+	
Oxidase	+	+	+	+	
Catalase	+	+	+	+	
O / F	+ / +	+ / +	+ / -	+ / -	
S / I / M	- / - / +	- / - / +	- / - / +	- / - / +	
Urease	+	+	+	+	
Nitrate reduction	W	+	-	W	
Simmons' Citrate	-	-	W	-	
TSI	A/A	K/A	K/K	K/K	
Methyl-red	-	-	+	-	
Voges-Proskauer	-	-	-	-	
Gelatinase activity	-	-	+	+	
Hydrolysis of Aesculin	-	-	-	-	
Lysine decarboxylase	-	-	ND	-	

<b>Table 1 continued:</b>					
Ornithine decarboxylase		-	-	ND	-
Argininedihydrolase		+	-	ND	-
<b>Tests</b>		<b>URM 10</b>	<b>URM 11</b>	<b>URM 12</b>	<b>URM 14</b>
Isolation media	Centrifuge of sample	Centrifuge NB	Centrifuge APW	Centrifuge APW	Centrifuge APW
	Enrichment medium	Mac A 5%	NA 5%	NA 5%	Mac A 5%
	Culture medium Salinity (w/v)				
	Colony pigmentation	Pink	Dark Cream	Cream	Pink
	Colony morphology	Round, Convex, Entire edges, Opaque	Round, Convex, Entire edges, Shiny	Round, Convex, Entire edges, Shiny	Round, Convex, Entire edges, Shiny
	Cell morphology	Short rod	Short rod	Short curved rod	Short curved rod
	Gram staining	Gram-negative	Gram-negative	Gram-negative	Gram-negative
	KOH	+	+	+	+
	Motility	+	+	+	+
	Oxidase	+	+	+	+
	Catalase	+	+	+	+
	O / F	+ / -	+ / -	+ / -	+ / +
	S / I / M	- / - / +	W / - / +	- / - / +	W / - / +
	Urease	W	W	W	W
	Nitrate reduction	-	-	-	+
	Simmons' Citrate	+	-	W	-
	TSI	A/A, H <sub>2</sub> S	K/K	K/K	K/A, H <sub>2</sub> S
	Methyl-red	-	+	+	-
	Voges-Proskauer	-	-	-	+
	Gelatinase activity	-	+	+	+
Hydrolysis of Aesculin	-	-	-	-	
Lysine decarboxylase	-	+	ND	-	
Ornithine decarboxylase	+	-	ND	-	
Argininedihydrolase	-	-	ND	-	
<b>Tests</b>		<b>URM 16</b>	<b>URM 20</b>	<b>URM A</b>	<b>URM B</b>
Isolation media	Centrifuge of sample	Centrifuge	Centrifuge	Centrifuge	Centrifuge
	Enrichment medium	APW	APW	APW	APW
	Culture medium	NA	Mac A	NA	Mac A
	Culture medium Salinity (w/v)	10%	10%	10%	10%
	Colony pigmentation	Dark cream	Pink	Cream	Dark pink
	Colony morphology	Round, Convex, Entire edges, Opaque	Round, Convex, Entire edges, Opaque	Round, Convex, Entire edges, Shiny	Round, Convex, Entire edges, Translucent
	Cell morphology	Rod	Rod	Short curved rod	Short curved rod

