

NUMERICAL STUDY ON SOME ACTINOMYCETES ISOLATED FROM BURULLOS LAKE IN EGYPT

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ABSTRACT

Twenty nine actinomycetes isolates were isolated from Burullos Lake and characterized taxonomically for 62 phenotypic traits including morphological; biochemical, nutritional, substrate utilization and anti-microbial activities. The results were analyzed by numerical techniques using the simple matching coefficient (S_sM) and UPGMA clustering. At 54% similarity level, the majority of the isolates were grouped into six phenon (A, B, C, D, E and F). Only two isolates were grouped separately and formed two single clusters at the same level of similarity. A representative isolate from each phenon was identified. The isolates were found to be *Streptoveriticillum morookaense*, *Nocardia brasiliensis*, *Streptomyces alanosinicus*, *Streptomyces globosus* and *Streptomyces gancidicus*

Keywords: Burullos Lake, Actinobacteria, Numerical taxonomy, Fresh water habitats

INTRODUCTION

Class Actinobacteria (high G+C content, gram positive bacteria) has been proposed by **Stackebrandt et al (1997)** and includes members with unparalleled ability to produce diverse secondary metabolites (**Mellouli et al 2003**). They represent an ubiquitous group of microbes widely distributed in natural ecosystem including soils (**Xu et al 1998**), marine sediments (**Ghanem et al 2000, Sabry et al 2004**) and fresh water (**Hahn et al 2003**). Actinomycetes are not adapted for

growth in aquatic habitats but can nevertheless be recovered readily from fresh water, sea water and sediments samples (**McCarthy and Williams, 1990**).

Several studies demonstrated by cultivation independent methods revealed that the freshwater actinobacteria are present in a wide spectrum of ecologically different and globally distributed freshwater habitats (**Glöckner et al 2000; Lindström and Leskinen, 2002; Hahn et al 2003**). **Zwart et al (2002)** recently identified 34 putative phylogenetic clusters of bacteria which seem to contain

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typical freshwater inhabitants, 5 of the 34 clusters were affiliated with the class Actinobacteria.

Distribution of Actinobacteria in fresh water has been reported by **Glöckner *et al* (2000)** who performed in situ hybridization. On the other hand, **Pernthaler *et al* (2001)** successfully enriched a phylo-type of one freshwater cluster of actinobacteria in a continuous culture system. Thus, despite the high number of cells of actinobacteria observed in fresh water samples no representatives of the freshwater actinobacterial lineages have been isolated so far (**Zwart *et al* 2002**). **Zwart *et al* (2003)** found that actinobacterial cluster ACK-M1 is being well represented in most fresh water habitats. They detected this cluster in all of the 81 lakes screened (from Belgium, The Netherlands, Denmark, Sweden and Norway), using reverse line blot hybridization. In the most recent published data (**Van der Gucht *et al* 2005**) the ACK-M1 (Actinobacteria) was the most representative in each of the studied lakes. These data confirm those previously reported (**Crump *et al* 1999**; **Urbach, *et al* 2001**).

Up to our Knowledge, few studies dealt with the distribution of class Actinobacteria in aquatic habitats in Egypt (**Al-Diwany and Cross 1978**; **Ghanem *et al* 2000**). Lake Burullous is shallow brackish water that serves as fishery resources and reservoirs for drainage water. It is contaminated with anthropogenic materials (**Abu-Elela *et al* 2004**). Therefore, this study is an attempt to evaluate the distribution of members of class Actinobacteria in Lake Burullous using phenotypic, physiological and biochemical data, depending on numerical taxonomic procedures.

MATERIAL AND METHODS

Actinomycetes isolates

Twenty nine actinomycetes isolates were selected from a total of 130 isolates previously isolated from sediments and water samples of Lake Burullous (**Abu-Elela *et al* 2004**). They were maintained on a basal medium prepared with 1 liter of Lake water of the following composition (g/l): soluble starch, 10.0; KNO₃, 2.0; K₂HPO₄, 2.0; MgSO₄ .7H₂O, 0.05; CaCO₂, 0.02; FeSO₄, 0.01 and agar, 20.0 at 28 °C (**Küster and Williams, 1964**). Culture stocks were maintained at -20 °C in starch nitrate broth with 20% glycerol (**Kieser *et al* 2000**).

Characterization and numerical analysis

Numerical analysis was not used for a direct taxonomical purpose but also to facilitate data handling and strains grouping. A total of 62 characters were coded as negative (0) or positive (1). The simple matching coefficient (SsM) (**Sokal and Michener, 1958**) and the Jaccard coefficient (Sj) (**Sneath, 1957**) were used and clustering was achieved by unweighted pair group average linkage (UPGMA), **Sneath and Sokal, (1973)** and **Sneath, (1979)**. The computations were performed by using SYSTAT-PC program V 7 (**Wilkinson *et al* 1992**) on an IBM computer.

The selected isolates were phenotypically characterized as described by **Pridham and Gottlieb (1948)**; **Shirling and Gottlieb, (1966)** and **Tresner *et al* (1968)**. All tests were carried out at 28°C for up to 3 weeks.

For morphological characterization, the following media were used: Krasslinikov SRL agar, Glycerol- nitrate agar, Czapek- Dox agar after slight modification, Glycerol- asparagine agar and Inorganic salts- starch agar (**Shirling and Gottlieb, 1966**).

Additional phenotypic characterization was performed using the standard procedures, catalase and urease production were detected, melanin production according to **Shirling and Gottlieb, (1966)**, nitrate reduction (**Williams et al 1983**), sulphide precipitation (**Cowan, 1974**) and indole production (**Molin and Trenstrom, 1986**). Lecithinase was performed on egg-yolk medium according to the method of **Nitsch and Kutzner, (1969)**, Lipase (**Elwan et al 1977**), protease (**Ammar et al 1991**), pectinase, amylase (**Ammar et al 1998**) production was investigated. Also Degradation activities of some substance such as starch, cellulose, chitin, gelatin and tyrosine were performed according to **Nonomura and Ohara (1969)**.

The antagonistic activity of the tested isolates against *Escherichia coli* HP101, *Staphylococcus aureus* ATTC 29523 and *Bacillus subtilis* was determined as described by **Goodfellow et al (1990)**. Zones of inhibition were scored as positive results after 24 h at 28C°. Resistance against phenol (0.002, 0.01 w/v), crystal violet (0.0001, 0.001 w/v), and sodium azide (0.01w/v) was tested. Antibiotic resistance was examined as the ability to grow on medium supplemented with antibiotics one at a time using gentamycin (998 µg.ml⁻¹), erythromycin (804 µg.ml⁻¹), tetracycline (990 µg.ml⁻¹) and cefixine (997 µg.ml⁻¹) according to **Williams et al (1983)**.

Chemotaxonomic study was performed by the detection of diaminopimelic acid (DAP) isomers and diagnostic sugars according to **Becker et al (1964)** and **Lechevalier et al (1977)** in AL-Azhar University Fermentation Biotechnology and Applied Microbiology Center. Identification was carried out in comparison to reference strains.

Scanning electron microscopy

Electron micrographs were made with Jeol model ISM 5300 scanning electron microscope (SEM) operating at 15 KeV (**Molitoris et al 1996**).

RESULTS AND DISCUSSION

In a previous survey on the distribution of *Actinobacteria* in Lake Burullos, 130 isolates were obtained (**Abu-Elela et al 2004**). Based on colony morphology, 29 isolates were selected which represent different morphotypes. In this study, we aimed to cluster those isolates for the purpose of identification and classification.