**Table 1 continued:**

	Gram staining	Gram-negative	Gram-negative	Gram-negative	Gram-negative
	KOH	+	+	+	+
	Motility	+	+	+	+
	Oxidase	-	-	+	+
	Catalase	+	+	+	+
	O / F	+ / -	+ / -	+ / -	+ / -
	S / I / M	W / - / +	- / - / +	+ / - / +	- / - / +
	Urease	-	-	W	-
	Nitrate reduction	-	-	-	+
	Simmons' Citrate	+	+	W	-
	TSI	K/A	A/A	K/K	K/A
	Methyl-red	+	+	+	-
	Voges-Proskauer	-	-	-	-
	Gelatinase activity	-	-	+	+
	Hydrolysis of Aesculin	+	+	-	+
	Lysine decarboxylase	-	-	+	-
	Ornithine decarboxylase	-	-	-	-
	Argininedihydrolase	+	-	-	-
	<b>Tests</b>	<b>URM C</b>	<b>URM E</b>	<b>URM F</b>	<b>URM I</b>
Isolation media	Centrifuge of sample	-	Centrifuge	-	Centrifuge
		NB	APW	NB	NB
	Enrichment medium	Mac A	Mac A	NA	NA
		5%	10%	5%	5%
	Culture medium				
	Salinity (w/v)				
	Colony pigmentation	Pink	Light pink	Light cream	Cream
	Colony morphology	Round, Convex, Entire edges, Opaque	Round, Convex, Entire edges, Opaque	Round, Convex, Entire edges, Opaque	Round, Convex, Entire edges, Opaque
	Cell morphology	Rod	Rod	Rod	Short curved rod
		Gram-negative	Gram-negative	Gram-negative	Gram-negative
	KOH	+	+	+	+
	Motility	+	+	+	+
	Oxidase	+	-	+	+
	Catalase	+	+	+	+
	O / F	+ / -	+ / -	+ / -	+ / -
	S / I / M	- / - / +	- / - / +	- / - / +	- / - / +
	Urease	W	W	+	W
	Nitrate reduction	+	W	+	+
	Simmons' Citrate	+	-	+	-
	TSI	A/A, H <sub>2</sub> S	K/A	K/A	A/A
	Methyl-red	+	+	-	+
	Voges-Proskauer	-	-	-	-
	Gelatinase activity	-	-	-	+
	Hydrolysis of Aesculin	-	-	-	+
	Lysine decarboxylase	-	-	-	-

**Table 1 continued:**

Ornithine decarboxylase	-	-	-	-
Aargininedihydrolase	-	-	-	-

NB: Nutrient Broth; NA: Nutrient Agar; APW: Alkaline Peptone Water; Mac A: MacConkey Agar; W: Weak