Actinomycetes isolates were studied for 62 characters listed in Table (1). Analysis of the 62 characters using the simple matching coefficient (S_SM) and UPGMA clustering yielded the dendrogram in Fig. (1). Data showed that the majority of the isolates were grouped at 54% similarity level, into six phenon (A, B, C, D, E and F). Only two isolates were grouped separately and formed two single clusters at this level.

Phenon A: The four isolates grouped in phenon A possess several features that are common to members of genus *Streptovorticillum*. It is thus suggested that the

Table 1. Comparison of the frequencies of positive and negative characters for the six clusters of actinomycetes obtained by numerical taxonomy analysis

Character	Phenon	A	B	C	D	E	F
	No. of isolates	4	3	6	5	3	6
Growth on							
Krasslinikov SRL agar		100	100	100	0	33	83
Glycerol-asparagine agar		100	100	100	100	100	100
Glycerol –nitrate agar		100	100	100	100	100	100
Czapek-Dox agar		100	100	100	100	100	100
Inorganic salts –starch agar		75	100	100	100	100	100
Substrate mycelium							
Yellow brown		50	67	33	80	0	33
Reddish brown		25	0	50	0	67	50
Blue		25	0	0	0	0	0
Violet		0	33	17	20	33	17
Aerial mycelium							
Gray yellowish pink		0	33	0	0	67	0
Pink gray		0	67	17	0	0	0
Gray		0	0	67	80	0	50
Whitish gray		50	0	0	0	0	0
Blue		25	0	0	0	0	0
Violet		0	0	17	20	0	17
Yellow		25	0	0	0	33	33
Diffusible pigment							
Unpigmented		0	100	100	80	67	50
Reddish brown		0	0	0	20	33	0
Yellowish brown		100	0	0	0	0	50
Growth at °C							
20		50	100	100	100	100	100
30		100	100	100	100	100	100
40		100	100	100	100	100	100
50		25	0	0	0	0	17
Growth at pH							
5		50	100	50	20	0	50
6		75	100	100	100	100	100
7		100	100	100	100	100	100
8		75	100	100	100	100	100
9		75	67	100	100	100	100

Table 1. continued

Character	Phenon	A	B	C	D	E	F
	No. of isolates	4	3	6	5	3	6
Utilization of							
D-Glucose		75	100	83	80	100	100
D-Fructose		75	67	100	40	33	100
L-Arabinose		0	0	50	0	67	50
Lactose		75	0	100	100	100	50
Raffinose		50	0	67	20	0	50
Rhamnose		50	33	67	40	67	83
Mannitol		25	33	67	80	67	67
Galactose		25	33	50	60	33	50
Xylose		25	33	67	80	67	100
Growth in presence of NaCl %							
0		100	100	100	100	100	100
4		100	100	100	100	100	100
7		75	33	50	20	33	50
10		0	0	0	0	0	0
Degradation of							
Starch 1%		100	100	100	100	100	100
Cellulose 1%		50	67	100	0	0	100
Chitin 0.25%		0	0	33	0	0	0
Gelatin 0.4%		0	0	0	0	0	67
Tyrosine 0.1%		100	67	100	100	67	100
Enzyme activities							
Catalase		25	67	33	100	100	50
Urease		50	67	0	40	0	50
Nitrate reduction		50	100	17	40	33	67
Sulfide precipitation		25	0	0	0	33	50
Melanin production		0	0	0	100	33	0
Antibiosis against							
<i>E.coli</i> HP101		0	33	17	0	33	50
<i>S.aureus</i> ATCC 29523		25	0	100	20	33	83
<i>B.subtilis</i>		0	0	17	0	67	100

Table 1. continued

Character	Phenon	A	B	C	D	E	F
	No. of isolates	4	3	6	5	3	6
Resistance to							
Phenol (0.002 w/v)		100	100	100	100	100	100
Phenol (0.01w/v)		0	0	0	0	0	0
Crystal violet (0.0001 w/v)		50	100	100	80	100	100
Crystal violet (0.001 w/v)		25	67	33	40	100	100
Gentamycin at (998 µg.ml ⁻¹)		0	67	0	0	0	17
Erythromycin (804 µg.ml ⁻¹)		0	0	0	0	0	17
Tetracycline (990 µg.ml ⁻¹)		0	0	0	0	0	0
Cefixine (997 µg.ml ⁻¹)		0	0	0	0	0	0

isolates are different species of this genus (Table 1). SEM observation of the representative strain (Fig. 2A) showed phenotypic features clearly related to the reference strain *Streptovercillium morookaense* (Hensyl, 1994) the characters shared with both strains are shown in (Table 2) aerial mycelia are gray while substrate mycelia are yellow brown, the aerial mycelia consisted of long, straight filaments of conidia showing side branches at regular intervals and whitish to gray spore masses. The cell wall hydrolysates contained LL-DAP with no diagnostic sugars. Spores are not motile. Absence of melanin pigments but gray yellowish brown diffusible pigments was produced. Several carbon sources were utilized (Table 2). Positive results were recorded with urease, nitrate reductase and capable degrading starch, tyrosine, xanthine and esculin .Our isolate differs only from the reference strain in the fact

that it could not utilize raffinose and in this respect, it was suggested that the isolate is closely related to *Streptovercillium morookaense* as recommended by the international Keys (Williams *et al* 1989 and Hensyl, 1994).

Phenon B: This phenon contained three isolates grouped at 55% similarity level. They had morphological and biochemical characteristics similar to members of non-cardioform actinomycetes (Lechevalier, 1989).

Selected isolate from phenon B (Fig.2 B) was similar to the reference strain *No-cardia brasiliensis* in having pink gray aerial mycelium and the aerial mycelium moderately fragmented into a chain of conidia. The cell wall hydrolysate contained meso-DAP with arabinose and galactose as diagnostic sugars. The experimental strain shared some properties with *N. brasiliensis* in being non motile,