**Table 2: The comparison of 16S rRNA gene sequences in ExTaxon.**

Strain	Closest strain in GenBank (EzTaxon)	Accession	Similarity
URM 1	<i>H.hydrothermalis</i> Slthf2(T)	AF212218	97.7
	<i>H.alkaliphila</i> 18bAG (T)	AJ640133	97.6
	<i>H.venusta</i> DSM 4743(T)	AJ306894	97.5
URM 3	<i>H. hydrothermalis</i> Slthf2(T)	AF212218	98.3
	<i>H.alkaliphila</i> 18bAG(T)	AJ640133	98.2
	<i>H.venusta</i> DSM 4743(T)	AJ306894	98.1
URM 5	<i>I.loihiensis</i> L2TR(T)	AE017340	98.9
	<i>I.abysalis</i> KMM 227(T)	AF052740	98.5
	<i>I.ramblicola</i> R22(T)	AY526862	98.5
URM 7	<i>S.costicolasubsp. alcaliphilus</i> DSM 16359(T)	AJ640132	95.8
	<i>S.sharmensis</i> BAG(T)	AM279734	95.6
	<i>S.costicola subsp. costicola</i> NCIMB 701(T)	X95527	95.3
URM 10	<i>H.andesensis</i> LC6(T)	EF622233	97.4
	<i>H.venusta</i> DSM 4743(T)	AJ306894	97.3
	<i>H.alkaliphila</i> 18bAG(T)	AJ640133	97.1
URM 11	<i>I.loihiensis</i> L2TR(T)	AE017340	99.5
	<i>I.abysalis</i> KMM 227(T)	AF052740	99.1
	<i>I.ramblicola</i> R22(T)	AY526862	99
URM 12	<i>I.loihiensis</i> L2TR(T)	AE017340	99.6
	<i>I.ramblicola</i> R22(T)	AY526862	99.3
	<i>I.abysalis</i> KMM 227(T)	AF052740	99.2
URM 14	<i>S.sharmensis</i> BAG(T)	AM279734	92.1
	<i>S.costicola subsp. alcaliphilus</i> DSM 16359(T)	AJ640132	92
	<i>S.costicola subsp. costicola</i> NCIMB 701(T)	X95527	91.4
URM 16	<i>H.gomseomensis</i> M12(T)	AM229314	99.4
	<i>H.arcis</i> AJ282(T)	EF144147	97.6
	<i>H.subterranea</i> ZG16(T)	EF144148	97.5
URM 20	<i>H.gomseomensis</i> M12(T)	AM229314	99.5
	<i>H.arcis</i> AJ282(T)	EF144147	98
	<i>H.subterranea</i> ZG16(T)	EF144148	97.4
URM A	<i>I.loihiensis</i> L2TR(T)	AE017340	99.4
	<i>I.abysalis</i> KMM 227(T)	AF052740	99
	<i>I.ramblicola</i> R22(T)	AY526862	98.9
URM B	<i>S.costicolasubsp. alcaliphilus</i> DSM 16359(T)	AJ640132	99.3
	<i>S.costicola subsp. costicola</i> NCIMB 701(T)	X95527	98.6
	<i>S.costicola subsp. vallismortis</i> DSM 8285(T)	AF057016	97.4
URM C	<i>H.boliviensis</i> LC1(T)	AY245449	98
	<i>H.neptunia</i> Eplume1(T)	AF212202	97.9
	<i>H.variabilis</i> DSM 3051(T)	AJ306893	97.5
URM E	<i>H.janggokensis</i> M24(T)	AM229315	99.9
	<i>H.subterranea</i> ZG16(T)	EF144148	99.7
	<i>H.arcis</i> AJ282(T)	EF144147	98.5
URM F	<i>H.andesensis</i> LC6(T)	EF622233	94.3
	<i>H. venusta</i> DSM 4743(T)	AJ306894	94.2
	<i>H. hydrothermalis</i> Slthf2(T)	AF212218	94.2
URM I	<i>S.costicolasubsp.alcaliphilus</i> DSM 16359(T)	AJ640132	99.7
	<i>S.costicola subsp. costicola</i> NCIMB 701(T)	X95527	98.9
	<i>S.sharmensis</i> BAG(T)	AM279734	97.8



From total strains, 50%, 25% and 25% of strains belonged to members of the genera