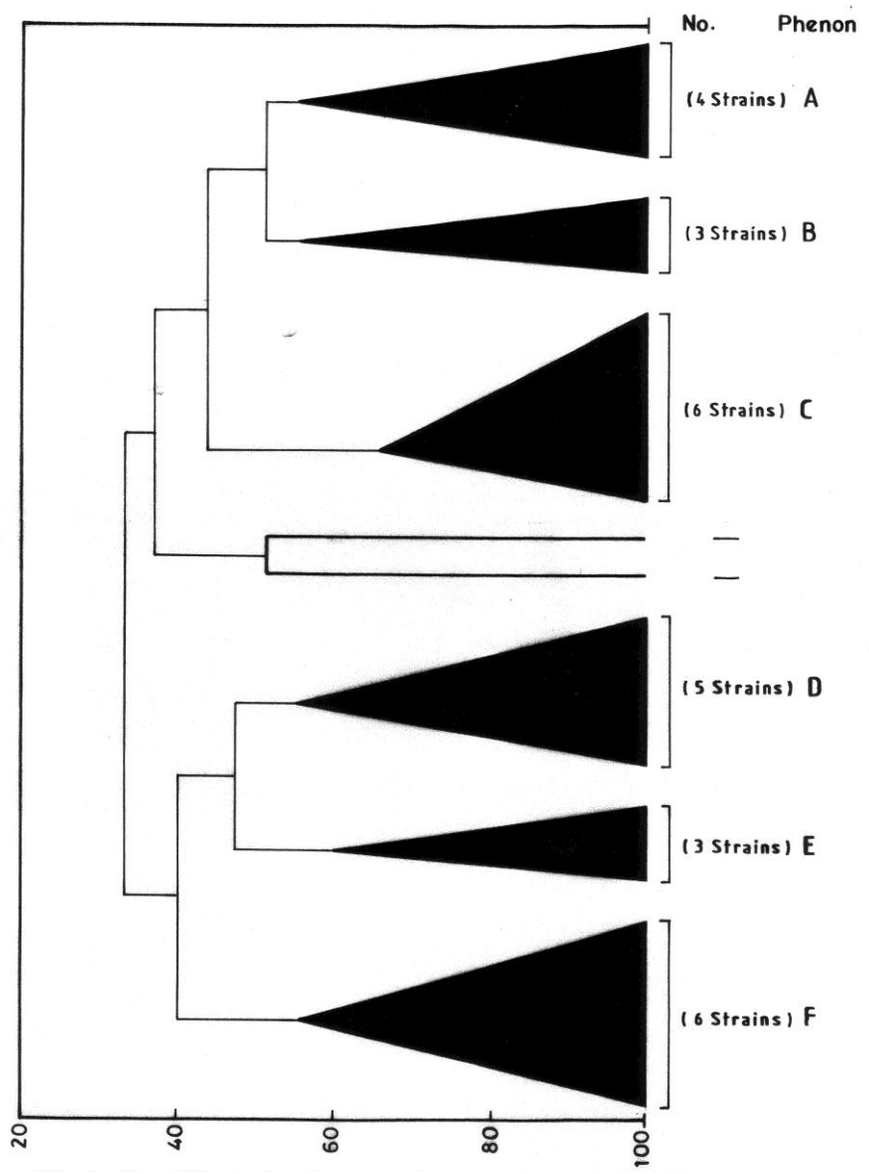


Fig. 1. Simplified dendrogram showing the relationships among phena based on the Ss μ - UPGMA analysis.

Table 2. Characteristic features of the identified actinomycetes isolates

Characteristic	St.m	N.b	S.a	S.g	S.r	S.gn
Morphological characters						
Substrate mycelium	yellow brown	Light yellow brown	reddish brown	yellowish brown	reddish brown	gray yellowish brown
Aerial mycelium	gray	Pink gray	gray	gray	gray yellowish pink spiral	gray
Spore chain /conidia	Long straight filament of conidia showing at regular intervals side branches	Aerial conidial chain	spiral	spiral and rectiflexible		spiral
Diffusible pigments	gray yellowish brown	-	-	-	-	-
Melanin production	-	-	+	-	-	-
Cell wall hydrolysate						
Diaminopimelic acid	LL-DAP	meso-DAP	LL-DAP	LL-DAP	LL-DAP	LL-DAP
Sugar pattern	nd	arabinose & galactose	nd	nd	nd	nd
Utilization of carbohydrates						
D-glucose	+	+	+	+	+	+
D-fructose	+	+	+	-	+	+
L-arabinose	-	-	+	+	+	+
Lactose	-	-	+	+	+	-
Raffinose	-	-	+	-	-	-
Rhamnose	+	-	+	-	+	+
Mannitol	+	-	+	-	+	+
Galactose	+	+	+	-	+	+
Xylose	+	+	+	+	+	+
Sucrose	+	-	-	-	-	-
Trehalose	nd	+	+	nd	+	+
Maltose	+	-	+	+	+	-
Cellobiose	+	nd	nd	-	nd	nd
Sodium citrate	+	+	-	-	-	+
Sodium acetate	+	nd	nd	+	nd	nd
Ribose	-	-	-	nd	-	-
Mannose	nd	+	nd	nd	+	+

Table 2. Continued

Characteristic	St.m	N.b	S.a	S.g	S.r	S.gn
Utilization of amino-acids						
L-cysteine	+	-	-	+	-	+
L-valine	+	+	-	-	+	+
L-alanine	+	+	+	+	+	-
L-phenyl-alanine	+	-	-	-	-	-
L-arginine	+	+	+	+	+	+
L-tryptophane	+	-	-	+	-	-
Biochemical activities						
Catalase	-	-	-	+	-	-
Urease	+	+	+	+	+	+
Nitrate reductase	+	+	+	+	+	+
Sulfide precipitation	+	-	-	-	+	-
Indole test	-	-	-	-	-	-
Degradation of						
Starch	+	+	+	+	+	+
Cellulose	-	-	+	-	-	+
Chitin	-	-	-	-	-	-
Gelatin	-	-	-	-	-	+
Tyrosine	+	+	-	+	+	+
Xanthin	+	+	+	+	+	+
Esculin	+	+	+	+	-	+
Degradation of						
Protein	+	+	+	-	+	+
Lipid	+	+	+	-	+	+
Pectin	+	nd	+	-	nd	nd
Lecithin	+	+	+	nd	+	-
Antibiosis against						
E.coli	-	-	+	-	+	+
S.aureus	-	-	-	-	+	+
B.subtilis	-	-	+	-	-	+

Table 2. Continued

Characteristic	St.m	N.b	S.a	S.g	S.r	S.gn
Resistance to						
Phenol (0.01w/v)	-	-	-	-	-	-
Phenol (0.002 w/v)	+	+	+	+	+	+
Crystal violet (0.001 w/v)	-	+	-	-	+	+
Crystal violet (0.0001 w/v)	+	+	+	+	+	+
Sodium azide (0.01 w/v)	-	-	-	-	-	+
Gentamycin (998 µg.ml-1)	-	+	-	-	-	-
Erythromycin (804 µg.ml-1)	-	-	-	-	-	-
Tetracycline (990 µg.ml-1)	-	-	-	-	-	-
Cefixine (997 µg.ml-1)	-	-	-	-	-	-

St.m : *Streptovercillum morookaense*

N.b : *Nocardia brasiliensis*

S.a : *Streptomyces alanosinicus*

S.g : *Streptomyces globosus*

S.r : *Streptomyces rubber*

S.gn : *Streptomyces gancidicus*

nd : not detected

DAP : Diaminopimelic acid

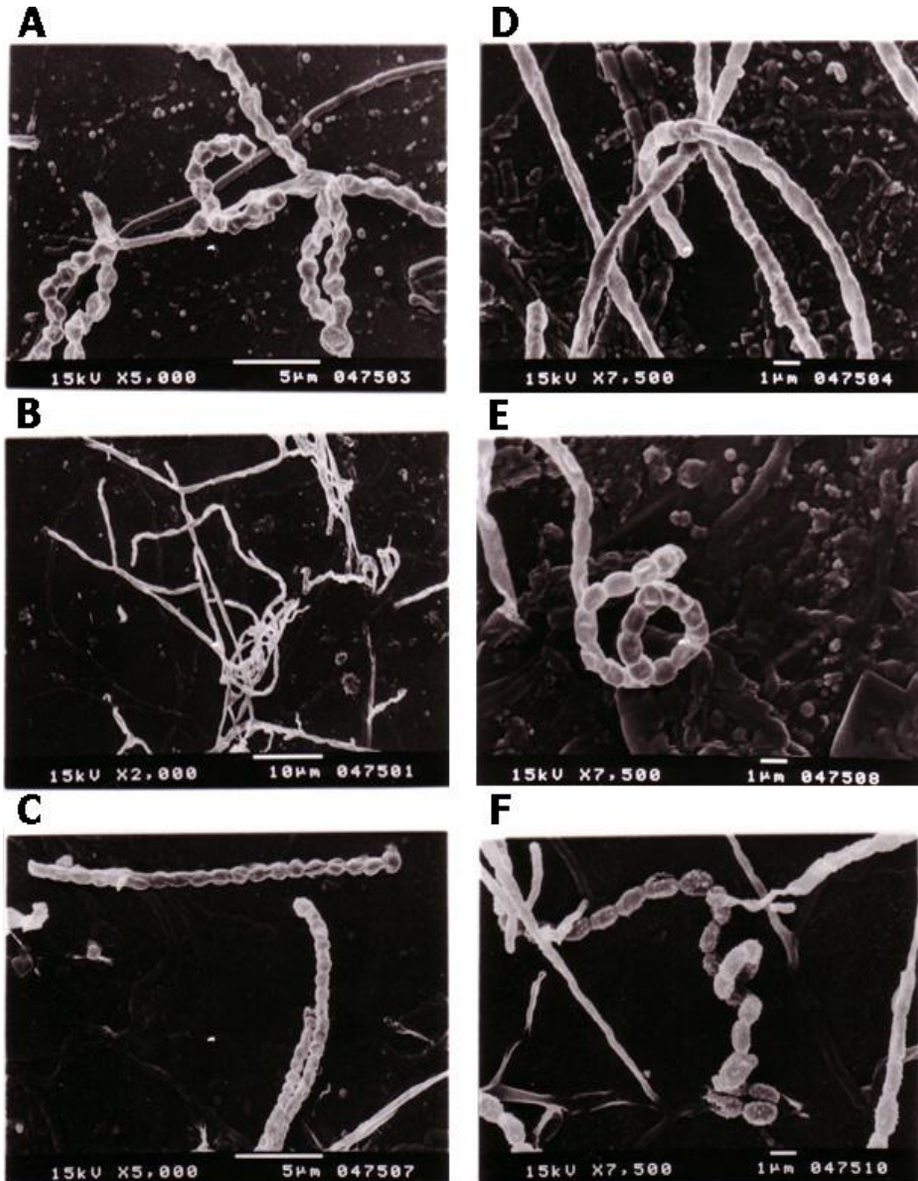


Fig. 2. Scanning electron micrographs of representative strains from each phena *Streptovercillum morookaense* (A); *Nocardia brasiliensis* (B); *Streptomyces alanosinicus* (C); *Streptomyces globosus* (D); *Streptomyces ruber* (E); *Streptomyces gancidicus* (F)

lacking melanin pigments, positive for urease and nitrate reductase, degraded starch, tyrosine, xanthine and esculin, in addition to the number of carbon sources utilized (Table 2). So it could be stated that actinomycete members of this phenon are closely related to *Nocardia brasiliensis*.

Phena C, D, E, and F: It is widely accepted that the genera *Streptomyces* and *Streptovercillum* are closely related. Both have cell-wall I (Lechevalier 1989), a high content of GC in their DNA (Pridham and Tresner 1974) and high similarities in DNA homology (Kroppenstedt *et al* 1981), and are lysed by the same phages (Wellington and Williams 1981). Additionally, they contain similar menaquinones, fatty acids and polar lipids (Lechevalier 1989). The only support for distinguishing between *Streptomyces* and *Streptovercillum* has come from DNA-DNA pairing studies (Gladek *et al* 1985).