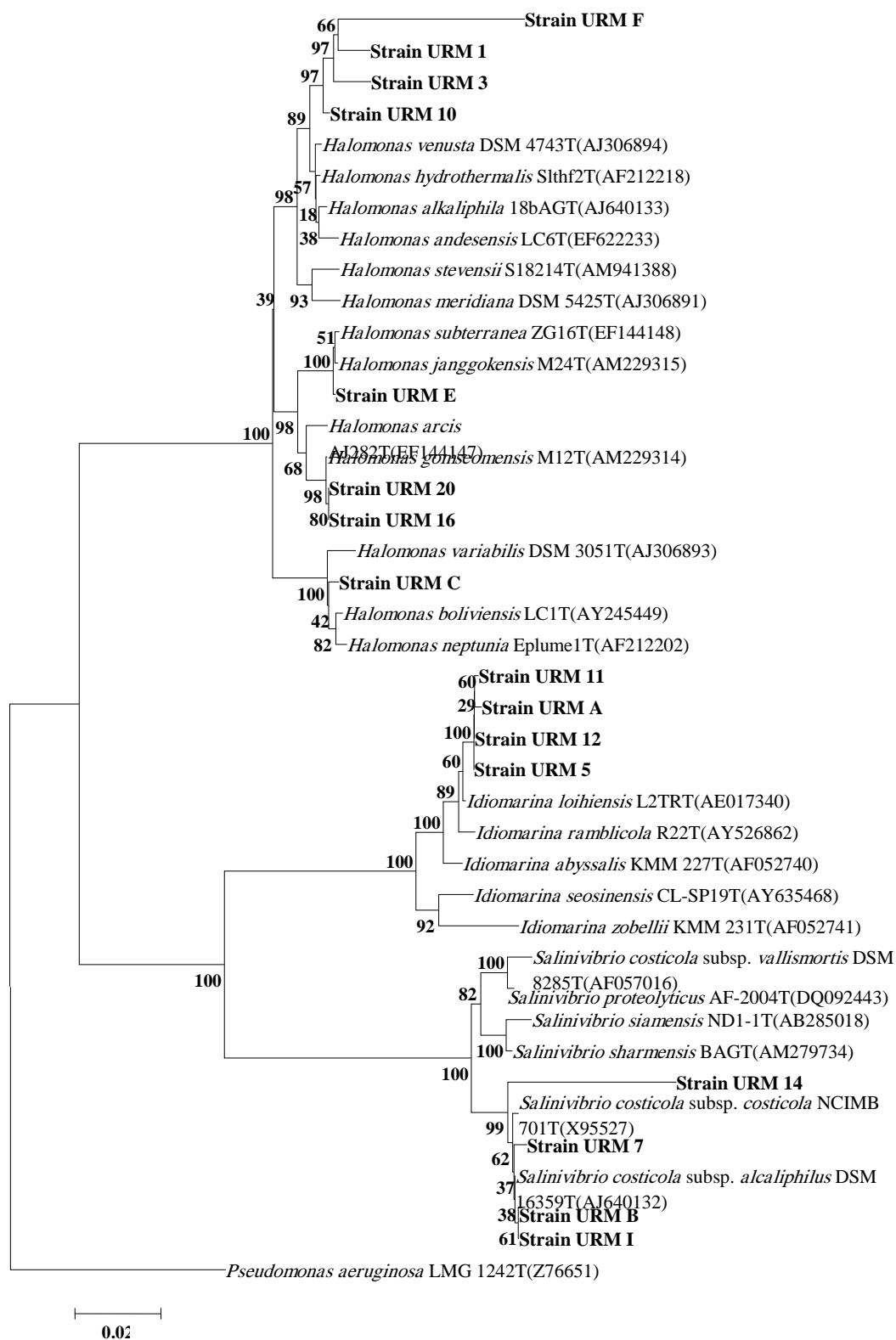
*Halomonas*, *Salinivibrio* and *Idiomarina*, respectively.

The genus *Halomonas* [50%] contained *H.andesensis* LC6(T) [12.5%], *H.gomseomensis* M12(T) [12.5%], *H.hydrothermalis* Slthf2(T) [12.5%], *H.boliviensis* LC1(T) [6.25%] and *H.janggokensis* M24(T) [6.25%]). The genus *Salinivibrio* [25%] included *S.costicola* subsp. *alcaliphilus* DSM [18.75%] and *S.sharmensis* BAG(T) [6.25%]. The identified genus *Idiomarina*

[25%] only belonged to *I.loihiensis* L2TR(T).

The study of 16S rRNA gene similarities showed more than 99% similarity for 50% of strains, 98.7–99% similarity for 6.25% of strains and less than 98.7% similarity for 43.75% of strains.

The analysis of 16S rRNA gene similarity along with phenotypic characteristics suggested that some of these strains can be representatives of new species in Urmia Lake. Furthermore, phylogenetic relationships of the studied strains are shown by phylogenetic tree in Fig.4.