The presence of LL-diaminopimelic acid in the cell wall of isolates belonging to phena C, D, E and F with no diagnostic sugars is consistent with the proposal that these strains belong to the genus *Streptomyces* and related genera (Williams *et al* 1989; Hensyl 1994). As pointed out by Williams *et al* (1983), there is no simple, rapid procedure for objective identification of streptomycetes. The problems of streptomycetes identification are largely a reflexion of the difficulties in streptomycetes taxonomy. Morphological or pigmentation characteristics were not included in streptomycetes taxonomy by Kampfer *et al* (1991). It is generally accepted that more than one category of spore chains can be observed in the same species and the distinction between recti-

flexibles and spirals is not clear (Williams and Wellington 1980; Williams *et al* 1989). The determination of the color of the spore mass is also not easy (Kutzner, 1981), especially with respect to spore color and other morphological criteria. These traditional characters, which are often difficult to determine (Williams and Wellington 1980; Kutzner, 1981), are inadequate for classification and identification. Carbon source utilization tests have been found in earlier studies to be characters with great differentiation potential. The utilization of sucrose, L-arabinose, inositol, mannitol, rhamnose and raffinose was recommended as an aid to species differentiation in the International *Streptomyces* Project (Shirling and Gottlieb, 1972) and these tests have been used in several identification schemes (Nonomura, 1974; Szabo *et al* 1975). The phenetic and diversity of streptomycetes are major problem and as pointed out by Kampfer and Kroppenstedt (1991), streptomycetes taxonomy is still developing.