**Figure 4: Neighbour-joining tree based on 16S rRNA gene sequences (Bootstraps were done using 100 replications and *Pseudomonas aeruginosa* LMG 1242T(Z76651) was used as an outgroup).**

## Discussion

Increased attention has been given in the last few years to moderately halophilic bacteria. Several studies have been conducted on their ecology, taxonomy and phylogeny as well as their biotechnological applications (Ramos-Cormenzana, 1991; Ventosa *et al.*, 1998a, b; Sanchez-Porro *et al.*, 2003).

In this study, the moderately halophilic Gram-negative bacteria members of the genera *Halomonas*, *Salinivibrio* and *Idiomarina* were isolated from Urmia Lake and their phenotypic and genotypic characteristics were studied. Most environmental isolates able to produce hydrolytic enzymes belonged to the Gram-negative genera *Salinivibrio* and *Halomonas*, two genera widely distributed in hypersaline environments (Ventosa, 1988; Ventosa, *et al.* 1998b). *Halomonas* is the largest genus in the family *Halomonadaceae*, with more than 50 recognized species until 2010 (Guzman *et al.*, 2010). This genus can tolerate or require a high salt concentration for growth (Ventosa *et al.*, 1998b). In addition, in our research, most of the reported strains belonged to the genus *Halomonas*. We isolated members of *Halomonas* from water samples of Urmia Lake but *Halomonas* have been isolated from water, soil, and seafood and depth-water samples from saline environments in the world (Oren, 2002b). Also, second abundance within the identified genera in the present research belonged to the strains of the genus *Salinivibrio*. Members of this genus are moderately halophilic bacteria distributed in salted meats, brines and

several hypersaline environments (Rao *et al.*, 1998).

In similar studies, *salinivibrio* was not isolated from Urmia Lake. Such as the results from ZununiVahed *et al.* (2011), strains of the genera *Halomonas* and *Idiomarina* were isolated but there was not any strain of *salinivibrio*. Maybe the present study is the first report of *salinivibrio* from Urmia Lake. Moreover, we isolated *Idiomarina loihiensis* sp. from surface water of Urmia Lake (depth of 30-50 cm), while the first isolation and identification of this species was from sediment at a depth of 11000 m in the Mariana Trench (Donachie *et al.*, 2003).

The said strains were only part of the microbial communities in Urmia Lake. The aim of this investigation was isolation and identification of halophilic bacteria from Urmia Lake that it led to identification of new strains from this lake. However, the presentation of new or unusual isolates, require a polyphasic approach. There were polyextremophile microorganisms within isolated strains too. The extremophiles that could tolerate more than one factor of harsh conditions are called polyextremophiles. They can withstand a variety of stresses (Chela-Flores, 2013).

*S.costicola* subsp. *alcaliphilus* and *S.sharmensis* are haloalkaliphilic bacteria (Romano *et al.*, 2005, 2011).

*H.boliviensis* is a halophilic, psychrophilic, alkalitolerant bacterium and *H.hydrothermalis* is a psychrotolerant halophile too (Kaye *et al.*, 2004; Quillaguaman *et al.*, 2004). Besides, the genus *Idiomarina* has been proposed as psychrotolerant heterotrophic halophilic

as well (Ivanova *et al.*, 2000). The 16S rRNA gene sequence similarity value has played an important role in delineating novel taxa and in the identification of isolates. Stackebrandt and Goebel (1994) suggested that a 16S rRNA gene sequence similarity of 97% should become the boundary for delineation of prokaryotic species, which has been well accepted among microbiologists.

More recently, Stackebrandt and Ebers (2006) proposed a more relaxed cut-off value of 98.7–99%, after inspection of a large amount of recently published data. Even though this new proposal requires further validation and discussion, it is evident that high quality of sequencing should be the prerequisite to the use of lower similarity cut-off values for the identification of prokaryotes.

Thus, the studied strains can be novel members of halophilic bacteria by supplemental studies such as DNA-DNA hybridization, G+C content and more chemotaxonomic and phenotypic studies. In addition, this article shows that Urmia Lake and its bacterial stores can have wide biotechnology applications.

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