Phenon C: The 6 isolates grouped in phenon C were clustered at 65% similarity level (Fig.1).The chemotaxonomic properties and morphological characteristics of our experimental isolates are consistent with the reference strain *Streptomyces alanosinicus* (gray series) Williams *et al* (1989). Both possess the same cell wall chemotype; produce gray aerial mycelia carrying gray spiral and rectiflexible spore chains (Fig. 3C). No observed motility and melanin pigments not produced. Both utilized arabinose, fructose, galactose, glucose, mannitol, raffinose and xylose but no utility to sucrose (Table 2).

Phenon D: Five isolates are grouped in this phenon at similarity level 55%. The strain selected from this phenon showed phenotypic characteristics (Fig. 2D) closely related to the reference strain *Streptomyces globosus*. Their aerial mycelia bear spiral and rectiflexible (straight to flexuous) spore chains forming gray spore masses. The cell wall contained LL-DAP with no sugars; absence of motility; glucose, L-arabinose, lactose and xylose were utilized by the selected strain but can not utilized fructose, galactose, mannitol, raffinose, and sucrose. (Table 2). On the basis of the previous collected data and in view of the comparative study of the recorded properties of the tested strain in relation to closet strain, the recommended identification may be closely related to *S. globosus*..

Phenon E: At 60% similarity level this phenon included three isolates. As reported in Table (1) the representative isolate (Fig. 2 E) was closer to the reference strain *Streptomyces rubber*, having grayish-yellowish pink spore masses, the spore chains were spiral, and the substrate mycelia were reddish brown. Both strains have the same cell wall chemotype. No detection to motility or melanin production. Both of them utilized glucose, D- fructose, L-arabinose, mannitol, rhamnose and xylose (Table 2).

Phenon F: At 55% similarity level, six isolates constituted this phenon, the selected isolate and the reference strain *Streptomyces gancidicus* shared in most characters. Their spore masses were in the gray series with gray yellowish brown substrate mycelia. Spore chains are spirals (Fig. 2F). Their cell hydrolysates contained LL-DAP with no diagnostic

sugars. No motility. Absence of melanin. Both of them were able to use arabinose, fructose, glucose, mannitol, galactose, and xylose where as they were unable to utilize raffinose and sucrose (Table 2).

Conclusion

The data obtained from examined fresh water habitat in Burullous Lake clearly showed that the Burullous Lake has its own distinct actinobacteria community. The lake is characterized by high representation of Streptomyces species from different series, and fewer numbers of nocardioforms were observed.

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مجلة اتحاد الجامعات العربية للدراسات والبحوث الزراعية ، جامعة عين شمس ، القاهرة ، ١٤ (١) ، ٨٧-١٠٣ ، ٢٠٠٦

دراسة عددية عن بعض الاكتينومييسيتات المعزولة من بحيرة البرلس بمصر

[٦]

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تجمعت وكونت ٦ مجموعات وجد فقط عزلتين منفصلتين عن هذه المجموعات وتم تعريف عزلة ممثلة لكل مجموعة وكانت العزلات المعروفة هي

Streptovercillum morookaense, *Nocardia brasiliensis*, *Streptomyces alanosinicus*, *Streptomyces globosus* and *Streptomyces gancticus*

تم عزل ٢٩ سلالة تنتمي الى مجموعة الاكتينومييسيتات من رسوبيات بحيرة البرلس وأجريت لها دراسة عددية وتصنيفية بعد اختبارها في ٦٢ صفة تصنيفية اشتملت على الشكل المورفولوجي والنمو الغذائي والخواص الفسيولوجية وتم تحليل النتائج احصائيا باستخدام الطرق العددية وعند ٥٤% من التماثل ووجد ان معظم العزلات

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